Extracting multiple surfaces from 3D microscopy images in complex biological tissues with the Zellige software tool.

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10 Abstract.

11 Efficient tools allowing the extraction of 2D surfaces from 3D-microscopy data are essential for studies 12 aiming to decipher the complex cellular choreography through which epithelium morphogenesis takes 13 place during development. Most existing methods allow for the extraction of a single and smooth manifold of sufficiently high signal intensity and contrast, and usually fail when the surface of interest has a rough 14 topography or when its localization is hampered by other surrounding structures of higher contrast. 15 16 Multiple surface segmentation entails laborious manual annotations of the various surfaces separately. As 17 automating this task is critical in studies involving tissue-tissue or tissue-matrix interaction, we developed the Zellige software, which allows the extraction of a non-prescribed number of surfaces of varying 18 19 inclination, contrast, and texture from a 3D image. The tool requires the adjustment of a small set of control parameters, for which we provide an intuitive interface implemented as a Fiji plugin. As a proof of principle 20 21 of the versatility of Zellige, we demonstrate its performance and robustness on synthetic images and on 22 four different types of biological samples, covering a wide range of biological contexts.

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24 Introduction.

The interplay between gene regulatory networks and physical forces in driving collective cell behaviors is key to tissue morphogenesis during development and to tissue homeostasis throughout life. Recent quantitative studies of epithelial morphogenesis have begun to unravel the basic cellular and physical principles of tissue development, by providing the tools to integrate multiple scales of tissue dynamics [1– 4]. These tools are instrumental to quantify how cell shape changes, cell divisions, cell rearrangements and cell extrusions contribute to tissue remodeling, and to establish data-driven computational models of tissue morphogenesis.

32 Quantitative analysis of an epithelium starts with the extraction of its apical surface from 3D-33 microscopy images (z-stacks of xy-optical sections) encompassing the volume immediately surrounding the 34 epithelium. However, this is a difficult task because this surface is usually not flat (it is best modelled as a 35 curved surface, or 2D submanifold embedded in 3D space), and is often surrounded by other biological 36 structures such as cell layers, acellular membranes, extracellular matrix, and vesicles that hamper its 37 visualization and reconstruction. To make the surface extraction tractable, these studies rely on specific 38 preparations of the specimen, allowing to expose the entire epithelial surface labeled with junctional 39 fluorescent markers to reveal the network formed by epithelial cell-cell contacts. Once the epithelial surface 40 has been extracted, automated cell segmentation and cell contour tracking tools can be used to follow the 41 dynamics of every cell within the epithelium.

Another challenging experimental limitation of these studies is that some of the structures surrounding the epithelium exert external physical constraints that are known to critically affect epithelial morphogenesis by directing cellular dynamics and signaling pathways [1, 5–7]. To understand the physical forces controlling tissue morphogenesis, it is thus essential to also characterize how the dynamics of these extra-epithelial surfaces relate to that of the epithelium (see [8] for review). This calls for the development of dedicated tools allowing the automated extraction of information from several surfaces of interest in a given sample, since the sheer volume of the data precludes any attempt at a manual analysis.

49 Several surface extraction tools have been developed, some of which are available as open access 50 software, such as PreMosa [9], FastSME [10], and LocalZProjector [11]. These tools focus on the extraction of a single, near-horizontal epithelial layer, which is assumed to (i) be sufficiently smooth, (ii) show enough 51 52 contrast against surrounding background signals, and (iii) cover the entire image field-of-view. Specifically, 53 it is assumed that the fluorescent marker used to label the epithelial cell network should provide the highest 54 contrast in the image and allow to select it out from autofluorescent extracellular structures such as the 55 cuticle in flies, or other acellular membranes in mammalian epithelia. The surface is then localized using 56 heuristic algorithms based on the detection of the pixels of maximum contrast and/or brightness. However, 57 applying these tools on more complex biological images with several epithelia of weaker contrast often 58 leads to incorrect localization of the surface of interest, and its blending with the nearby unwanted 59 biological structures.

60 MinCostZ on the other hand, is the only available open-source tool that allows the extraction of up 61 to two surfaces from a 3D stack, and imposes explicit continuity constraints on the reconstructed surfaces. 62 MinCostZ surface extraction relies on a previously developed formulation of the problem as a graph-cut 63 optimization [12]. It is implemented as an ImageJ plugin [13], taking as control parameters, the number of 64 surfaces to extract, the maximum slope and the range of distances allowed between the surfaces, as well 65 as some user-defined cost function that should reflect the characteristics of the surfaces in term of signal intensity, contrast and texture. Despite its interest, this approach remains computationally costly and 66 67 difficult to apply in practice due to the non-trivial choice of the cost function and the need to know 68 beforehand the relative positions of the surfaces to be extracted.

Alternatively, one can segment the surfaces of interest by using supervised machine learning tools 69 70 such as the software solutions Weka [14] or Ilastik [15], as proposed in the ImSAnE surface reconstruction 71 framework [16]. A deep learning approach, using a network of the U-net type to segment the pixels 72 belonging to a single surface of interest, has also recently been reported [17]. While promising as they can 73 provide state of the art segmentations of epithelial surfaces in difficult imaging conditions, machine learning 74 approaches require the prior manual annotation of a sufficiently large set of surfaces to generate suitable 75 training sets. This process can be very time consuming, often necessitating several rounds of trials and 76 errors to obtain satisfactory results, without guarantees to be generic, *i.e.*, to generalize to a wide range of 77 datasets. So far, no solution to the multiple surface extraction problem has been proposed, which is 78 satisfactory both in terms of genericity and ease of use.

79 However, such a tool is highly desirable for modern biology studies. Indeed, tissue organization in 80 the context of developmental biology emerges from the interaction of several neighboring structures [8] 81 through the interplay of molecular signals [18, 19], as well as electrical [20, 21], hydraulic [22, 23] and 82 mechanical contact interactions [24, 25]. The importance of such interaction is exemplified in embryonic 83 explanted tissue cultures that develop abnormally when separated from their neighboring structures [26]. 84 Similarly, in the context of tissue engineering, stem-cell derived aggregates harbor various types of tissues 85 surrounding the genuine organoid, and these tissues presumably influence organoid shape, fate and 86 differentiation (see [27] for review). The ability to simultaneously study the dynamics of neighboring 87 structures together with the structure of interest is therefore essential for an integrated understanding of 88 tissue development, and for any attempt to harness tissue self-organization in vitro.

89 Here, we introduce Zellige, a tool based on a novel constructive approach that allows the automatic 90 extraction of a non-prescribed number of surfaces from a 3D image. To do this, the user is only required to adjust a small set of intuitive control parameters, a task largely facilitated by a user-friendly interface 91 92 implemented as a plugin for the open-source Fiji platform [28]. We tested the performance and robustness 93 of Zellige for multiple surface extraction by applying it to synthetic images and 3D microscopy images from 94 four different types of biological samples, containing multiple surfaces of interest of widely varying texture 95 and contrast. These experiments demonstrate the ability of the approach to extract several (up to 4) 96 surfaces of potentially very low contrast, selectively from other highly contrasted and complex structures, 97 with a single set of reconstruction parameters. A sensitivity analysis also reveals a high robustness of Zellige 98 against small variations of these parameters. This will make it a tool of choice in terms of versatility and 99 ease of use for the investigation of biological surfaces.

100 Results & discussion.

¹⁰¹ Proof of concept of multiple surface extraction on a synthetic image.

The implementation of Zellige is summarized in Figure 1 and in the Methods section (Figure S1). Figure 2 102 shows the results produced by Zellige on a phantom 3D image [29] containing three distinct synthetic 103 104 surfaces generated as described in Supplementary note 2. The three surfaces are extracted with little errors. 105 We assessed the quality of the reconstruction by comparing each of the height-maps produced by Zellige to the corresponding ground truth (GT) height-map, which is exactly known in this case (Figure 2A-C, and 106 107 Supplemental note 3). For the three surfaces, the reconstruction has subpixel accuracy over >99% of the 108 GT pixels (Figure 2D-E), with a root mean square error (RMSE) of ≤ 0.6 in pixel units, showing that the 109 surface localization is highly accurate. In addition, the coverage, which measures the proportion of the 110 reconstructed surface relative to the GT, is near 100% for the three surfaces. To achieve these results, the 111 control parameters of the two steps of the surface extraction were adjusted manually to some adequate reference values (see Supplementary Table S1) using the Zellige Fiji interface. Only the parameters 112 controlling the pixel classification step (amplitude and Otsu threshold parameters T_A and T_{otsu} , minimal 113 island size S_{min} , and smoothing parameters σ_{xy} and σ_z) did actually require a modest adjustment. The 114 parameters of the surface assembly step (parameters T_{OSE1} , R_1 , C_1 and T_{OSE2} , R_2 , C_2 of the 1st and 2nd 115 construction rounds, respectively) were set to their default reference values (see Figure S2 and 116 117 Supplemental note 4) and did not need to be adjusted.

118 Thus, using a single set of control parameters, Zellige can extract multiple surfaces of various 119 shapes and textures with virtually no error, without requiring the user to provide information about their 120 number or relative position, nor about their shape or texture characteristics.

121 Performance of Zellige on biological samples.

122 Example 1: Extracting multiples surfaces from an image of a pupal fly specimen.

Over the past few decades, the Drosophila model has been invaluable to decipher the molecular and cellular 123 124 mechanisms underlying organ embryogenesis [30, 31]. Epithelium morphogenesis studies not only revealed 125 the importance of mechanical stresses (including stress boundary conditions) and planar polarity signaling 126 on cell dynamics to generate tissues of reproducible sizes and shapes, it also highlighted the importance of 127 extracellular matrix attachments in constraining the tissue stresses that guide patterning [1, 32]. At the 128 pupal stage, the fly undergoes dramatic remodeling of its larval organs into adult organs. Large scale tissue flows initiate at a timing that coincides with molting, when the epithelium contracts away from the 129 130 overlaying cuticular sac, a protective acellular membrane that imposes mechanical boundary conditions to 131 the tissue.

Figure 3 shows the results of applying Zellige on a 3D image of a Drosophila pupa acquired with a 132 spinning disk confocal microscope [33]. The sample expresses Ecadherin-GFP, a fluorescent marker of cell-133 cell junctions, and encompasses a portion of the pupa's abdomen and a small portion of its wing. Four 134 surfaces of interest can be identified, with varying signal intensities, noise levels and features (Figures 3A-135 136 B). The abdomen is formed of an epithelium (surface S2) overlaid by a cuticle (surface S1). Lying just beneath 137 these two surfaces, one can observe globular structures showing in some places a higher intensity than the 138 signal coming from the surfaces. The wing also consists of an epithelium of low intensity signal (surface S4), 139 and an overlying cuticle (surface S3). These two surfaces are relatively flat, except for surface S3 which is 140 very steep near one of its edges.

141 Figure 3C shows a 3D graphical representation of the height-maps reconstructed by Zellige (green) 142 and those reconstructed by an expert biologist (blue), taken as ground truth (GT). While these height-maps clearly show greater roughness than those of the synthetic surfaces presented earlier, they could again be 143 144 obtained with a single set of control parameters that were adjusted manually with the Zellige interface (see 145 Supplementary Table S1). We observe an excellent match between the four reconstructed and 146 corresponding GT height-maps, despite the rather complex topography of surfaces S1 and S2 (with slopes 147 reaching up to 45°), and the near-vertical inclination of surface S3 at its boundary. Yet, small deviations may 148 be seen in the regions of highest slope of the surfaces. Some of these deviations are likely attributable to 149 uncertainties in the definition of the GT height-maps, whose accuracy depends on the expert.

150 Figure 3D shows the differences between the reconstructed and GT height-maps, plotted as color-151 coded error maps. These differences are <2 (in pixel units) for most pixels, while some regions of higher error values can be seen locally in surfaces S1 and S2, and at the boundary of surface S3. Note that for 152 153 surfaces S2 and S4, which contain junctional epithelial meshes composed of larger and smaller cells, 154 respectively, the GT height-map encompass not only the mesh but also the interior of the cells, where no 155 junctional signal is detected. The distance calculated inside the cells is thus more subjected to intensity 156 fluctuations, especially for surface S2. Nonetheless, the RMSEs of surfaces S1, S2 and S4 are less than 1, 157 showing that on average the reconstructed height-maps match the corresponding ground truth with 158 subpixel accuracy. The higher RMSE (1.25) of surface S3 is largely due to the region of steep region at the 159 edge of this surface (yellow region on the error map for this surface, Figure 3D). The coverage of the reconstructed height-maps is excellent (≥ 96%) for surfaces S1 and S2, and slightly lower, but still very good 160 161 (≥ 93%) for the smaller surfaces S3 and S4. Figure 3E shows the 2D projections of the 3D image obtained 162 for each of the reconstructed surfaces and for the corresponding ground-truths. The inaccuracies visible on 163 the error maps (see Figure 3D) do not significantly impact these projections, which appear very similar to 164 the projections obtained with the corresponding ground-truths. Thus, while the biological sample contains significant noise and shows a much more variable contrast (especially with the presence of high intensity 165 166 globular structures near surfaces S1 and S2), Zellige makes it possible to segment these surfaces selectively, 167 with a quality of segmentation comparable to that obtained by manual expert segmentation.

168 This possibility brings several perspectives that are not offered by single surface extraction 169 algorithms. First it opens the possibility to systematically study the tissue axial movements (along z) relative 170 to the cuticle during molting, allowing for example to gain insights into the early tissue contraction of the 171 wing hinge that acts as a mechanical inducer over the wing blade [8]. Second, Zellige makes it possible to 172 automatically extract structures such as the abdomen epithelium, which is usually segmented manually 173 [34], due to the difficulty to separate the large larval cells from the cuticle mesh and from other globular 174 structures (such as fat bodies or macrophages) present underneath the epithelium. All these structures 175 become intertwined when using other extracting tools. In this context, Zellige opens new opportunities to 176 study collective cell behavior during epithelial morphogenesis in vivo, and to integrate in the analysis the 177 surrounding surface-like structures involved in the mechanics of the system.

178 Example 2: Extracting a thin cochlear epithelium surface from a multilayer dataset.

As the first model in which planar cell polarity signalling was shown to be conserved in vertebrates [35], the mammalian auditory organ, the cochlea, is arguably our most valuable model to study epithelial patterning and morphogenesis beyond the fly and zebrafish [36, 37]. Cochlear morphogenesis involves complex and tightly controlled patterning processes during which the cochlear sensory epithelium extends and develops its characteristic coiled snail shape, while adopting a striking cellular mosaic organization, with graded changes of morphogenetic parameters along the cochlea [38, 39]. These morphogenetic processes are well recapitulated in organotypic cultures, on the condition that the mesenchyme that underlies the epithelium

be preserved. The cultures are then amenable to live imaging [37, 40], pharmacological [41] and geneticmanipulations.

188 Figure 4A shows a confocal swept field microscope acquisition of an embryonic mouse cochlea [42]. The sample contains only one surface of interest, the cochlear epithelium, but this surface lays on top 189 190 of a thick tangled mesh of non-epithelial cells originating from the mesenchyme. The whole biological tissue 191 is stained for filamentous actin (F-actin) using phalloidin. The epithelium surface presents a non-uniform 192 signal included in a small z-range ($6 \le z \le 10$), and a mesh of very heterogeneous size. Between sections 193 z = 10 and z = 14 one can observe the basolateral region of the epithelial cells, also stained for F-actin. he 194 particularity of this sample is that the mesenchyme presents an intense and contrasted signal over a wide 195 range of z-values ($14 \le z \le 43$). This makes it challenging to extract the surface of the epithelium, which is 196 characterized by low intensity and low contrast.

Figure 4B shows a 3D representation of the height-map reconstructed by Zellige and the corresponding GT height-map (again reconstructed manually by an expert). On the corresponding error map (**Figure 4C**), most (~83%) pixels of the reconstructed height-map show subpixel accuracy (with distances < 1 to the corresponding pixels of the GT height-map). The errors are greater in regions where the cell size is larger, as well as in the area where the signal intensity is very low. However, they remain smaller than 2 for > 95% of the pixels. This result is consistent with the low value of the RMSE (1.1). The surface is also reconstructed with an excellent coverage (> 99%, **Supplementary Table S1**).

204 As the sample contains a single epithelial surface of interest, we compared the performance of 205 Zellige with three other software that can extract only a single surface (Figure 4D). The projections of PreMosa, FastSME and LocalZProjector completely miss the epithelium. Only regions of high contrast 206 207 corresponding to the mesenchyme are projected. In contrast, Zellige generates a projection very close to 208 the ground truth. This demonstrates the efficiency of Zellige to selectively extract a low contrast surface, 209 despite the presence of several structures of higher contrast. Indeed, Zellige detects every structure as a 210 possible surface seed without any assumption on its contrast, and only extends this seed into a surface if enough spatial continuity is found in the surrounding signal. This feature allows to separate individual 211 212 surfaces from other structures spatially, which should greatly facilitate the analysis of live imaging 213 experiments.

214 Example 3: Extracting a single bronchial epithelial surface rendered abnormally rough by SARS-CoV-2.

215 Recently, we used Zellige to extract the surface of a primary culture of bronchial epithelial cells following 216 infection by the SARS-CoV-2 virus [43]. The infection causes the surface of the epithelium to become 217 abnormally rough due to cell damages as seen from discontinuities within the cell layer. The sample we chose from this study is a 3D confocal image of the epithelium responding to SARS-CoV-2 infection [44] 218 219 (Figure 5A). The surface of interest in this image corresponds to the layer of epithelial cells stained for the 220 tight junction protein Zona Occludens-1 (ZO-1). The surface roughness causes the network of junctions to 221 extend over the height of the z-stack, with a signal of varying intensity (Figure 5A). In addition, the junctional 222 network remains non-planar even at the level of a single cell, hence violating the smoothness condition commonly assumed to hold in the context of epithelial surface extraction. We also observe the presence of 223 224 nearby punctiform structures of high contrast that are mainly located outside of the epithelium surface. 225 This sample therefore provides an example of a surface with a complex landscape, interspersed with a 226 constellation of signals which may interfere with the surface segmentation.

The 3D representation of the reconstructed and corresponding GT height-maps (**Figure 5B**) makes it possible to appreciate the roughness of the surface of interest. The two height-maps overlap quite satisfactorily. As shown on the error map (**Figure 5C**), a large majority (71.1%) of pixels of the reconstructed

230 height-map show errors smaller than 1 pixel (~ 96% of them showing errors smaller than 2 pixels). The error 231 is however larger in regions where the cell size is larger, as well as in areas where the ZO-1 signal intensity 232 is very low, preventing a complete reconstruction of the junctions. Nevertheless, the overall RMSE remains small, with a value of 0.81 (Figure 5). The coverage of the reconstructed surface is also excellent (98% of 233 234 the GT height-map), despite the above-mentioned discontinuities. Figure 5D shows the comparison of the 235 projection generated by Zellige to those produced by PreMosa, FastSME, and LocalZProjector. The 236 projection generated by PreMosa misses many junctions of the epithelial network, but it removes quite 237 well the punctiform signal originating from other optical sections. FastSME performs better than PreMosa 238 in reconstructing the junctions, but they produce a projection where the punctiform signal remains strong. 239 In contrast, Zellige and LocalZProjector manage to both reconstruct the surface well and to filter out the 240 punctiform signal quite effectively. This result demonstrates the efficiency of Zellige to extract a surface 241 with complex topography by excluding intense and contrasted spurious signals away from the epithelium 242 surface.

Example 4: Extracting the apical and basal layers of a dome-shaped epithelium (developing inner ear organoid).

245 Organoids are stem cell-derived and self-organizing 3D tissue structures that can mimic certain organ 246 structures. They have emerged as promising in vitro models for developmental biology research, as well as 247 biomedical translational research applications. Here we take the example of mouse stem cell derived inner ear organoids that form vesicular structures composed of an epithelium harboring sensory cells. These 248 249 organoids are part of a cellular aggregate that also contains other tissues such as the mesenchyme [45] 250 adjacent to the organoids. The epithelial cells of the forming inner ear organoids acquire a basal-apical 251 polarity, with their apical side facing the lumen of the organoid, and their basal side facing outwards. The 252 apical junctional network of the epithelium is difficult to visualize in microscopy images as it is seen from 253 below, through the basal layer. Another difficulty is the spherical geometry of the vesicle system, which 254 makes the epithelial surface of interest difficult to extract in regions of high inclination relative to the focal 255 plane.

Figure 6 shows the result of applying Zellige on a 3D confocal microscopy image of half of a developing inner ear organoid at 14 days of culture, a stage at which markers characteristic of the mouse otic vesicle can be detected [46]. The sample was fixed and stained for F-actin to visualize all cellular structures including the epithelium. Two surfaces of interest can be identified, namely the basal side of the epithelium and the apical junctional network (Figure 6A-B). Both surfaces are mesh-like structures of high inclination, high signal intensity and high contrast. The vesicle lumen also contains cell debris of high intensity and contrast that are not part of any surface of interest.

263 Figure 6C shows a 3D graphical representation of the height-maps reconstructed by Zellige and those reconstructed by an expert biologist, taken as ground truth (GT). Due to their dome-shaped 264 265 topography, the manual segmentation of these surfaces was rather laborious, and is more likely prone to 266 errors in the regions of high inclination. Despite this, we observe an excellent match between the two reconstructed and corresponding GT height-maps (Figure 6D). The distance between the two height maps 267 268 is <2 for the large majority (> 92%) of pixels, while larger error values occur locally in regions of near-vertical 269 inclination of the surfaces. The RMSEs of both apical and basal surfaces are close to 1, showing that on 270 average the reconstructed height-maps match the corresponding GT with about pixel accuracy. The 271 coverage of the reconstructed height-map is nearly 100% for the basal surface, and ~88% for the apical surface. Figure 6E shows the 2D projections of the 3D image obtained for each of the reconstructed 272 273 surfaces, as compared to the projected GT height maps. Thus, for the extraction for these highly inclined surfaces, Zellige produces height maps of a quality comparable to those obtained from manual expertsegmentation.

In this example, Zellige could be combined with a 2D cell tracking framework such as TissueMiner 276 277 [3] to perform cell dynamics analysis. Note that geometric distortions introduced in the projected surfaces 278 by the epithelium inclination could be corrected for, using complementary tools such as DProj [11]. This 279 approach could provide a means to quantitatively address how an inner ear organoid epithelium patterns 280 at the cellular and organoid scales, while quantifying the epithelial thickness changes due to cellular 281 intercalation or cell shape changes in the depth of the epithelium. This would also permit to better 282 characterize the variability of inner ear organoids within in a given aggregate, and it could allow one to 283 explore how the organoid interacts with surrounding tissues and how these interactions influence the 284 differentiation of their constituent sensory cells.

A sensitivity analysis reveals the robustness of Zellige in extracting surfaces from biological images.

To evaluate further the quality and robustness of the segmentation obtained by Zellige, we carried out a sensitivity analysis of the reconstruction on each of the samples tested. This analysis consisted in varying one control parameter at a time (**Figures S2-S6**, **Figure 7**, Supplemental note 4), while keeping the other parameters fixed at a nominal value (Supplementary Table S1). The RMSE and coverage of each of the reconstructed surfaces were evaluated and plotted as a function of the value of the parameter that was varied.

293 Figure S3 shows the results of the sensitivity analysis carried out on the image of Figure 3 (Example 1, pupal fly specimen), when varying the parameters controlling step 1, i.e. the selection of 294 295 putative surface pixels (parameters T_A , T_{otsu} , S_{min} , σ_{xy} and σ_z). As can be seen, the variations of the two 296 classification threshold parameters T_A, T_{otsu} and of the minimum island size S_{min} in their respective intervals 297 does not substantially modify the RMSE and the coverage of the reconstructed surfaces, whose values 298 remain roughly constant for $T_A \le 8$, $T_{otsu} \le 12$ and over the entire S_{min} interval (Figure S3A). Within these 299 intervals, the RMSEs of all the reconstructed surfaces remain \leq 1.5, while the coverage values are > 95% for 300 surfaces S1 and S2, and > 85% for surfaces S3 and S4. The surfaces S1 and S3 show lower signal intensity 301 and lower contrast than surfaces S2 and S3, making them more difficult to extract. Surface S4 has the lowest 302 contrast, and fails to be reconstructed if the classification threshold values are too stringent (namely for T_{otsu} > 12, or T_A > 8). Nevertheless, the intervals of stability of T_A and T_{otsu} (that is, the intervals of values 303 304 over which a high-quality extraction of all surfaces is obtained) remain relatively wide (cf. Figure 7). The 305 smoothing parameters σ_{xy} and σ_z also have some effect on the reconstruction of the surfaces. When σ_{xy} is 306 less than 3, the RMSE is higher for the mesh-like epithelial surfaces S2 and S4 (formed by the junctional 307 network of the epithelium). A minimal smoothing along the axial direction is also important to ensure that the reconstructed surfaces are not too fragmented, preventing their complete reconstruction. Yet, σ_z 308 309 should not be chosen too large either, to avoid merging nearby surfaces along the z axis. In this case, the 310 closely positioned surfaces S1 and S2 are well-separated if setting σ_z close to 1, but they become merged 311 when $\sigma_z > 2$. In general, the surface construction parameters have little effect on RMSE and coverage 312 (Figure S3B). The sensitivity analysis on this challenging specimen shows a good robustness of Zellige to 313 extract the four surfaces with a single set of parameters, each of which can be chosen in a reasonably wide 314 interval considering the other fixed.

The results of the sensitivity analyses performed with the other biological image stacks (examples 2, 3 and 4 described above) are shown in **Figures S4**, **S5** and **S6** (Supplemental note 4). These results are summarized in **Figure 7**, which show the stability intervals over which the extracted height-maps

318 satisfy the criteria RMSE \leq 1.5 and coverage \geq 85%, for which the reconstruction can be considered of high 319 quality. Overall, the stability intervals for the two classification threshold parameters (T_A and T_{otsu}) are 320 narrower for specimens containing a surface of low signal intensity and low contrast, while still covering 321 about 1/4 of their respective width. The graphical user interface of Zellige allows the user to adjust T_A and 322 T_{otsu} interactively, making it intuitive to search for reasonable values. We found that $2 \le \sigma_{xy} \le 3$ and $\sigma_z = 1$ 323 generally give high quality results for all tested specimens (Figure 7A). We therefore expect only little 324 adjustment to be required by the user on the smoothing parameters from their default permissive values 325 (set to $\sigma_{xy} = 1$ and $\sigma_z = 1$). Values of σ_z that are too large may lead to the merging of a surfaces with a nearby 326 structure of high contrast (surface or else), as it happens for the epithelium surface in the cochlea specimen, 327 which merges with the underlying mesenchyme signal when σ_z is greater than 2 (Figure 4 and Figure S4). 328 The effect is even more pronounced for surfaces of high inclination (Figures 2, 6 and Figures S2, S6), or 329 presenting a particularly rough texture (Figures 5 and Figures S5).

Regarding the control parameters of step 2 (the assembly step), for the four examples presented except for example 1, the quality of the reconstructed surfaces is stable and high over the main part of their respective intervals of variation, deteriorating occasionally only when extreme values (\simeq 1) are used for these parameters (see **Figures S2-S6, Figure 7B** and Supplemental note 4). Example 1 poses particularly stringent constraints on the control parameters of the reconstruction due to the requirement of reconstructing the 4 surfaces of different contrast and texture present in this sample using the same set of parameters.

Finally, the computation times for running Zellige on a given dataset ranged between a few seconds and a minute on a standard PC computer (see **Figure S7** and Supplementary note 5), except in a few exceptional cases corresponding to extreme values of the control parameters. As a safeguard, a stopping criterion could be implemented so as to exit the run (declaring the current parameter values invalid) if the surface assembly computation exceeds a user-prescribed duration.

342 Overall, the sensitivity analysis indicates that the surface extraction performed by Zellige is robust 343 to variations of the control parameters of step 1 (surface pixel selection step). In general, the reconstruction is more sensitive to the amplitude threshold parameter (T_A), and the Otsu threshold parameter (T_{otsu}) 344 should be kept sufficiently low for samples containing surfaces of intensity close to the background. 345 346 However, in some cases such as in our example 3, the opposite is true. Thus, the two threshold parameters 347 play somewhat complementary roles, and the possibility to adjust them independently is useful in practice 348 to be able to cover as many cases as possible. A smoothing along xy appears necessary to correctly 349 reconstruct the surfaces supported by a junctional mesh. Not surprisingly, best results are obtained when 350 the radius of the gaussian filter used for this (parameter σ_{xy}) is adapted to the mesh (or cell) size. Likewise, a smoothing along z is beneficial, but the extent of this smoothing (parameter σ_z) should not be too large 351 352 to avoid causing the fusion of nearby surfaces. With a few exceptions, the values of the RMSE and coverage 353 show little sensitivity to the values of the parameters controlling step 2 (surface assembly step), at least once putative surface pixels have been properly selected. In the presence of several surfaces of potentially 354 355 very different sizes, the parameter controlling the fraction of OSE sizes allowed for OSE seeds (parameter 356 T_{OSE1}) should be relatively large (≥ 0.5 or greater, *i.e.* allowing more than 50% of the largest OSE sizes for 357 seeds) to allow the extraction of a surface of small size (for example, to extract the surface n° 4 of 358 example 1, which covers less than 20% of the xy-field of view, T_{OSE1} must be larger than 0.6). Extreme values 359 (close to 1) for the connectivity rates (C1 and C2) are too stringent and lead to a drop in the coverage of the 360 reconstructed surfaces. To sum up, we see that the most critical parameters for a satisfactory extraction of 361 the different surfaces are those controlling step 1. In most cases the parameters controlling step 2 do not 362 need to be adjusted and can be fixed to their default reference values.

363

364 Conclusion.

We have developed Zellige, a new tool to extract multiple surfaces from 3D fluorescence microscopy 365 366 images. Zellige automatically finds surfaces by first identifying putative pixels that are likely to belong to a 367 biological surface, and second by assembling a surface through connection of adjacent pixels satisfying 368 natural proximity constraints. By using Zellige on synthetic epithelium images we have shown that it accurately reconstructs a surface with excellent performances in terms of both the distance to the ground-369 370 truth height-map and the surface coverage (Figure 2). Zellige can deal with complex images containing 371 multiple surfaces, with computation times not exceeding a few tens of seconds on a standard computer. 372 Importantly, the user is not required to specify the number of surfaces to be extracted. In the Drosophila 373 specimen (Figure 3), the software readily extracts the 4 surfaces of interest that could be identified. Since 374 Zellige detects putative surface pixels in the first step by combining local and global thresholds, it can deal 375 with images where the multiple surfaces display different features, such as in the mouse cochlear embryo 376 (Figure 3). With this difficult dataset, we could also confirm Zellige's robustness against very low signal-to-377 noise levels. The constructive approach of surface region growing used by Zellige in its second step enables 378 it to circumvent the surface smoothness requirement, that is classically assumed by other surface extraction 379 tools. For instance, it could reconstruct the highly irregular surface of a bronchial tissue infected by SARS-CoV-2 (Figure 5). 380

The robustness and flexibility of Zellige come at a price, namely, the requirement to specify 12 parameters when running the surface extraction. However, the sensitivity analysis we performed shows that adjusting only 4 of these parameters is enough in practice to handle a wide range of image types. These parameters correspond to intuitive notions (*e.g.*, thresholding and smoothing), which makes Zellige particularly easy to use. The Fiji interface that we implemented to perform this adjustment should make Zellige even more user-friendly and effective for biological applications.

To our knowledge, Zellige is the only open-source tool that can extract an unspecified number of epithelial surfaces from a 3D volume, possibly larger than two. This unique feature is especially useful in complex images that could be processed only by specialized tools before. For instance, Zellige can extract surfaces with projections on the *xy* plane that completely overlap, such as the basal and apical epithelia in the organoid image of **Figure 6**. Previously, such an image could be processed only by tools that relied on segmenting a mesh around the object surface, such as MorphoGraphX or ImSAnE [16, 47].

393 The flexibility and robustness of Zellige should allow to considerably relax the constraints that were 394 previously imposed on the sample preparation and the image acquisition steps by the subsequent analysis. 395 Indeed, Zellige can accommodate any number of surfaces in the acquired volume, overlapping or not, and 396 of different contrast features. Zellige also showed excellent robustness against image noise. This should 397 make it particularly useful in imaging contexts that are not easily amenable to automated analysis, such as 398 intravital imaging. Finally, it is worth noting that Zellige is a generalist and modular method. With some 399 adaptation of the surface pixel selection step, it could be used with imaging modalities beyond the scope 400 of this article, for instance, in extracting the irregular and noisy surfaces of biological objects imaged with 401 3D electron microscopy images.

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403

404 Methods.

405 Implementation.

406 Zellige was devised with the goal of achieving accurate segmentation of multiple biological surfaces from 407 3D confocal images. Unlike other existing surface extraction tools, it makes no assumption on the number 408 of surfaces to be extracted and does not require the surfaces of interest to be the structures of highest 409 contrast in the image. Zellige is written in Java, relying on the ImgLib2 library [48] and is distributed as a Fiji 410 plugin, with a graphical user interface (GUI) designed to allow users to quickly find a good set of extraction 411 parameters for a given image.

- 412 Zellige extracts each surface present in the image in the form of a height-map (or *z*-map), that is, a 413 mapping
- 414 $z: (x,y) \rightarrow z(x,y),$

415 which associates to each point (x,y) over which the surface projects, the z-coordinate of the unique pixel 416 (x,y,z) belonging to the surface. Each extracted height-map is then used to produce a 2D projection of the 417 3D stack restricted to a small sub-volume (of user-selected width) centered around the corresponding 418 surface. To achieve a robust extraction, Zellige proceeds in two algorithmic steps, which are only outlined 419 below (see Supplementary note 1 for implementation details).

420 In the first step, or *surface pixel selection* step, a segmentation is applied to the 3D image to select 421 pixels that likely belong to a surface of interest (Figure 1 and Figure S1A). These putative surface pixels are 422 detected as local maxima of image intensity along the z-axis, after using two independent binary classifiers, 423 one based on pixel contrast and the other one on pixel intensity. Five adjustable parameters control the 424 selection step: two threshold parameters (T_A and T_{otsu}) control the strength of the binary classifiers applied 425 on contrast and intensity, respectively, and three parameters (S_{min} , σ_{xy} and σ_z) control clean-up operations 426 applied at the end of the classification (removal of small isolated spots, and local averaging along the xy 427 plane and the z axis, respectively).

428 In the second step, or surface assembly step, an iterative algorithm is used to extract the height-429 maps of each of the surfaces present in the image (Figure 1 and Figure S1A). The assembly starts by 430 grouping neighboring putative surface pixels together within each orthogonal (xz or yz) section of the 3D 431 image, in order to form a set of building blocks referred to as orthogonal surface elements (OSEs). These building blocks are then used to assemble the surfaces, in a process analogous to jigsaw puzzles, where 432 433 OSEs adjacent to the surface boundary are added if they match this boundary, and rejected otherwise, until 434 no matching OSE can be found (Figure S1). In order to increase the robustness of the assembly step, Zellige applies it in two rounds, proceeding along different axes during the first and second rounds. Each round is 435 controlled by 3 adjustable parameters: a threshold parameter ($0 \le T_{OSE} \le 1$) sets a minimum size for the 436 building blocks that can be used as seeds to initiate the assembly of a surface; and two other parameters 437 438 $(0 \le R \le 50 \text{ and } 0 \le C \le 1)$ set the matching constraints used to accept or reject the addition of OSEs to a 439 surface. The assembly step is thus controlled overall by 6 parameters, *i.e.* two groups of 3 parameters (T_{OSE1} , R_1 , C_1) and (T_{OSE2}, R_2, C_2) controlling the first and second assembly rounds, respectively. 440

441 Finally, the height-maps of each of the reconstructed surfaces are used to obtain a corresponding
442 2D projection (Figure 1). In practice a maximum projection restricted to a subvolume of width δz (δz being
443 a user-defined parameter) centered around the surface of interest is performed.

444 Availability of data and code.

- The project homepage below contains the source code, installation instructions, and documentations.
- 446 Zellige
- 447 ٠ Project name: Zellige. 448 Project homepage: https://gitlab.pasteur.fr/ida-public/zellige-core 449 URL for the Fiji plugin (Update > Manage update sites): https://sites.imagej.net/Zellige/ . 450 • Gitlab Branch: master Operating systems: Platform independent. 451 • 452 • Programming language: Java. Compiled in Java8 453 • Other requirements : Runs from Fiji [28]. 454 • License: BSD 2 455 • Any restrictions to use by non-academics: None. 456 • Scripts to create Phantoms 457 Project name: Phantoms. 458 • Project homepage: https://doi.org/10.5281/zenodo.6414596 459 .
- 460 Operating systems: Platform independent.
- Programming language: MATLAB.
- 462 License: BSD 2
- Any restrictions to use by non-academics: None.
- 464 Data sets.

Image 3D stacks, ground-truth height map and height maps produced by Zellige are available on Zenodo
 under the CC-BY license: <u>https://zenodo.org/communities/zellige/</u> [29, 33, 42, 44, 46].

467 Human bronchial epithelium imaging.

The data used in Figure 5 were taken from the recent study [43] to which we refer for the preparation and imaging of human bronchial epithelium cultures. In brief, MucilAirTM were purchased from Epithelix (Saint-Julien-en-Genevois, France) and cultured for at least 4 weeks to reconstruct a differentiated human bronchial epithelium *in vitro* and stained as previously described. Images of the cultures were acquired using an inverted Zeiss LSM 710 confocal microscope controlled by the ZEN pro 2.3 software. Z-stack images of whole-mount samples were acquired with a Zeiss Plan Apochromat 63x oil immersion lens (NA=1.4). The image used here was published in Robinot *et al.* [43] under the CC-BY-4 license.

475 Drosophila imaging.

476 Flies were raised at 25°C under standard conditions. Pupae were collected for imaging as described

- 477 previously [49]. Ecad::GFP flies [50] were used for live imaging as previously described [1]. In brief, images
- 478 were acquired with a spinning disk microscope from Gataca Systems driven by the MetaMorph software.
- 479 The system is equipped with an inverted Nikon TI2E stand, a motorized XYZ stage, and a Nikon Plan Apo 60x
- 480 oil immersion (NA=1.4) lens and with a Prime95B camera.

481 Cochlea imaging.

The inner ears from wild-type (C57BL/6) mice were rapidly dissected from temporal bones at embryonic stages E14.5 in HEPES-buffered (10 mM, pH 7.4) Hanks' balanced salts solution and fixed in 4% paraformaldehyde, 1 hour at room temperature. Specimen were permeabilized and stained for phalloidin-Atto 565 (Sigma) as previously described [39]. Fluorescence images were obtained with a swept-field confocal microscope (Opterra II) from Brucker. This system is equipped with a Nikon Plan Fluor 60x oil immersion lens (NA=1.4).

Biologists in this study hold a designer certificate of animal experimentation (level 1), allowing them to perform experimental work on animals in strict accordance with the European directive 2010/60/EU, and French regulations. The Ethics Committee of the Institut Pasteur (Comité d'Ethique en Experimentation Animale - CETEA) has approved this study with the project identifier dha170006. This approval is based on careful compliance to the 3Rs principle in the care and use of animals (Annex IV - 2010/60/EU).

493 Inner ear organoid imaging.

ESCs derived from blastocyst-stage embryos of R1 mice (mESCs) (ATCC, SCRC-1036) were maintained in 494 495 feeder-free culture on 0.1% w/v gelatin (Sigma) coated substrates using LIF-2i medium as established previously [51]. The organoids were generated following the previously published protocol [45, 51]. 496 Aggregates were harvested at day 14 and fixed in 4% v/v PFA (Electron Microscopy Sciences) overnight at 497 498 4°C. After blocking (PBS; 10% v/v normal goat serum; 0.1% v/v Triton X-100), the aggregates were stained 499 for phalloidin Atto 565 (1:1000) (Sigma) overnight at RT on a shaker, and washed three times with PBS 500 containing 0.1% v/v Triton X-100 for 1 h each at RT. Prior to imaging, the aggregates were incubated in a modified version of ScaleS solution containing 4 M Urea (Sigma), 40% w/v D-Sorbitol (Sigma), and 0.1% v/v 501 502 Triton X-100, for 3-5 days to clarify the tissue. Finally, the aggregates were whole-mounted using the ScaleS 503 solution and imaged using a confocal laser scanning microscope (A1R HD25, Nikon) equipped with a Nikon 504 25x silicon oil immersion lens (NA=1.05).

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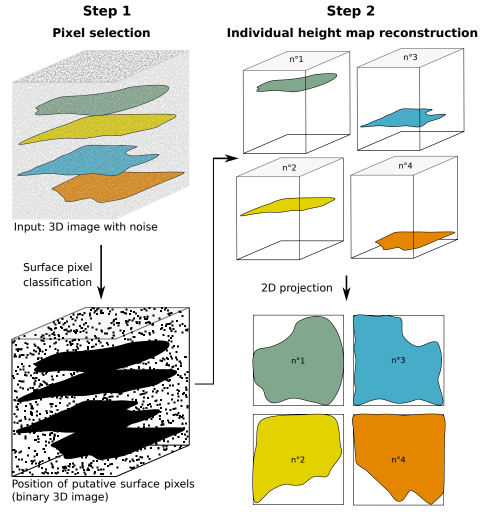


Figure 1. Flowchart of Zellige's algorithmic steps. Surface pixel selection (step 1), surface assembly in the form of a height map (step 2), and subsequent projection localized to the height map, are schematically depicted in the case of a 3D image containing 4 surfaces of interest.

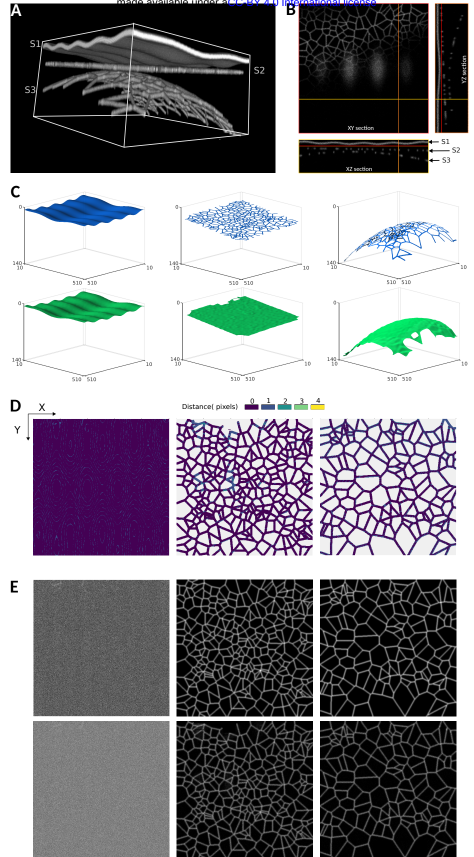


Figure 2. Multiple surface extraction on a synthetic 3D image. (A-B) The image contains 3 phantom surfaces (S1, S2, S3) of different shapes (sinusoidal, flat, and paraboloidal, respectively), and different textures (surface S1 has constant intensity, while surfaces S2 and S3 are supported by Voronoi meshes of different cell-sizes). (C) 3D representations of the height maps extracted by Zellige (in green) and of the ground truth (GT, in blue) height maps of surfaces S1, S2, and S3. (D) Error maps displaying the distance along the z-axis between the reconstructed and GT height maps for surfaces S1, S2, and S3. (E) Projections of the 3D image localized to the different surfaces S1-S3 (maximum intensity projections over a subvolume of a width $\delta z=1$ pixel above or below the corresponding height-maps). Upper and lower panels show the projections based on the GT and the reconstructed height-maps, respectively.

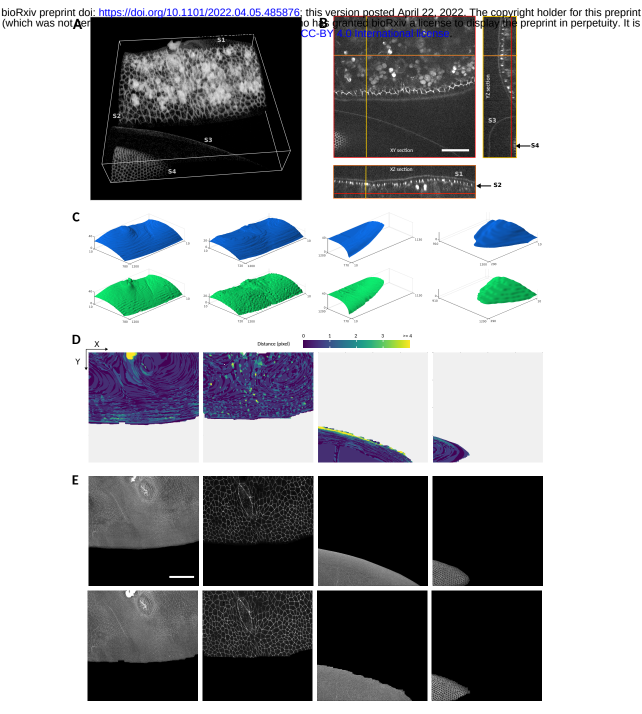
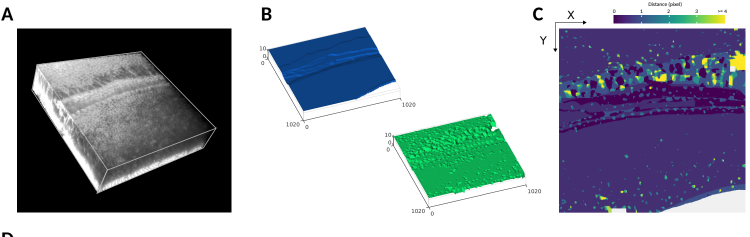


Figure 3. Fly specimen. (A,B) Volume rendering (A) and orthogonal sections (B) of a 3D image of fly embryo taken around 24h after puparium formation, covering a portion of the abdomen (showing histoblast cells and larval cells), and a portion of the developing wing. Scale bar 50 µm. Four surfaces of interest may be identified in the dataset (of dimensions 1200 × 1200 × 51 pixels): surfaces S1 and S2 are relatively close to one another and located within overlapping z-ranges ($8 \le z \le 50$ and $20 \le z \le 50$, respectively). Surfaces S3 and S4 (located in the z-ranges $42 \le z \le 50$ and $9 \le z \le 50$, respectively) are relatively far from each other and can nearly be separated by a plane. (C) 3D representations of the height maps extracted by Zellige (in green) and of the ground truth height maps (GT, in blue) of surfaces S1-S4. The reconstructed height-maps of all surfaces S1-S4 cover >93% of the area of the corresponding GT (cf. Figure S2 and Table S1). To reduce the staircase artifacts (more or less visible depending on the surface) due to the digitization of the GT and reconstructed height-maps, all height-maps were smoothed with a 2D gaussian filter with a standard radius of 5 pixels (cf. Supplemental note 1). (D) Error maps (color-coded distance along the z-axis between the reconstructed and the GT height-maps) plotted for each of the reconstructed surfaces. The large majority of pixels on the reconstructed height-maps (98%, 96%, 91%, and 99% for surfaces S1 to S4, respectively) display errors of <2 pixels. The height-maps of surfaces S1, S2, S4 show subpixel accuracy on average (RMSE < 1), while that of surface S3 is slightly less accurate (RMSE = 1.25). (E) Projections of the 3D image localized to the different surfaces S1-S4 (in this and all subsequent figures, these are maximum intensity projections over a subvolume of width $\delta z = \pm 1$ pixel above or below the corresponding height-maps). Upper and lower panels show the projections based on the GT and the reconstructed height-maps, respectively. Scale bar 50 μ m.



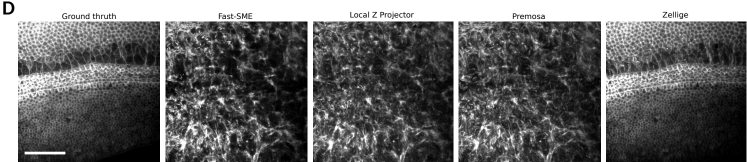


Figure 4. Cochlea specimen. (A) Volume rendering of a 3D confocal swept-field image of the mouse cochlear embryo on embryonic day E14.5. The dataset (of dimensions $1024 \times 1024 \times 45$ pixels) shows a portion of the sensory epithelium (at the topmost sections of the stack) and the underlying non-cellular layer of mesenchyme on which the organ develops. Both structures are stained with phalloidin to reveal F-actin. Scale bar 40 µm. The surface of interest is the epithelium surface, harboring the sensory and supporting cells under differentiation. The mesenchyme layer is not (strictly speaking) assimilable to a surface, but it produces a strong background signal nearby the surface of interest, hampering its extraction. **(B)** 3D representations of the height map extracted by Zellige (in green) and the GT height map (in blue), of the epithelium surface. **(C)** Color-coded error map of the reconstructed height-map, which shows subpixel accuracy (errors <1) over a large majority (83%) of pixels, as well as on average (RMSE ~ 1.1). **(D)** Projections localized to the GT height-map of the epithelium surface (left most panel), and to the height-maps extracted with the four different algorithms: FastSME, LocalZProjector, PreMosa, and Zellige. Only Zellige correctly extracts the surface of the epithelium in this example. Scale bar 40 µm.

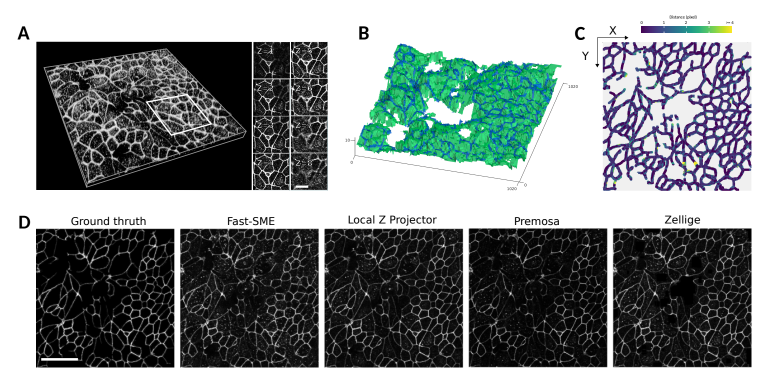


Figure 5. Human bronchial epithelial cells infected by SARS-CoV-2. (A) Volume rendering and individual sections of a confocal 3D image of a primary culture of bronchial epithelial cells 4 days after it was infected by the SARS-CoV-2 virus. The dataset (of dimensions $1024 \times 1024 \times 15$ pixels) covers a portion of the epithelium immunostained for the tight junction protein ZO-1. Notice the roughness of the epithelium surface and the presence of anomalous bulges (arrows) resulting from the SARS-CoV-2 infection. Scale bar 10 µm. (B) 3D representations of the height map extracted by Zellige (in green) and the GT height map (in blue), of the epithelium surface. (C) Color-coded error map of the reconstructed height-map. Despite its roughness, the surface of interest is reconstructed with subpixel accuracy over the majority (71%) of pixels, as well as on average (RMSE ~ 0.8). (D) Projections localized to the GT height-map of the epithelium surface (leftmost panel), and the height-maps extracted with the four different algorithms: FastSME, LocalZProjector, PreMosa, and Zellige. Scale bar 30 µm.

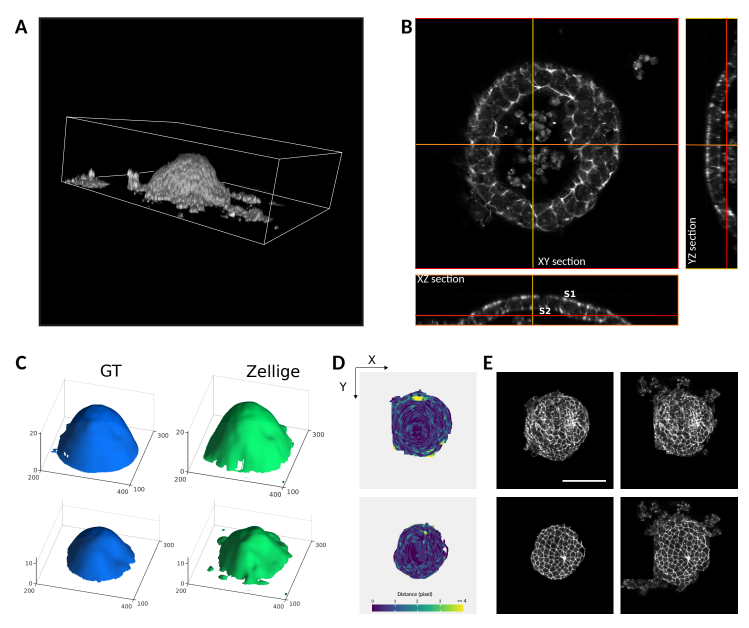


Figure 6. Organoid specimen. (A-B) Volume rendering (A) and orthogonal sections (B) of a confocal 3D image of a (half of) inner ear organoid, which has been fixed and stained with phalloidin to reveal F-actin. The dataset (of dimensions $520 \times 465 \times 35$ pixels) includes two dome-shaped epithelial surfaces of interest, forming the apical (inward) and basal (outward) sides of the organoid. (C) 3D representations of the height map extracted by Zellige (in green) and the GT height map (in blue), of the epithelium surface. (D) Color-coded error maps of the reconstructed height-maps for the apical (left) and basal (right) epithelial surfaces of interest are reconstructed with an error of < 2 pixels over a large majority (96% and 93% for the apical and basal surfaces, respectively) of pixels, as well as on average (RMSE ~ 0.8 and 1.1 for the apical and the basal surfaces, respectively). (E) Projections localized to the GT height-maps of the epithelium surface (panels on the left), and the height-maps extracted by Zellige (panels on the right). Scale bar 100 µm

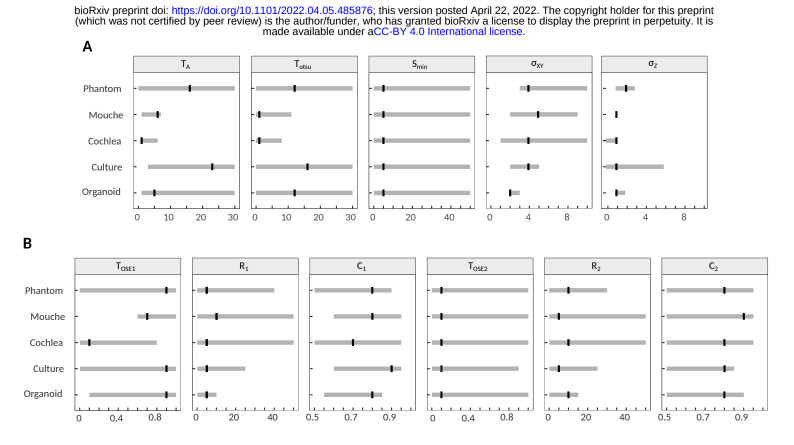


Figure 7. Summary of the sensitivity analysis. The intervals indicated in grey for each parameter and each of the images tested correspond to the parameter values for which the reconstruction satisfies high quality criteria defined by RMSE \leq 1.5 and coverage \geq 85%. Black marks indicate the reference value obtained by manual adjustment for each image (cf. Supplemental note 2). (A) Parameters of the surface selection step. (B) Parameters of the surface assembly step.

FIGURE 1

Flowchart of Zellige's algorithmic steps. Surface pixel selection (step 1), surface assembly in the form of a height map (step 2), and subsequent projection localized to the height map, are schematically depicted in the case of a 3D image containing 4 surfaces of interest.

FIGURE 2

Multiple surface extraction on a synthetic 3D image. (A) The image contains 3 phantom surfaces (S1, S2, S3) of different shapes (sinusoidal, flat, and paraboloidal, respectively), and different textures (surface S1 has constant intensity, while surfaces S2 and S3 are supported by Voronoi meshes of different cell-sizes). (B) 3D representations of the height maps extracted by Zellige (in green) and of the ground truth (GT, in blue) height maps of surfaces S1, S2, and S3. (C) Error maps displaying the distance along the z-axis between the reconstructed and GT height maps for surfaces S1, S2, and S3. (E) Projections of the 3D image localized to the different surfaces S1-S3 (maximum intensity projections over a subvolume of a width $\delta z=1$ pixel above or below the corresponding height-maps). Upper and lower panels show the projections based on the GT and the reconstructed height-maps, respectively.

FIGURE 3

(A,B) Volume rendering (A) and orthogonal sections (B) of a 3D image of fly embryo taken around 24h after puparium formation, covering a portion of the abdomen (showing histoblast cells and larval cells), and a portion of the developing wing. Scale bar 50 μ m. Four surfaces of interest may be identified in the dataset (of dimensions $1200 \times 1200 \times 51$ pixels): surfaces S1 and S2 are relatively close to one another and located within overlapping z-ranges ($8 \le z \le 50$ and $20 \le z \le 50$, respectively). Surfaces S3 and S4 (located in the z-ranges $42 \le z \le 50$ and $9 \le z \le 50$, respectively) are relatively far from each other and can nearly be separated by a plane.

(C) 3D representations of the height maps extracted by Zellige (in green) and of the ground truth height maps (GT, in blue) of surfaces S1-S4. The reconstructed height-maps of all surfaces S1-S4 cover >93% of the area of the corresponding GT (cf. Figure S2 and **Table S1**). To reduce the staircase artifacts (more or less visible depending on the surface) due to the digitization of the GT and reconstructed height-maps, all height-maps were smoothed with a 2D gaussian filter with a standard radius of 5 pixels (cf. Supplemental note 1).

(D) Error maps (color-coded distance along the z-axis between the reconstructed and the GT height-maps) plotted for each of the reconstructed surfaces. The large majority of pixels on the reconstructed height-maps (98%, 96%, 91%, and 99% for surfaces S1 to S4, respectively) display errors of <2 pixels. The height-maps of surfaces S1, S2, S4 show subpixel accuracy on average (RMSE < 1), while that of surface S3 is slightly less accurate (RMSE = 1.25).

(E) Projections of the 3D image localized to the different surfaces S1-S4 (in this and all subsequent figures, these are maximum intensity projections over a subvolume of width $\delta z = \pm 1$ pixel above or below the corresponding height-maps). Upper and lower panels show the projections based on the GT and the reconstructed height-maps, respectively.

FIGURE 4

(A) Volume rendering of a 3D confocal swept-field image of the mouse cochlear embryo on embryonic day E14.5. The dataset (of dimensions $1024 \times 1024 \times 45$ pixels) shows a portion of

the sensory epithelium (at the topmost sections of the stack) and the underlying non-cellular layer of mesenchyme on which the organ develops. Both structures are stained with phalloidin to reveal F-actin. Scale bar 40 μ m. The surface of interest is the epithelium surface, harboring the sensory and supporting cells under differentiation. The mesenchyme layer is not (strictly speaking) assimilable to a surface, but it produces a strong background signal nearby the surface of interest, hampering its extraction.

(B) 3D representations of the height map extracted by Zellige (in green) and the GT height map (in blue), of the epithelium surface.

(C) Color-coded error map of the reconstructed height-map, which shows subpixel accuracy (errors <1) over a large majority (83%) of pixels, as well as on average (RMSE \sim 1.1).

(D) Projections localized to the GT height-map of the epithelium surface (left most panel), and to the height-maps extracted with the four different algorithms: FastSME, LocalZProjector, PreMosa, and Zellige. Only Zellige correctly extracts the surface of the epithelium in this example.

FIGURE 5

(A) Volume rendering and individual sections of a confocal 3D image of a primary culture of bronchial epithelial cells 4 days after it was infected by the SARS-CoV-2 virus. The dataset (of dimensions $1024 \times 1024 \times 15$ pixels) covers a portion of the epithelium immunostained for the tight junction protein ZO-1. Notice the roughness of the epithelium surface and the presence of anomalous bulges (arrows) resulting from the SARS-CoV-2 infection. Scale bar $10 \,\mu$ m.

(B) 3D representations of the height map extracted by Zellige (in green) and the GT height map (in blue), of the epithelium surface.

(C) Color-coded error map of the reconstructed height-map. Despite its roughness, the surface of interest is reconstructed with subpixel accuracy over the majority (71%) of pixels, as well as on average (RMSE ~ 0.81).

(D) Projections localized to the GT height-map of the epithelium surface (leftmost panel), and the height-maps extracted with the four different algorithms: FastSME, LocalZProjector, PreMosa, and Zellige. Scale bar $30 \,\mu$ m

FIGURE 6

(A-B) Volume rendering (A) and orthogonal sections (B) of a confocal 3D image of a (half of) inner ear organoid, which has been fixed and stained with phalloidin to reveal F-actin. The dataset (of dimensions 520 × 465 × 35 pixels) includes two dome-shaped epithelial surfaces of interest, forming the apical (inward) and basal (outward) sides of the organoid.

(C) 3D representations of the height map extracted by Zellige (in green) and the GT height map (in blue), of the epithelium surface.

(D) Color-coded error maps of the reconstructed height-maps for the apical (left) and basal (right) epithelial surfaces of the organoid. The surfaces of interest are reconstructed with an error of < 2 pixels over a large majority (96% and 93% for the apical and basal surfaces, respectively) of pixels, as well as on average (RMSE ~ 0.8 and 1.1 for the apical and the basal surfaces, respectively).

(E) Projections localized to the GT height-maps of the epithelium surface (panels on the left), and the height-maps extracted by Zellige (panels on the right). Scale bar 100 μ m.

FIGURE 7

Summary of the sensitivity analysis. The intervals indicated in grey for each parameter and each of the images tested correspond to the parameter values for which the reconstruction satisfies high quality criteria defined by $RMSE \le 1.5$ and $coverage \ge 85\%$. Black marks indicate the reference value obtained by manual adjustment for each image (cf. Supplemental note 2). (A) Parameters of the surface selection step. (B) Parameters of the surface assembly step.