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Digital biology approach for macroscale studies of biofilm growth and biocide effects with electron microscopy†

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Microbial interactions are one of the major topics of current research due to their great societal relevance. It is now established that biofilms—associations of microorganisms, exchanging various chemical compounds, including proteins and nucleic acids—are capable of promoting horizontal transfer of resistance genes. However, our understanding of the processes occurring in biofilms is rather limited. A possible method to partly overcome this problem is the implementation of highly efficient imaging and mapping of these structures. This work proposes a combination of automated scanning electron microscopy (SEM) and a comprehensive software system that uses deep neural networks to perform an in-depth analysis of biofilms. Time-dependent, high-throughput mapping of biofilm electron microscopy images was achieved using deep learning and allowed microscale data analysis of visible to the eye biofilm-covered area (*i.e.*, at the macroscale). For this study, to the best of our knowledge, the first matrix and cell-annotated biofilm segmentation dataset was prepared. We show that the presented approach can be used to process statistical data investigation of biofilm samples in a volume, where automation is essential (>70 000 separate bacterial cells studied; >1000 times faster than regular manual analysis). To evaluate the approach, multiple time steps of biofilm development were analyzed by first-to-date kinetic modeling of biofilms with SEM, revealing the complex dynamics of biofilm formation. Moreover, it was shown that the described procedure is capable of capturing differences between antibiotics and antimicrobial compounds applied to studied biofilms.

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Introduction

There is a growing crisis of antibiotic resistance that is causing several health issues worldwide.¹ The reason for this may be both the irrational use of antimicrobials and their application to prevent a combined bacterial–viral infection. One of the ways for bacteria to endure antibiotic treatment is to form biofilms.^{2,3} Understanding the mechanisms of biofilm formation and the contribution of these processes to increasing levels of antibiotic resistance are critical public health priorities in developing new antibiofilm agents and methods to combat antimicrobial resistance.

The implementation of high-tech innovations in biological research is currently making a significant contribution to the development of life science.⁴ Digital biology solutions provide insights into various scientific problems, such as climate change,⁵ genome annotation,⁶ biological image analysis,^{7,8} protein folding,⁹ drug discovery,¹⁰ cancer detection,^{11,12} biology laboratory virtualization,¹³ and the problem of antibiotic discovery.^{14,15} In fact, the combination of models, that predict antimicrobial activity of molecules or generate novel compounds with the high-throughput mapping of experimental microscopic data may become a powerful strategy for the accelerated discovery of antibiofilm agents in the near future.^{16–18}

There are several definitions of biofilms.^{19–21} Generally, biofilms are highly structured associations of microorganisms attached to the surface (which can be both biotic or abiotic) or forming floating mats on liquid surfaces.²² These associations are contained within a self-producing matrix of extracellular polymeric substances (EPS) consisting of proteins, lipids, nucleic acids, and polysaccharides. These compounds play an important role in enhancing adhesion to the surface,

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aggregating microorganisms, and ensuring the structural integrity of the biofilm.^{23–25}

Biofilms form on industrial production lines, heat exchangers, and work surfaces, leading to corrosion and damage to mechanisms and contamination of raw materials and products.²⁶ For the food industry, contamination with biofilms can lead to much more severe effects,²⁷ contributing to outbreaks of foodborne infections.^{28,29} Biofilms are among the drivers of the growing antibiotic resistance crisis and account for two-thirds of all infections.³⁰ The National Institute of Health estimates that biofilms cause 65–80% of all microbial infections and 80–90% of all chronic infections, making biofilms a major public health problem.³¹ Diseases associated with biofilms include upper and lower respiratory tract diseases, endocarditis, chronic otitis media, eye infections, chronic wounds, diabetic foot ulcers, urinary tract infections, and periodontitis.^{32,33} Bacterial colonization of medical devices such as intravascular and urinary catheters, pacemakers, heart valves, contact lenses, breast implants, endotracheal tubes, and orthopedic implants can lead to device-associated infections.^{34,35}

Various methods carry out the detection of biofilms: staining with further detection using photometric methods; various methods of microscopy, including staining, using fluorescent labels; molecular genetic techniques for detecting the expression of biofilm-forming genes, *etc.*³⁶ There are several established techniques for the automated computational analysis of biofilm images using confocal microscopy,^{37–39} some of which can even handle statistical area analysis.⁴⁰ Automated detection of stalked bacteria is also available.⁴¹ However, there is currently no single method or test system for the comprehensive AI-based automatic detection of biofilms and their conditions with Scanning Electron Microscopy (SEM), which can perform statistical analysis on a vast amount of imaging data, where manual analysis cannot be performed (including single cell counting and matrix quantification). In fact, there are prior studies that have measured cellular properties inside biofilms⁴² and the matrix distribution⁴³ based on fluorescence images. However, the mechanisms and processes of biofilm formation are not well understood, and studies that detail the morphological changes in the bacterial population in the process of bacterial colonization are still limited.

With the development of automation techniques in electron microscopy, it has become an appealing imaging method. Currently, SEM imaging is widely used as a quality control method in semiconductor manufacturing and steel production.⁴⁴ Furthermore, the combination of energy-dispersive X-ray spectroscopy with automated SEM imaging can be used for the elemental analysis of materials⁴⁵ or pharmaceutical products.⁴⁶ Contaminant identification, surface topography investigation, and defect detection are also offered with electron microscopy and find application in different areas of science and technology, *e.g.*, solar panels.⁴⁷

Microbiology is the area where electron microscopy can also be employed.^{48–50} SEM has been shown to be a suitable tool for tracking bacterial adhesion and biofilm formation on biotic⁵¹ and abiotic surfaces.^{52,53} SEM has the levels of magnification

and resolution required to visualize individual cells of microorganisms and their communities or biofilms, as well as their spatial organization.⁵⁴ In addition, SEM is a valuable tool for studying the effect of antimicrobial agents on the cell morphology of bacterial populations in various biofilms.⁵⁵ The in-depth method to study biofilms would allow *in silico* high-throughput mapping and detection of the key features of the studied systems: separate cells, channels, and matrix (Fig. 1a). To the best of our knowledge, there are still no computer-aided solutions for SEM image analysis of biofilms that can provide the recognition of matrix and single cell detection without preliminary data annotation with similar or above quality.

The data acquisition rate is currently very high and would not allow a manual analysis of images without increasing research budgets and potential loss of analysis quality due to human errors. Recently, a solution to this problem—the application of deep learning algorithms—has become increasingly widely adopted. Examples of usages include biomedical applications,⁵⁶ analysis of pharmaceutical powders,⁵⁷ protein nanowires,⁵⁸ catalysts,⁵⁹ and analysis in a liquid phase.⁶⁰ Besides segmentation and detection tasks, deep learning-based image inpainting has also performed.⁶¹ Despite the active application of deep learning algorithms in cell imaging,^{62–70} there is a significant lack of knowledge in AI-based biofilm SEM image analysis. State-of-the-art techniques only perform segmentation of whole biofilms without recognition of the matrix⁷¹ with the Trainable Weka Segmentation plugin,⁷² segmentation of cellular compartments,^{73–75} and cell segmentation on SBF-SEM images.⁷⁶ The main reason for this small coverage of possible objects is most likely the unavailability of well-annotated large amounts of data for researchers compared to other types of microscopy.⁷⁷ With the development of frameworks for semi-automatic microbiological data annotation^{78–80} or advanced segmentation deep learning architectures,^{81,82} it is possible to partially reduce the bottleneck with image annotation. However, higher quality data still significantly improve the results and allow other researchers to train their models more effectively.

An alternative method to study biofilm structures is using fluorescence confocal microscopy.⁸³ Unlike SEM, it allows 3D visualization of biofilm architecture and can distinguish between living and dead microbial cells. Although spatial organization of bacterial cells and EPS in the biofilms can be determined by confocal laser scanning microscopy (CLSM), only SEM can show the architecture of biofilms at a single bacterial cell resolution level.⁸⁴ In addition, CLSM lacks the necessary magnification and resolution for detailed observation of individual cells in a biofilm and their morphology.⁸⁵ It also requires fluorescent dyes to evaluate biocide effects, which can affect biofilm physiology (*e.g.* reduction levels of resazurin that sometimes used in CLSM⁸⁶ can be decreased in the presence of antibiotics,⁸⁷ improve the breakthrough of cell membranes,⁸⁸ affect the viability,⁸⁹ lead to decrease of elongation rates⁹⁰) or even damage living cells.⁸⁷ Precise cell counting is a limiting factor for confocal microscopy, as most protocols, which use fluorescent pigments, allow only semiquantitative analysis.^{71,87} Since only 2D imaging with SEM is available, it is possible to



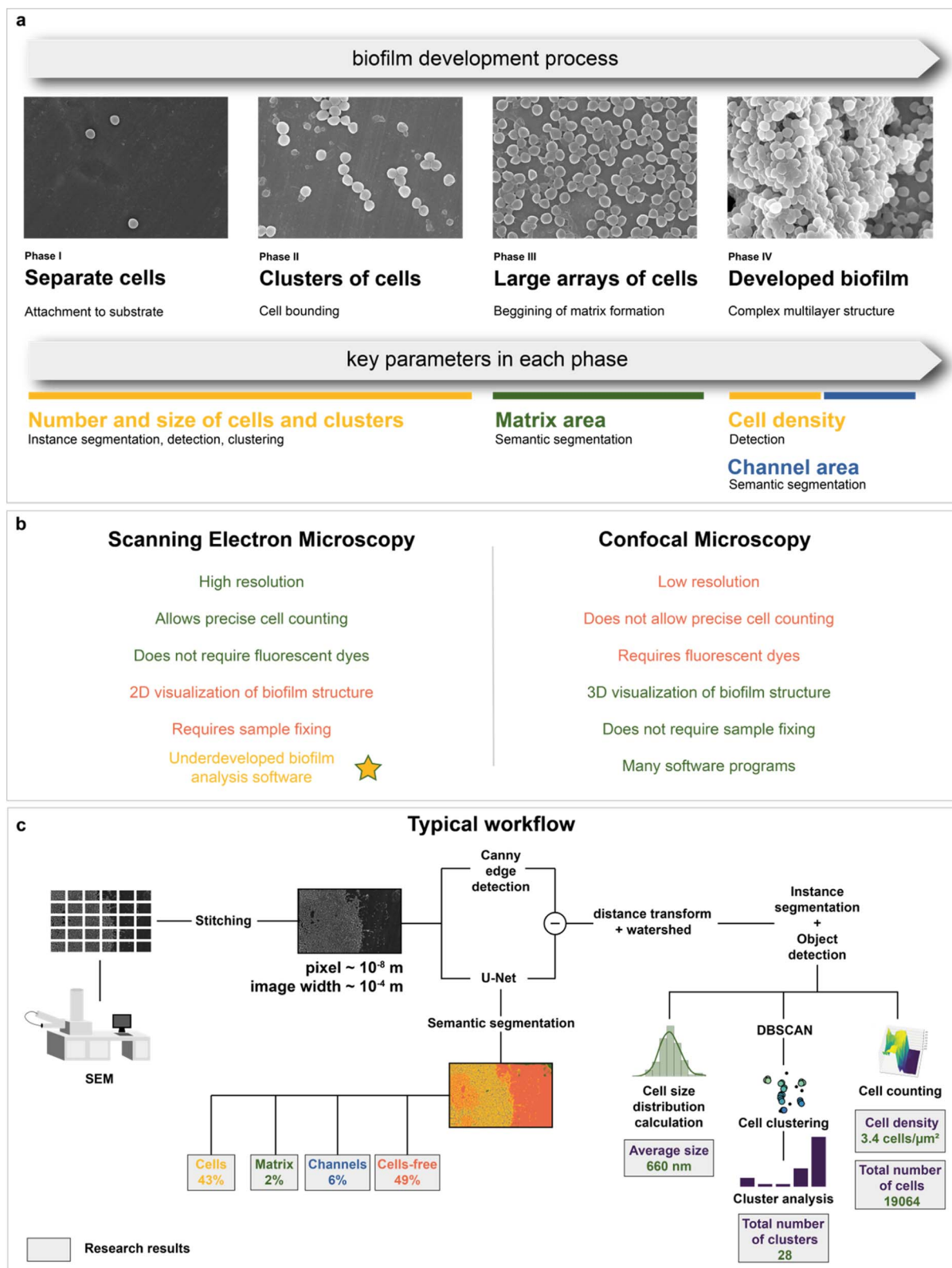


Fig. 1 Biofilm formation and image analysis: (a) biofilm development process; each phase is characterized by key parameters, which can be automatically determined with computer vision techniques; (b) brief comparison of SEM and confocal microscopy for biofilm imaging. Yellow star highlights one of the aims of the research: novel software development for quantitative biofilm analysis with SEM; (c) a typical workflow for deep neural network analysis; a combination of models and algorithms offers ample research opportunities.



calculate only the area density of cells. Nevertheless, it is likely to correlate with the bulk cell density, allowing evaluation of growth and biocidal effect (see “Kinetic modeling of biofilm growth” and “Automated mapping of large volumes of SEM images for the investigation of antimicrobial compound impact” in the Results and Discussion section for more information). A comparison of SEM and confocal microscopy is shown in Fig. 1b. Both methods have their own pros and cons⁹¹ and are often used together.⁹² However, there is much more software for quantitative analysis using confocal microscopy compared to electron microscopy.

Despite the fact that deep learning-based approaches are more efficient than the popular image thresholding, neural network solutions are not common at the moment. It slows down the digital transformation in biofilm studies. Implementation of novel computer vision techniques allows fast calculation of cells and matrix on images making possible to evaluate biofilm growth and biocide effects with a greater statistical accuracy.

Therefore, the main purpose of this work is to propose a digital approach to studying biofilm structures with SEM. It combines automated scanning electron microscopy with deep neural network analysis (Fig. 1c). As a result, it became possible to analyze biofilms with the use of machine intelligence at the macroscale and derive conclusions about their composition (cell, matrix, and channel area calculation) and morphology (cell size distribution, number, and size of cell clusters in the biofilm formation area, total number and density of bacterial cells in the area visible to the human eye). These data were used to study biofilm growth dynamics from SEM images. The composition of biofilms after antibiotic and antimicrobial compound application was also investigated with neural network image segmentation. This work not only proposes a method to perform automated image analysis but also shows how it can be implemented on case studies, where the analysis of large volumes of microscopic data is required.

Results and Discussion

Object of study

Staphylococcus aureus was chosen as a model microorganism, as it is one of the main etiological agents of nosocomial infections and is well known for its ability to form biofilms on host tissues and implants.⁹³ Biofilm formation of *S. aureus* often leads to chronic infections in patients suffering from osteomyelitis, endocarditis, cystic fibrosis, or in patients undergoing medical procedures such as catheterization.^{94,95} All of the above makes *S. aureus* one of the main human pathogens and the major microorganism for biofilm research. Moreover, *Staphylococcus* is a convenient object of study due to its cell shape, which facilitates its image processing (Fig. 2a and b).

To demonstrate differences between nascent and mature biofilms, Fig. 2c shows biofilms formed by the *S. aureus* ATCC 6538 strain at different cultivation times. After 24 hours of cultivation, spherical cell conglomerates with a small amount of matrix and clearly visible channels were observed. After 72 hours of biofilm cultivation, the cells were observed to be

almost completely covered with matrix, and the outlines of individual cells were blurred.

The total biofilm formation dynamics are shown in Fig. 2d. After 3 hours of cultivation, single cells or groups of cells (cell clusters) were observed. After 6 hours, cell clusters connected by intercellular bridges appeared. After 12 hours of cultivation, connected cell layers formed multilayer structures with a clearly distinguishable matrix. A complex multilevel structure with a large amount of matrix was observed the following day. After 72 hours, the formed biofilm became a multidimensional structure, where numerous channels were observed. Cells immersed in the extracellular matrix were clearly distinguishable. Visible differences in biofilm morphology at different cultivation times allow us to recognize biofilm morphology without additional methods.

As an example of a typical biofilm development process, area analysis of example biofilm images was performed (see section “Model predictions” in the ESI† for the segmentation results). The results showed a sharp increase in bacterial area and a decrease in cell-free area. Matrix enlargement did not occur until 12 hours after the start of cultivation and reached a plateau after three days. Channel areas also showed an increase after 24 hours; however, their area share was insignificant compared with 2 dominant classes: cells and matrix.

Deep learning methodology

As mentioned above, deep learning is actively used in pattern recognition tasks, such as semantic segmentation and object detection. Convolutional neural networks have made remarkable progress in computer vision.⁹⁶ They are based on convolution operations, which convert the original image into a set of feature maps. This set stores correlation information between closely located pixels and provides detection of distinctive local motifs, making it easier to train the model. Moreover, the nature of the convolution operation makes it easy to use graphical processing units to speed up the training process. Convolutional neural networks dominate computer vision tasks over classical machine learning and image processing algorithms.

A typical neural network-aided computer vision pipeline includes the following steps: labeling the data, preprocessing the images, optimizing the network hyperparameters, and training the final model. Data labeling is a step of identifying specific objects in research data. SEM images of biofilms were manually labeled into masks using a special platform by research scientists with good domain knowledge (Fig. 2a). Segmentation masks included bacteria, matrix, channels, and support areas without cells (cell-free zone) (Fig. 2b). Channel zone differs from cell-free zone by relatively small size and specific location inside large clusters of cells (see section “Channel zone recognition” in the ESI† for detailed explanation). The main task of the segmentation network is to predict the areas of each class on images (Fig. 3a). The training dataset size was equal to 72. Although, it is possible to train a segmentation model using only 4 dissimilar high-resolution training images and proper augmentations with less than 10% loss in

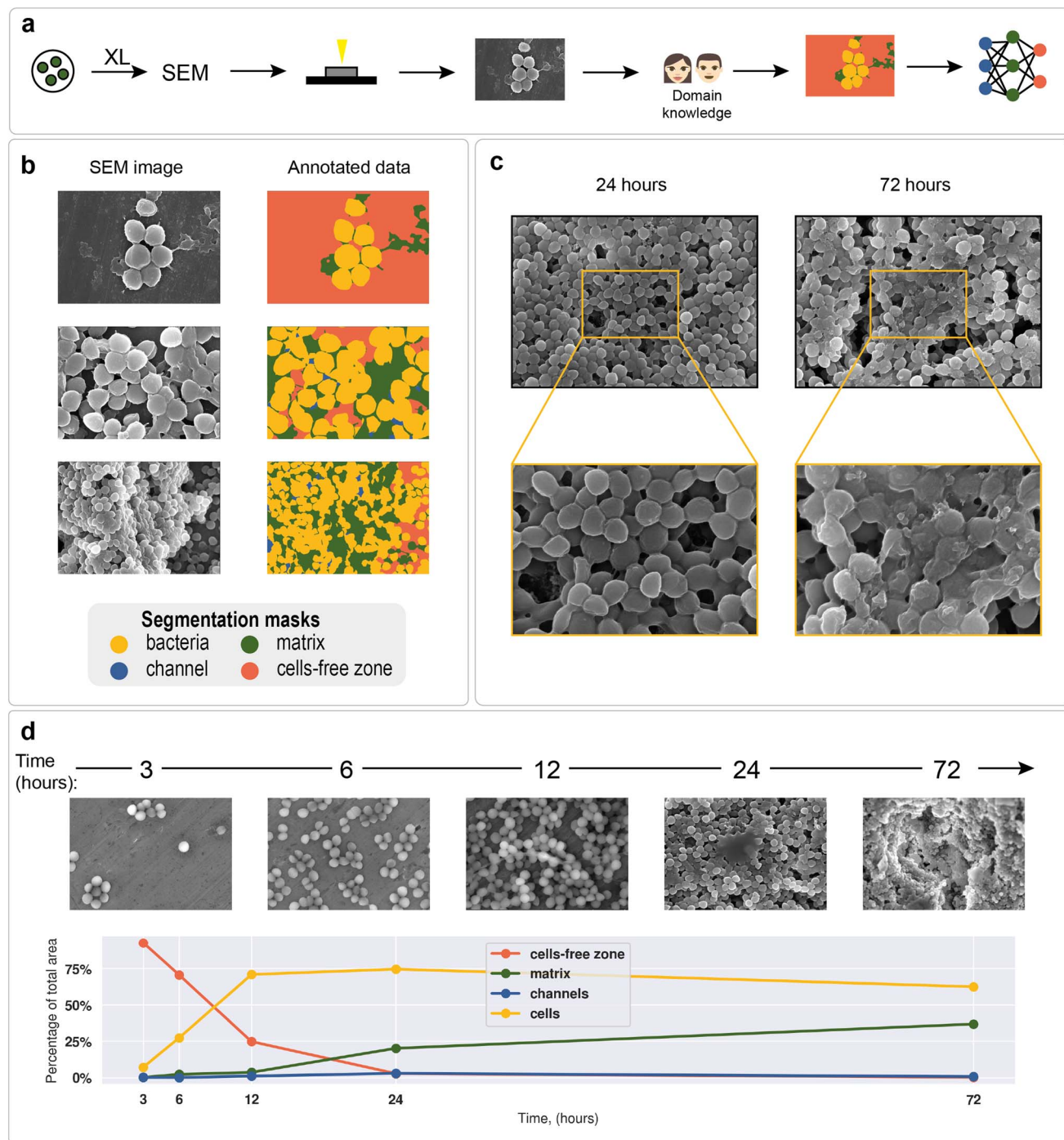


Fig. 2 SEM images of biofilms: (a) neural network training data preparation process; (b) examples of annotated data on different images; (c) *S. aureus* ATCC 6538 biofilms; biofilm obtained after 24 hours of cultivation on the surface of a dense nutrient medium (left); biofilm obtained after 72 hours of cultivation on the surface of a dense nutrient medium (right); (d) *S. aureus* biofilm formation dynamics with cell, matrix, channel, and cell-free zone area statistics; results are obtained with a U-Net segmentation neural network.

quality. High resolution and rich content of training images leads to convergence of the IoU score (Fig. 3b) vs. training dataset size curve despite the seeming lack of training images (see “Neural network implementation and training” in the Methods section for more information). It is worth mentioning that bacterial image labeling is not a clearly defined task.

Different specialists can obtain different results, impacting the performance of the model. Image preprocessing includes modification techniques to facilitate model training (normalization, resizing, augmentations, *etc.*). Hyperparameter optimization is a step of choosing the best combination of hyperparameters (model architecture, encoder, learning rate,



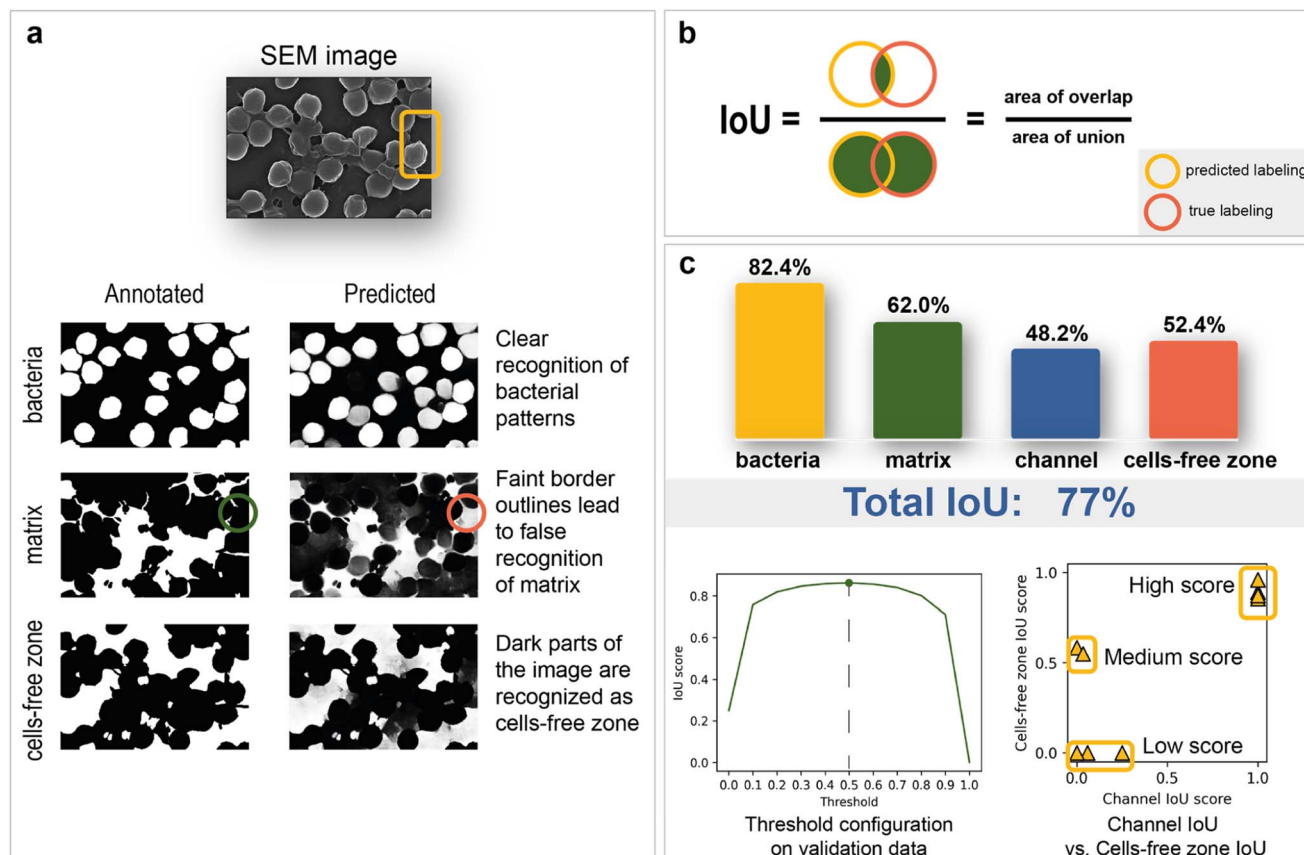


Fig. 3 Training and inference results: (a) predicted images and interpretation; (b) Intersection over Union (IoU) score explanation; (c) model results grouped by segmentation masks with channel IoU vs. cells-free zone IoU scatter plot and optimization plot of binarization threshold.

etc.) for the neural network. The final step is to train a neural network with optimal hyperparameters by minimizing the chosen loss function using a backpropagation algorithm (see “Neural network implementation and training” in the Methods for detailed information).

Model inference results

The total IoU score on the test data is 77%. The disparity between scores on validation (85%) and test is caused by the variations between image sets. The IoU scores for each segmentation class are shown in Fig. 3c. The best result of 82.4% is observed for bacterial cells, while channel areas are predicted to be the worst, with a result of 48.2%. The matrix obtained the second-best metric of 62%. It is worth noting that the IoU score is sensitive to the imbalance of segmentation classes, which can underestimate the true quality of the network. As the channels on images are not clearly defined and can easily be confused with areas without cells, the correlation between channel and cell-free zone segmentation quality was investigated with an IoU score scatter plot (Fig. 3c). Test data are divided into three groups (low score, medium score, high score):

1. The low-score group is distinguished from the others by having no cell-free zones, which leads to many false positives.

2. The medium-score group—lack of channel zones along with a small area of support.

3. The high-score group tends to have a small channel area and thus reduces false negatives associated with the predisposition of the model to label segments as cell-free zones.

The relationship between the IoU score and binarization threshold was investigated on validation images. As seen in Fig. 3c, the optimal threshold has a value of 0.5.

The segmentation network also shows significantly better results on test images compared to classical computer vision algorithms (Table 1). In summary, this work's final model is

Table 1 Cell area segmentation results for CV algorithms on test images

Algorithm	IOU score
Global thresholding ($\nu = 127$)	36.3%
Otsu's thresholding	45.6%
Adaptive Gaussian thresholding	49.5%
Adaptive mean thresholding	49.5%
Adaptive mean thresholding + closing	51.2%
Edge-based segmentation	27.9%
CNN-based segmentation (our approach)	82.4%



devoid of significant systematic errors, making it a viable tool for morphology analysis.

Microscale analysis of macroscale area

Microscopy is a local method, that is, the data obtained depend on the choice of the site for image recording. The site chosen may not be relevant to the characterization of the entire sample nor statistically representative. In turn, subjective factors, such as operator fatigue, aesthetic preferences, and many others, can influence the choice of an area for recording an image. The automatic digital operation mode has become available only recently, which makes it possible to eliminate the human factor and subjectivity in choosing the region for image registration. Automated scanning allows the sequential recording of multiple images according to a statistically relevant scheme. After each recording, the imaging area is shifted in a software-controlled manner (without human operator participation) to the neighboring region, followed by automatic adjustment of focus, brightness and contrast. The operator sets only the recording scheme: the required number of images along the *x*-axis and *y*-axis, as well as the overlapping of the recording areas, which can be negative. In contrast to manual human operation, software-controlled microscopy characterization can be performed nonstop for sufficient time (24–96 h, for instance). This is a great step forward, which opens new opportunities in studying biofilms. Here, in the present study, we used automatic image recording in connection with ML automation.

Thus, automated scanning with further SEM image stitching provides the ability to scan macroscale areas (which are more than 0.05 mm in width and distinguishable by the naked eye) at the scale of 1 pixel being equal to 400 nm². The developed neural network was used to characterize the morphology of the biofilm. Image analysis of the 0.14 × 0.08 mm² area included the following:

1. Matrix, cells, channels, and cell-free zone mapping (Fig. 4a)
2. Cell detection (Fig. 5a and b)
3. Statistical analysis of bacterial cell size (Fig. 4b) and population (Fig. 4c)
4. Cell clustering in the region of biofilm formation (Fig. 5c)

The aim of the biofilm microscopy preparation in the work was to maintain the biofilm's structure to the greatest extent possible (see "Sample preparation" in the Methods section). But this technique leads to reduction of cell size when water is removed in the EM vacuum and to an underestimation of the amount of matrix in the biofilm.⁹⁷ However, this distortion is invariant to biofilm samples. Considering this, kinetic modeling and biocide effect evaluation should remain valid, since the decrease of the matrix as a result of dehydration will be observed in all samples approximately equally. However, to achieve higher accuracy with the developed approach, correction factor should be included.

Mapping provides information on how matrix and bacterial cells are distributed on the support. Two dominant segmentation classes are depicted: cells (43% of the total area) and cell-free zones (49% of the total area). The slight presence of the

matrix (2% of the total area) can be explained by the fact that before the registration of images, the cells have not yet had time to release a significant amount of matrix to form a cohesive biofilm. An additional possible reason is the dehydration of the obtained biofilm during sample preparation. Channels are observed over a large area of the image, confirming the process of incipient biofouling.

However, information about the total area of bacteria on the image does not allow us to estimate how many cells are located on support. Cell detection is a key problem for the statistical characterization of biofilms. Unfortunately, single-cell image annotation takes considerable time, as the density in biofilms may reach 3.4 cells per μm². This disadvantage can be eliminated by applying mask postprocessing, which solves the problem of single-cell overlapping and obviates the need for annotation of individual bacteria. The approach is based on unsupervised edge detection with subsequent watershed algorithm implementation, which is actively used for separating different objects in an image, *e.g.*, single cells⁹⁸ (Fig. 5a).

First, Canny edge detection with additional dilation was employed to obtain an edge mask. Then, we inverted the mask and applied a cell semantic segmentation mask obtained with a neural network to ignore noncellular zones (matrix and channel areas are also ignored).

The distance transform and subsequent local maxima detection on the transformed image allows us to find the geometric centers of cells. Finally, watershed segmentation was used to estimate individual cell areas. The results for cell detection of regions with different fields of view are shown in Fig. 5b. Some cells are merged into one bounding box. This outcome is probably due to the low difference in intensities at the edges, which breaks the correct edge detection, or because of incomplete cell division. The overall procedure makes it possible to count the total number of cells in the biofilm and collect statistics.

For example, bacterial cell size analysis in the biofilm can be performed. It is important to avoid regions with severe cell overlap to eliminate the bias caused by the superimposition of one cell on another. The resulting histogram is shown in Fig. 4b. The bacterial size was normally distributed with an average of 660 nm.

Population density heat maps make it possible to estimate the number of cells per unit area in different parts of the biofilm. Automated creation of visualization maps may be useful for future biofilm formation studies. In the example, the 3D plot surface was constructed (Fig. 4c). The positive correlation between area cell density and channel zones helps to establish when and where the biofilm developed. The area cell density on the image varies from 0 cells per μm² (bare support areas) to 3.4 cells per μm² (central zones with an abundance of channels). The total number of cells visible in the image reaches 19 064, according to the model results.

Although computer-aided counting of cells gives a lower bound, since it does not consider the multilayer structure of the biofilm, the approach could be scaled to 3D images, so the error will be corrected. Even though, the area and bulk cell densities should correlate. Thus, the biofouling trend should remain



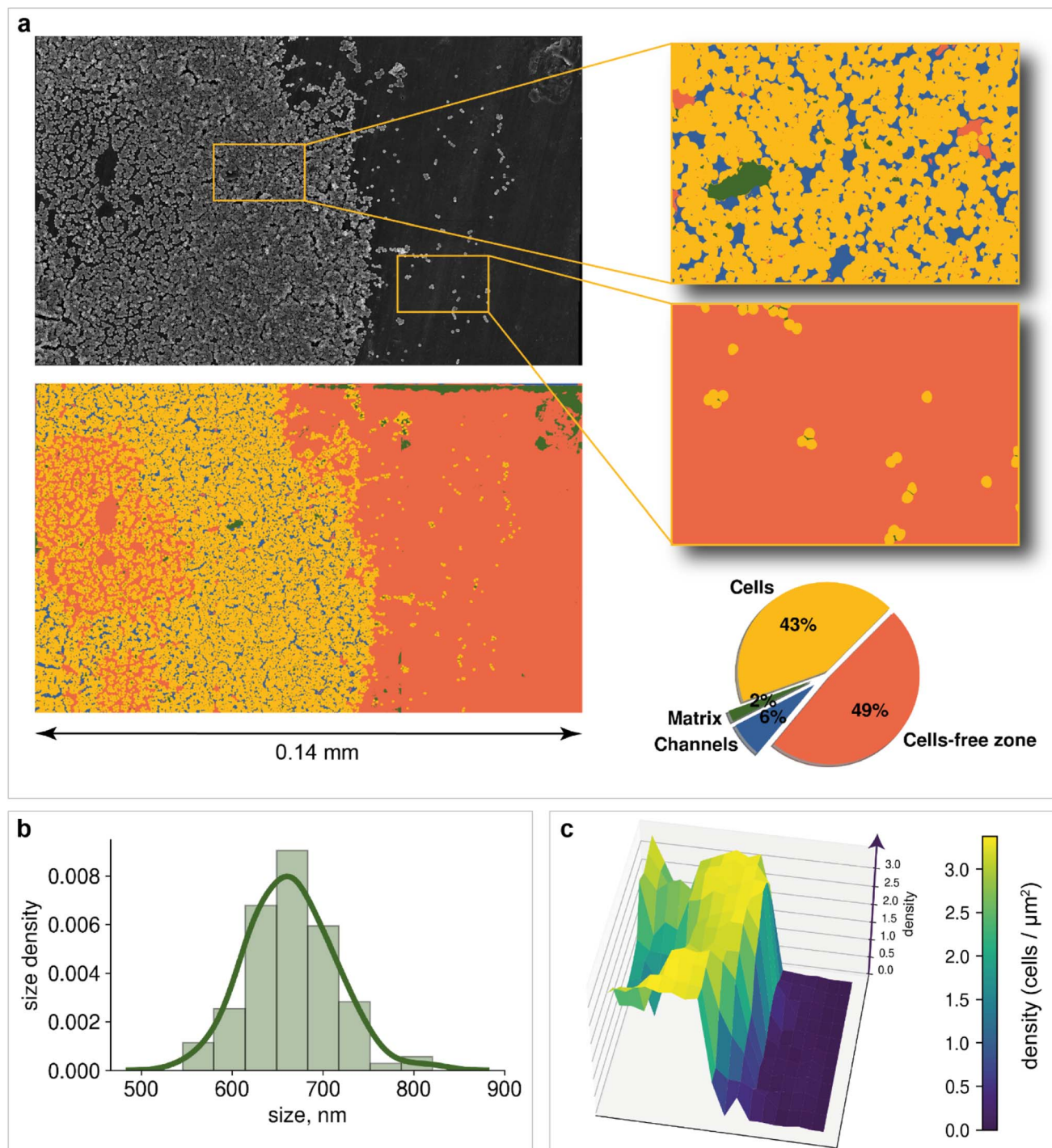


Fig. 4 Morphology analysis: (a) macroscale segmentation of biofilm with surface area calculation; (b) cell size distribution in the region of biofilm formation; diameter sizes were calculated as the principal axes of ellipses, approximated on single-cell segmentation masks; (c) population density heatmap. The z-axis corresponds to the density; the x-axis and y-axis are the image width and height, respectively.

unchanged. The differences in ratios of cells, matrix and cells-free zone should have the same tendency either (see section "Method limitations" in the ESI†).

The creation of techniques for mining statistical data in microbiology research is important for the investigation of biofilm evolution over time. Cell clusters (groups of cells located close to each other) are initiators of biofilm creation, making

their detection a particularly important task. The application of clustering algorithms on top of preprocessed cell detection allows the automatic calculation of the total number of cell clusters and their size (number of cells in a cluster). The approach in this work is based on DBSCAN (density-based spatial clustering of applications with noise). Its main advantage is that one does not need to specify a specific number of

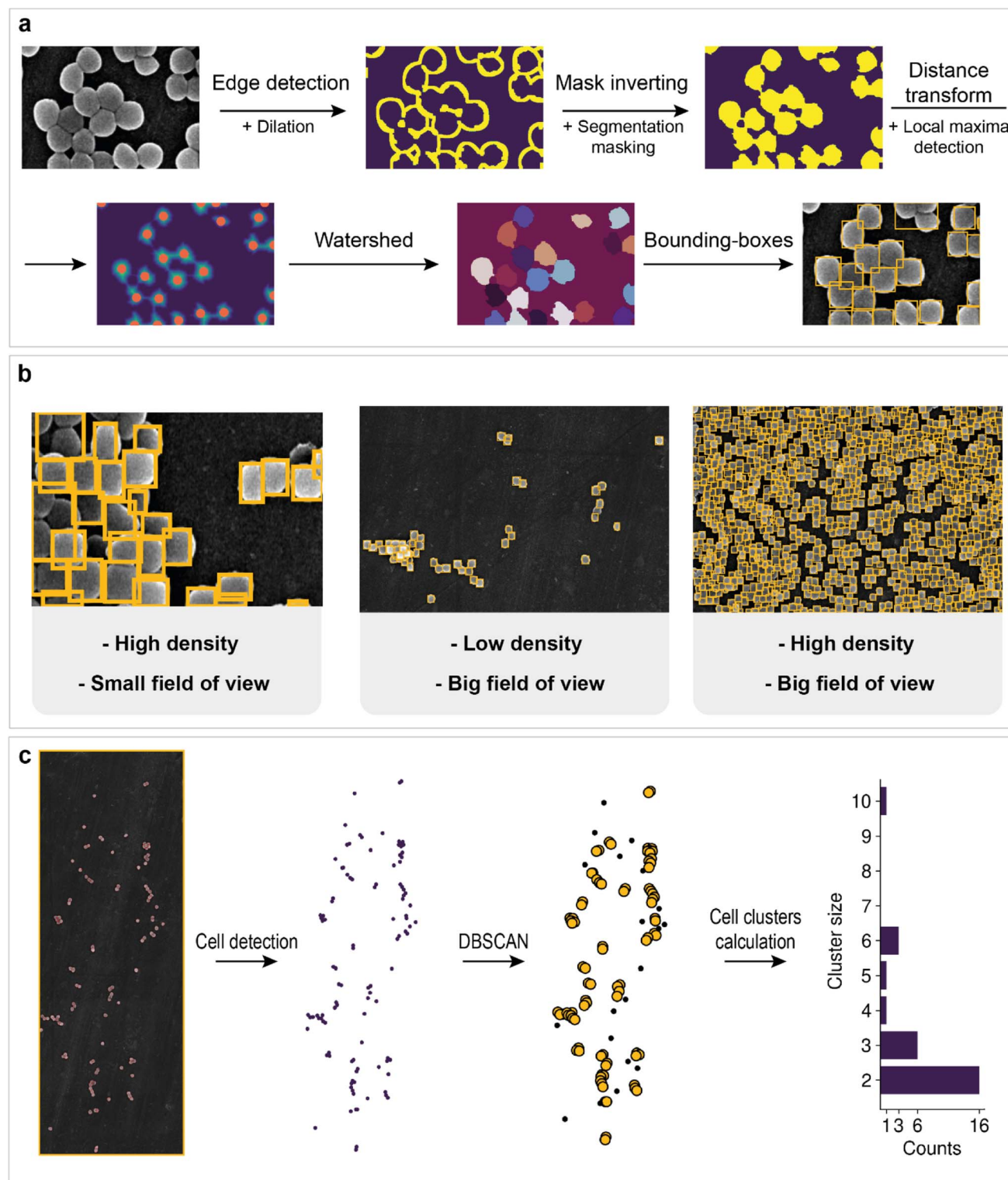


Fig. 5 Cell detection: (a) postprocessing of the cell segmentation mask for detecting individual bacteria; (b) inference of the detection algorithm on images with different fields of view and cell densities; (c) cell clustering approach and results at the site of biofilm formation; black dots on the scatter plot after DBSCAN implementation are separated cells; orange circles are cells in clusters; the final result is a bar chart with numbers of cell clusters of different sizes.



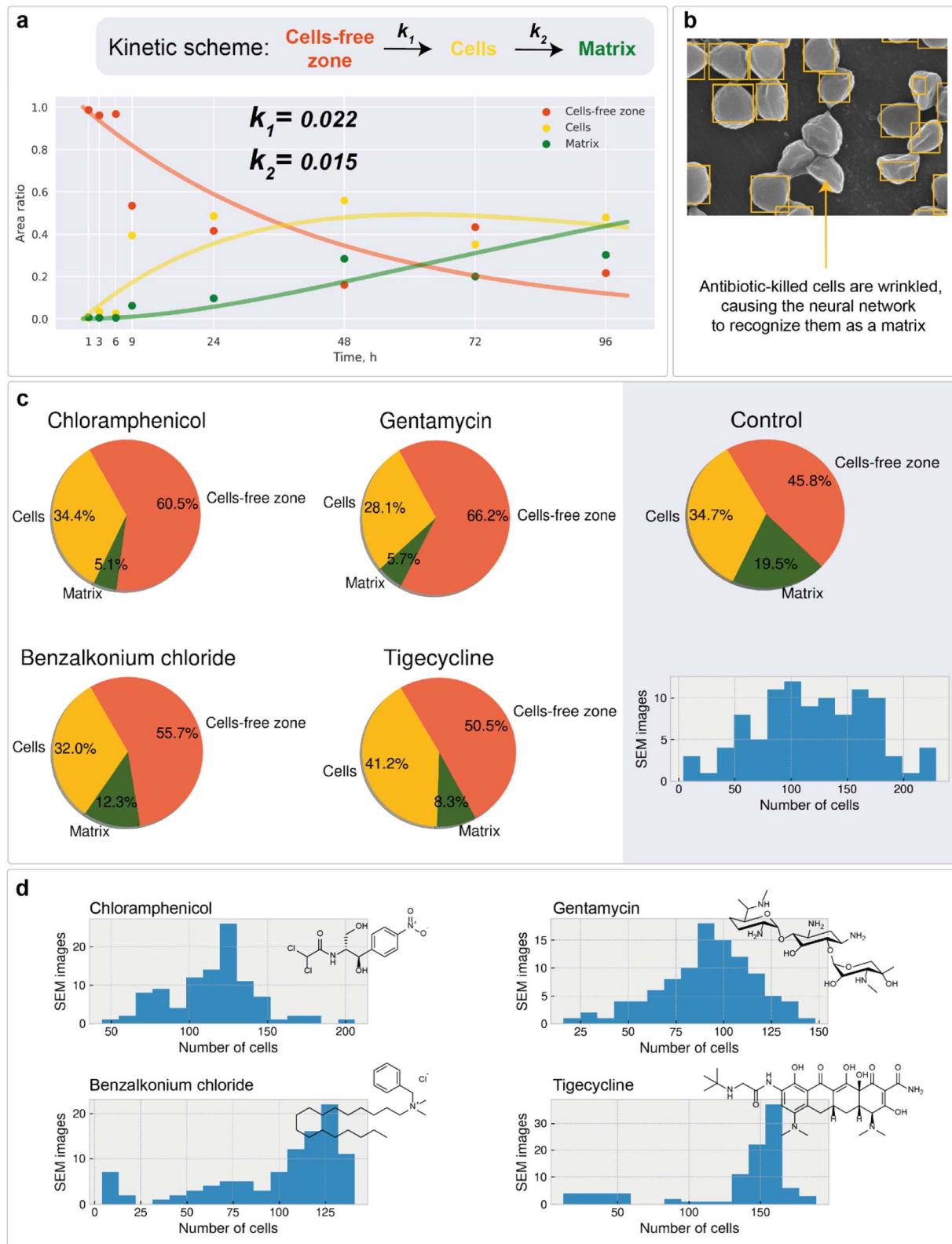


Fig. 6 Kinetic modeling and the study of antibiotic impact on bacteria: (a) kinetic scheme and concentration–time curves for different segmentation classes with calculated rate constants of cell and matrix formation; (b) dead bacterial cell patterns cause bias in cell recognition toward the matrix; (c) area calculations for the biofilm samples treated with different antibiotics; (d) cell counting statistics.

clusters to implement the algorithm. In addition, the possibility of filter noise makes it possible to separate single cells from cells in clusters. The workflow is shown in Fig. 5c. The number of clusters decreases as the cluster size increases, which may indicate the aggregation process in biofilms.

Kinetic modeling of biofilm growth

High-throughput analysis of biofouling requires quantitative characterization of the biofilm growth rate. The biofilm growth kinetics of *Staphylococcus aureus* were investigated by implementing a sequential first-order reaction kinetics scheme (Fig. 6a). 100 biofilm SEM images (square 10×10 with negative overlap) were registered and processed with the segmentation DL model for each period of time (1, 3, 6, 9, 24, 48, 72, and 96 hours, totaling 800 SEM images with a total analyzed area of 0.0832 mm^2 and image pixel size near 100 nm^2). Then, the median ratio of cells, matrix, and cell-free zone for each time period was calculated using segmentation data (median was chosen as a measure of central tendency due to stability towards statistical outliers).

The calculated data show the expected tendency of cell-free zone ratio reduction and cell area enlargement. As the cells emerge, they begin to secrete matrix. The obtained results are then used to estimate the rate constants of cell and matrix formation (Fig. 6a). The rate constants of cell formation ($k_1 = 0.022$) and cell coating with matrix ($k_2 = 0.015$) were established. k_1 is numerically equal to the rate (h^{-1}) of cell area formation at the beginning of the biofilm development process. k_2 is numerically equal to the rate (h^{-1}) of cell coating with the matrix when the support substrate is fully covered with bacterial cells.

These constants can be applied to quantify how biofilms develop over time. For example, the half-life of the cell-free segmentation class (i.e., the time when 50% of the studied area is covered with cells and matrix) amounts to approximately 32 hours. In addition, the time at which the cell area ratio becomes close to the maximum possible is equal to 62 hours (all kinetic calculations can be found in "Kinetic calculations" in the Methods section).

Automated mapping of large volumes of SEM images for the investigation of antimicrobial compound impact

To ensure that the developed algorithms can be applied to study the antibiotic susceptibility of biofilms or the antibiotic activity of novel compounds, high-throughput mapping of SEM images with antibiotic-treated bacteria was performed. The work included the study of chloramphenicol (inhibitor of peptidyl transferase), gentamycin (disturbs tRNA and mRNA interactions), benzalkonium chloride (inhibitor of hyaluronidase), and tigecycline (protein synthesis inhibitor) impacts. For each compound, 100 SEM images of the developed biofilms treated with the antibiofilm agent were recorded. After that, segmentation with the following cell detection was performed on all images. The full data set under study was of the size of 500 images (including control sample) with an approximate total

analyzed area of 0.052 mm^2 having an image pixel size approximately equal to 100 nm^2 .

Unlike normal cells, cells that have died as a result of antibiotic exposure have a curved shape with wrinkles. Despite this, the vast majority of such cells are correctly recognized by the system. However, there are examples where the shape deformation led to the recognition of cells as a matrix (Fig. 6b).

Area calculations (Fig. 6c) on SEM images of antibiotic-treated biofilms show insignificant (less than 1%) areas of channels and dominant cell-free zone areas, indicating an early stage of development. The distribution of cell counts on images (Fig. 6d) for chloramphenicol and gentamycin are symmetrical, with averages of 120 and 90 cells per image, respectively. The samples have almost similar matrix ratios. The distribution of cell counts on images for benzalkonium chloride is skewed to the right, indicating the presence of regions with uneven distribution of cell density. The tigecycline sample is also characterized by a rightly skewed cell per image distribution. In all samples, the number of images with a large number of cells (150+ units) is less than in the control sample. These results prove the applicability of the investigated antibacterial drugs in combating the formation of biofilms.

All samples showed a lower biofilm (cell + matrix) area than the control sample (developed biofilm, 72 hours after cultivation), indicating the presence of biocide effects. The value of the effect decreased consistently in the following series: gentamycin (-20.4%), chloramphenicol (-14.7%), benzalkonium chloride (-9.9%), and tigecycline (-4.7%). The protocol allows the differentiation of such samples, confirming the applicability of the described approach.

Considering the biofilm suppression effect, the following relative order can be suggested based on the reduced matrix area: gentamycin, chloramphenicol ($\sim 5\%$), $>$ tigecycline ($\sim 8\%$) $>$ benzalkonium chloride ($\sim 12\%$). For the combined biofilm and individual cell suppression effect, gentamycin showed the best performance (Fig. 6c) with a more uniform suppression profile (Fig. 6d). It is important to note that these observations should be considered preliminary, since many more samples of different biofilms should be studied to make general conclusions. In addition, killing curve measurements should also be performed. However, the computer vision approach, which is the main topic of the article, will remain unchanged.

The present study developed a methodology for high-speed automatic analysis, including obtaining data for cell growth/inhibition kinetics over time (Fig. 6a), distinguishing individual cells and the matrix (Fig. 6c), and establishing the shape/uniformity of profiles (Fig. 6d). It is a versatile approach for complex analysis and comprehensive treatment of visualization abilities provided by electron microscopy.

Conclusions

In the present study, an automated biofilm SEM image analysis protocol was reported for the first time with direct biofilm characterization at the macroscale. The challenge of macroscale statistical characterization is not only the amount of data but also the depth-insight level needed to reveal morphology



changes. All computational techniques used in the study can be scaled in the future to provide analysis of 3D data recorded with volume electron microscopy. To highlight the difference, we compared the automated approach developed here with a regular manual analysis.

In summary, the pipeline allowed processing, mapping and detection of the following:

1. 1330 SEM images (800 images for kinetic modeling of biofilm growth, 500 images for investigation of biocide effects, and 30 images for mapping of the macroscale biofilm region)
2. 1.15 cm^2 of total analysed area, registered with negative overlaps and 0.1472 mm^2 of total SEM image area.
3. 74 736 separate bacteria cells with the establishment of the size for the fraction of the recognized cells.

The development of a baseline segmentation neural network takes approximately 24 hours for an experienced data scientist. The training procedure with known hyperparameters can be estimated at 5 hours.

Manual data annotation of one SEM image (without labeling of separate cells) takes an average of 25 minutes. Therefore, the annotation of all images that were used would have taken approximately 558 hours for one person, whereas one neural network can provide the mapping of one image with an approximate time of 1 second (1500 times faster than a specialist; see section "Neural network processing time" in the ESI†).

As a result, a combination of fully automated scanning electron microscopy measurements with the use of machine intelligence was developed. Automated SEM enables imaging and mapping of large sample areas, acquiring reliable statistics about sample composition with significantly less time compared to manual labeling. The developed protocol, which combines an image stitching algorithm, a deep neural network for image segmentation, and robust computer vision techniques for object detection and clustering, provides automated analysis of SEM images, yielding areas of cells, cell-free zones, channels, and a matrix, number of separate cells, their sizes, and the number of cell clusters.

The described approach was tested against more than 1000 images of *Staphylococcus aureus*, allowing us not only to study macroscale areas of the sample with nanoscale image resolution but also to study the complex dynamics of biofilm formation and antibiotic tolerance. The digital solution is supposed to be used in SEM biofilm studies, helping to validate novel anti-biofilm compounds with high-throughput computing and statistical significance. Of course, it should be mentioned that the SEM used as an experimental technique and the computational software developed have their own limitations in terms of accuracy, reliability and algorithms. We expect that further studies will elucidate these limitations in more detail and lead to the development of improved techniques.

Methods

Sample preparation

Bacterial strains and cultivation conditions. We used the type strain *S. aureus* ATCC 6538 obtained from the State

Collection of Pathogenic Microorganisms SRCPMB-Obolensk. The cultivation of bacterial biofilms was carried out at a temperature of 37 °C for 24 and 72 hours in Petri dishes on the surface of a dense nutrient medium Muller-Hinton agar (Himedia, Mumbai, India).

Antibacterial drugs. Antibiotics – Gentamicin (30 mg l⁻¹), Chloramphenicol (30 mg l⁻¹), Tigecycline (30 mg l⁻¹); antiseptic drug benzalkonium chloride (1 mg l⁻¹).

Obtaining bacterial biofilms. The surface of an agar plate was inoculated with 0.1 ml of a bacterial culture suspension at a concentration of 10^7 cfu ml^{-1} . The cultures were incubated at 37 °C for 24 and 72 hours to form a biofilm. At the end of cultivation with sterile tweezers, the surface of the bacterial lawn was covered with sterile cellulose applicators with a size of 7 × 7 mm and kept for 2–3 minutes. Then, the applicators with biofilm imprints were transferred with sterile tweezers to the surface of an aluminum plate with a size of 20 × 20 mm for 30 seconds. After that, the cellulose applicator was removed with tweezers, moving upwards without shifting and pressing. Next, the sample was fixed as described below. Despite transfer, several fixations and rinses, we observed *S. aureus* biofilms, which are similar to biofilms that were published before in the literature.⁹⁹

Research of tolerance of biofilms to antibacterial drugs. The biofilm sample, transferred to the surface of an aluminum plate as described above, was covered with 20 µl of an antibacterial drug. For antibiotics (Gentamicin, Chloramphenicol and Tigecycline), exposure was maintained for 30 minutes. For an antiseptic (benzalkonium chloride), the exposure was kept for 1 min and 5 min. Then, the sample was washed with 100 µl of sterile water and dried, and the sample was fixed as described below.

Preparation of biofilm samples for imaging using scanning electron microscopy (SEM). To prepare the obtained samples for analysis using SEM, a biofilm sample was fixed on an aluminum plate by immersing the plate in 20% glutaraldehyde for 1 hour. After the time had elapsed, the plate with the biofilm fixed on it was washed three times with deionized water and dried in air. Fixation with glutaraldehyde is now considered the optimal procedure for preventing structural damage to bacterial cells due to its ability to quickly and irreversibly cross-link proteins, fixing the biofilm to the surface.¹⁰⁰ Before SEM measurements, the samples were placed on the surface of an aluminum sample stage with a diameter of 25 mm and fixed with clamps, and a conductive layer of platinum/palladium alloy 10 nm thick was deposited on them. To select the best way to visualize the biofilm, samples were taken with and without a conductive layer of carbon. Modification of the classical fixation technique by changing the concentration of glutaraldehyde and exposure time and avoiding dehydration with alcohol was required because of the emerging sample cracking and complete delamination of the sample.

Electron microscopy measurements

The microstructure of the samples was studied by field emission scanning electron microscopy (FE-SEM) on Hitachi SU8000 and Hitachi Regulus 8230 electron microscopes. The automatic

acquisition mode of data was implemented using the “Zigzag function”. A 10×10 image square pattern (100 images for each sample; the average recording time for one sample was approximately 1.5 h) with negative overlap was used to study each sample. The images were recorded at a magnification of 10 000 times in the secondary electron mode at an accelerating voltage of 10–30 kV and a working distance of 8–10 mm. SEM image stitching was performed using ImageJ software with a grid/collection stitching plugin based on the Fourier shift theorem.¹⁰¹

Neural network implementation and training

Data labeling was performed using Labelbox infrastructure.¹⁰² All networks were implemented with PyTorch¹⁰³ and trained using the PyTorch Lightning python package.¹⁰⁴ The training was performed on a single NVIDIA 1080 Ti graphics card. Each model was trained on 1000 epochs and validated on 8 SEM images.

Segmentation model PyTorch implementations were used.¹⁰⁵ Augmentation was performed using the Albumentations python package.¹⁰⁶ Basic augmentations include flips, rotations, shifts, scaling, sharpening, blurring, grid distortion, and cropping.

Hyperparameters include model architecture, encoder, optimizer, batch size, learning rate, crop height in

augmentations, and special parameters in the loss function (α). In our model, the loss function is in the form:

$$L = \alpha \cdot \text{BCE loss} + (1 - \alpha) \cdot \text{Dice loss}$$

The hyperparameter optimization consisted of sequential model training with different possible values of the selected hyperparameter and fixed values of other hyperparameters. The optimization scheme and optimal hyperparameters are shown in Fig. 7.

The training dataset size/validation IoU score relationship (Fig. 7c) was constructed to study the dependence of model quality on the number of training SEM images. As a result, an intersection over union score of 78.9% on validation was reached using only 4 training images. With further increases in the training size, the metric slowly reaches a value of approximately 85%. Such a good performance of the neural networks is most likely achieved due to the variety and large image size (2560 \times 1760 px).

In addition, the use of elastic transformations (Fig. 7c) of the training images resulted in a 2.7% increase in the IoU score (0 + 4 vs. 4 + 4 elastic transformed images + original images). The computational cost of elastic transformation critically affects

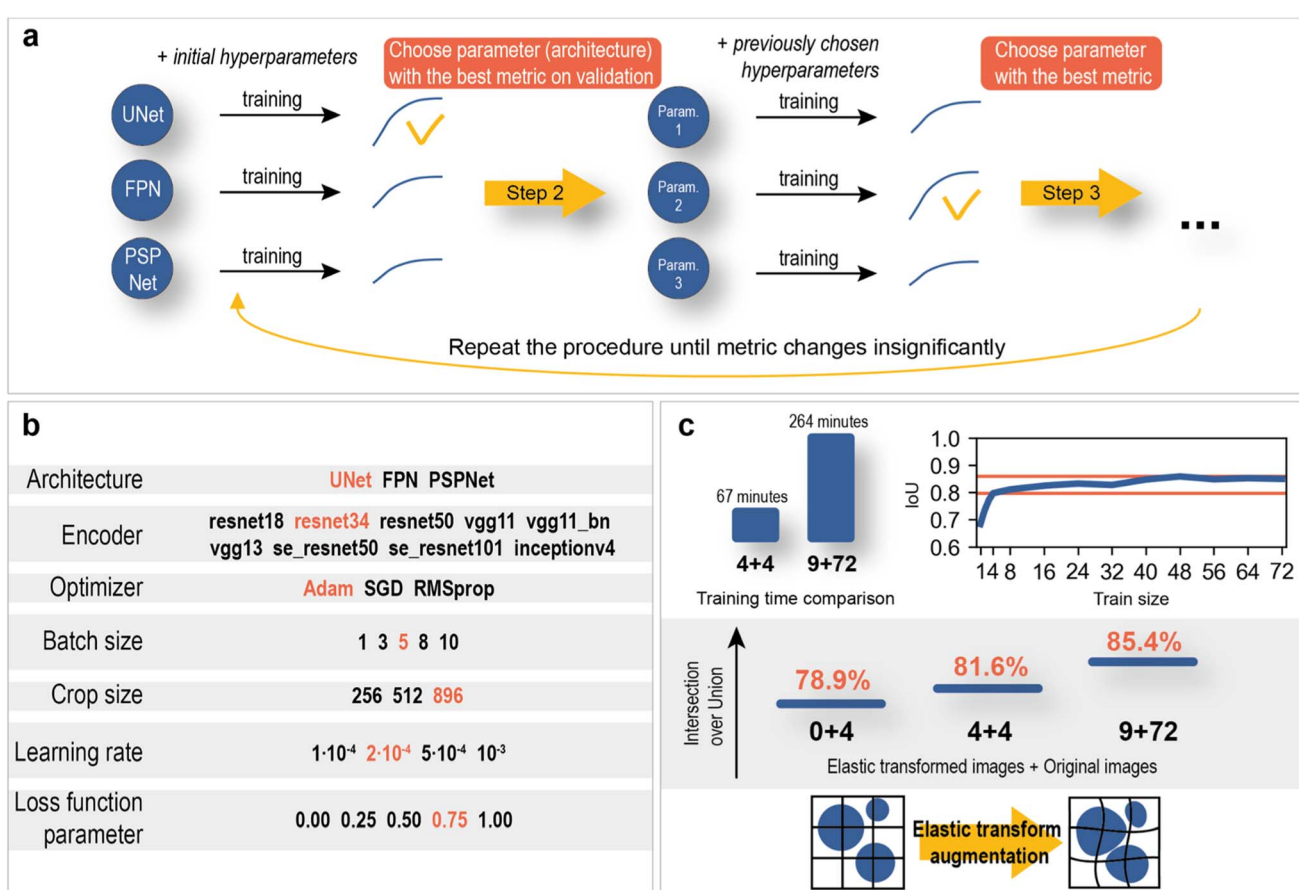


Fig. 7 Neural network optimization. (a) hyperparameter optimization algorithm. (b) Possible hyperparameters (red – optimal parameters). (c) Train size and elastic transform augmentations effect.



the learning time, which is why a set of pregenerated augmented images was used during training instead of real-time elastic transformations.

The final model consisted of 72 training images with 9 elastically transformed images. The training time averaged 4 hours and 24 minutes, which is almost 4 times slower than the training time on 4 original images with 4 elastic transformed images. The difference in IoU score was 3.8% (Fig. 7c). The results are statistically proven with the Mann–Whitney U test on two samples of models with a size of 4 (see section “Statistical significance of using larger amount of training data” in the ESI†). It can be concluded that segmentation models in this work do not demand significant amount of data, which greatly facilitates the data labeling process and reduces training time.

Computer vision methods

Image processing for detection was carried out with the OpenCV-Python¹⁰⁷ package and included the following:

1. Image reading.
2. Gaussian blurring for the following edge detection with a 3 × 3 kernel.
3. Canny edge detection with thresholds of 100 and 200.
4. One iteration of dilation with kernel 5 × 5.
5. Mask inversion.
6. Distance transform (distance type = L2)
7. Connected component computation (connectivity = 4, type = cv2.CV_32S)
8. Contour finding (cv2.RETR_LIST, cv2.CHAIN_APPROX_SIMPLE)
9. Ellipse fitting for bacterial size estimation.

Bacterial size data filtering was carried out with quantiles of 0.01 and 0.99.

Local maxima search (minimum distance between local maxima = 30, *maximum_filter* and *minimum_filter* with neighborhood_size = 25 and threshold difference of 5) and DBSCAN (eps = 0.1 after standard scaling procedure) were carried out with the SciPy Python package.¹⁰⁸ Watershed segmentation was performed using the scikit-image Python package.¹⁰⁹

Kinetic calculations

The neural network used for the computational study of biofilm development kinetics was trained with the same hyperparameters as for the main model but with additional images of an empty support and contrast and brightness augmentations to decrease model data uncertainty.

The k_1 rate constant was estimated by linear regression between the logarithm of cell-free zone ratios and time with sample weights (1/8, 1/8, 1/8, 1/8, 2/8, 1/8, 1/16, 1/16) as measures of prior knowledge certainties for each period of time.

The k_2 rate constant was calculated using minimization of the function:

$$\sum_i^8 [\text{cell area ratio after } t(i) \text{ hours} - f(k_2, t = t(i))]^2,$$

$$\text{where } f(k_2, t) = \frac{k_1}{k_2 - k_1} \cdot (e^{-k_1 \cdot t} - e^{-k_2 \cdot t})$$

The half-life of the cell-free zone area was calculated with the following formula:

$$T_{1/2} = \frac{\ln 2}{k_1} = 32 \text{ hours}$$

Time when the maximal cell area is reached:

$$t_{\max} = \frac{\ln \frac{k_2}{k_1}}{k_2 - k_1} = 62 \text{ hours}$$

Data availability

The code and a link for the files used for the analyses presented in this paper are available at <https://github.com/Ananikov-Lab/cv4biofilms>.

Author contributions

Konstantin S. Kozlov, Daniil A. Boiko – algorithm development, programming, data processing, data analysis; Elena V. Detusheva, Konstantin V. Detushev, Anatoly N. Vereshchagin – microbiological study; Evgeniy O. Pentsak – electron microscopy study; Valentine P. Ananikov – conceptualization, algorithm development, data analysis, supervision. All authors – manuscript preparation and revisions.

Conflicts of interest

The authors declare no competing financial or non-financial interests.

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