1 **Overburdened Ferroptotic Stress Impairs Tooth Morphogenesis** H.S. Wang^{1, #}, X.F. Wang^{1,2, #}, L.Y. Huang^{1,2}, C.L. Wang², F.Y. Yu^{1,2, *}, L. Ye^{1,2, *} 2 ¹ State Key Laboratory of Oral Diseases & National Center for Stomatology & 3 National Clinical Research Center for Oral Diseases, West China Hospital of 4 5 Stomatology, Sichuan University, Chengdu, China. ² Department of Endodontics, West China Hospital of Stomatology, Sichuan 6 7 University, Chengdu, China # These authors contributed equally to this work 8 9 10 * Corresponding Author: 11 Prof. Fanyuan Yu 12 Phone: +86 15882456022 13 Fax: +86 028 855003585 14 Email: fanyuan_yu@outlook.com Address: No.14, 3rd Section, Ren Min Nan Road, Wuhou, Chengdu, Sichuan, China 15 16 17 Prof. Ling Ye 18 Phone: +86 028 855003585 19 Fax: +86 028 855003585 20 Email: yeling@scu.edu.cn Address: No.14, 3rd Section, Ren Min Nan Road, Wuhou, Chengdu, Sichuan, China 21 22 23 24

25 Abstract

26	The role of regulated cell death (RCD) in organ development, particularly the impact
27	of non-apoptotic cell death, remains largely uncharted. Ferroptosis, a non-apoptotic
28	cell death pathway known for its iron dependence and lethal lipid peroxidation, is
29	currently being rigorously investigated for its pathological functions. The balance
30	between ferroptotic stress (iron and iron-dependent lipid peroxidation) and ferroptosis
31	supervising pathways (anti-lipid peroxidation systems) serves as the key mechanism
32	regulating the activation of ferroptosis. Comparing to other forms of regulated
33	necrotic cell death (RNCD), ferroptosis is critically related to the metabolism of lipid
34	and iron which are also important in organ development. In our study, we examined
35	the role of ferroptosis in organogenesis using an ex vivo tooth germ culture model,
36	investigating the presence and impact of ferroptotic stress on tooth germ development.
37	Our findings revealed that ferroptotic stress increased during tooth development,
38	while the expression of Gpx4, a crucial anti-lipid peroxidation enzyme, also escalated
39	in dental epithelium/mesenchyme cells. The inhibition of ferroptosis was found to
40	partially rescue erastin-impaired tooth morphogenesis. Our results suggest that while
41	ferroptotic stress is present during tooth organogenesis, its effects are efficaciously
42	controlled by the subsequent upregulation of Gpx4. Notably, an overabundance of
43	ferroptotic stress, as induced by erastin, suppresses tooth morphogenesis.
44	Key Words: ferroptosis, tooth development, organ culture, cell death, non-apoptotic
45	cell death

46

48 Introduction

49 A delicate balance between cell division and regulated cell death (RCD) is crucial for 50 the development of tissues and organs [1]. In organs such as the nervous, immune, 51 and reproductive systems, cells are initially produced in excess and are subsequently 52 removed by RCD. During organ development, structures with transient functions are 53 also eliminated by RCD when they are no longer necessary [2]. The best-known 54 example of this is the formation of digits in higher vertebrates, where the interdigital 55 webs are eliminated primarily through the apoptotic machinery [3]. RCD plays a vital 56 role in animal development and adult life by eliminating abnormal and potentially 57 harmful cells [4]. Apoptosis is the most extensively studied form of RCD which 58 shrinks the nucleus and buds the plasma membrane (without rupturing it), making it 59 essential in achieving cell number regulation, tissue remodeling, and sculpting 60 structures, driving morphogenesis during organ development [5].

61 However, several new types of cell death, including pyroptosis, necroptosis, and 62 ferroptosis, have been discovered [6]. Unlike apoptosis, these newly discovered RCDs 63 are characterized by pore formation and/or rupture of the plasma membrane, thus 64 termed as regulated necrotic cell death (RNCD) [7]. While apoptosis benefits organ 65 morphogenesis in certain circumstances, whether RNCD and its regulatory 66 mechanisms are biologically required in organ development remains poorly 67 understood. Pyroptosis and necroptosis are mainly activated by external pathogens or 68 damage-associated molecular patterns [8], whereas ferroptosis is a distinct form of 69 iron-dependent, lipid peroxidation-driven programmed cell death [9]. Ferroptosis is a 70 particular type of cell death that links lipid metabolism, ROS (reactive oxygen species) 71 biology, iron regulation, and disease [10]. Despite the pathological role of ferroptosis 72 and/or ferroptotic stress in multiple diseases [11,12], recent studies have identified the 73 occurrence of ferroptosis in embryonal erythropoiesis and aging [13] and aged 74 skeletal muscle [14], indicating its unrecognized role in physiological processes. 75 Unlike autonomous activation of apoptosis during organ development, the broken 76 balance between ferroptotic stress and its suppressing mechanisms is now considered 77 the main reason for ferroptosis. Recent studies demonstrated that MYSM1 deficiency

78 causes human hematopoietic stem cell loss by ferroptosis, highlighting the broader 79 developmental and regenerative role of ferroptosis [15]. The investigation into the 80 potential involvement of ferroptosis and/or ferroptotic stress, other than rest forms of 81 non-apoptotic cell death like pyroptosis and necroptosis, in organ morphogenesis is an 82 area of great interest. However, conducting high-throughput analysis through in vivo 83 studies presents challenges as it relies heavily on generating specific transgenic mice, 84 since erastin, the classic small molecule used to induce ferroptosis is feasible in vivo 85 [16]. Thus, there's a crucial need for a more adequate model to explore the 86 ferroptosis/ferroptotic stress involved in development.

87 The process of tooth development, which contains cell number regulation 88 (proliferation, differentiation, and extinction of ameloblast), tissue remodeling 89 (elimination of transient structure called enamel knot), and sculpting structures (cusp 90 formation) [17,18], is an ideal model for the investigation of RCD in organ 91 development. Moreover, ex vivo culture of tooth germ is a well-established study 92 system allows for the controlled manipulation of environmental, genetic and 93 pharmacological factors (like erastin), which can influence developmental events of 94 tooth [19], providing compelling evidence for its usefulness as an efficient research 95 model in studying ferroptosis/ferroptotic stress within developmental process [20].

96 In the present study, erastin was used in an *ex vivo* culture model of tooth germ to 97 investigate the possible role of ferroptosis during tooth morphogenesis. To definitively 98 identify ferroptosis, multiple markers are needed, including lipid peroxidation, iron 99 accumulation, mitochondria injury, and the expression of ferroptosis-related genes [9]. 100 According to our results, Gpx4 (glutathione peroxidase 4, a peroxidase to counter the 101 oxidation of lipids in membranes) [21] is upregulated according to odontogenesis and 102 amelogenesis together with obvious accumulation of iron. Erastin significantly 103 suppressed the morphogenesis of cultured tooth germ and showed a dose-dependent 104 manner. And meanwhile, erastin-treated tooth germ has elevated lipid peroxidation 105 indicated by expression of 4-HNE (4-hydroxy-2-nonenal, end-product of lipid 106 peroxidation), enhanced iron accumulation, shrunken mitochondria, and alternated 107 expression of ferroptosis related genes. However, all these phenomena could be

108 partially rescued by Fer-1, a classical inhibitor of ferroptosis.

In summary, our results suggest that erastin induces morphologic defects of tooth germ is conducted by the activation of ferroptosis. Ferroptotic stress and its regulatory mechanism participate in tooth morphogenesis, which borders our understanding of the physiological function of non-apoptotic regulated cell death in organ development.

114 **Results**

Spatiotemporal characterization of Gpx4 expression and iron accumulation in tooth morphogenesis

117 In the different developmental stages of the mouse first mandibular molar, the 118 expression of Gpx4 was detected by IHC (Fig 1A). These data revealed gradually upregulated expression of Gpx4 in tooth germ from E13.5 to E17.5 (Fig 1A-a to 1A-119 120 c), but started to decrease from PN1 to PN3 (Fig 1A-d to 1A-e). An enlarged view of 121 each group indicated a different expression level of Gpx4 between dental epithelia 122 and mesenchyme (Fig 1A-a' to 1A-e'). After calculating the IOD (Integrated optical 123 density) of Gpx4 in epithelia, mesenchyme, and total area of tooth germ respectively, 124 we found that Gpx4 in dental epithelia is higher than that in mesenchyme although 125 they both increased from E13.5 to E17.5, which could be observed in Fig 1A-a' to 126 1A-d'. Negative controls are listed in Fig S1A.

127 Iron accumulation, one of the core risk factors for ferroptosis, were detected by TPE-o-Py (*ortho*-substituted pyridinyl-functionalized tetraphenylethylene) [22] using 128 129 IF (Fig 1B). The results showed accumulation of iron is upregulated within dental 130 epithelium and mesenchymal cells during differentiation of odontoblast and 131 ameloblast (Fig 1B-a to 1B-c) then decreased at PN1 which is consistent to the expression pattern of Gpx4. However, at PN3, the active secretory period of both 132 133 ameloblast and odontoblast, the accumulation of iron rebound to a high level and 134 distributed within enamel (Fig 1B-e). Collectively, our results demonstrated that at early stage of tooth development, 135 enhancing risk factors of ferroptosis (iron accumulation) was accompanied by 136

- 137 strengthening anti-ferroptosis mechanism (upregulation of Gpx4), the accumulation of

138 iron and the expression of Gpx4 in dental epithelium/mesenchymal cells and

139 extracellular matrix are critically related to the developmental stage.

140 Erastin impairs tooth morphogenesis especially within dental mesenchyme

141 To investigate the possible function of ferroptosis in tooth development, an *ex vivo* 142 culture model of the molar germ was established as described before [23], with or 143 without treatment of erastin. Successful ex vivo culturing of tooth germ from D0 to 144 D7 is presented in Fig S1B. All the molar germs were dissected from E15.5 mousses' 145 mandibles and cultured in medium with or without erastin $(10\mu M)$ for 1, 3, and 5 days 146 (Fig 2A). Comparing Fig 2A-d and 2A-h, gross anatomy showed apparent tiny tooth 147 germ in the erastin treated group than that of CTRL (Control group) in D5 (Day 5). 148 Histological analysis of D5 was performed in Fig 2B; treatment of erastin ($10\mu M$) 149 elevated the number of necrotic-like cells (NLCs), especially in dental mesenchyme 150 (Fig 2B-b' and -d'). We also found a dose-dependent manner of erastin in suppressing 151 tooth morphogenesis (Fig 2C). In D5 incubation of different concentrations (1.5, 5, 10, 152 and 20µM) of erastin, tooth germ deteriorated morphologically (decreasing size, Fig 153 2C-a to -e) and histologically (elevating number of NLCs within dental mesenchyme, 154 Fig 2C-f' to -j') according to the increasing concentration of erastin of erastin 155 treatment. Relative development suppression of tooth germs was then calculated by 156 height, width, and coronal area compared to the CTRL (Fig 2D); The radar graph 157 showed significant dose-dependent impairment of erastin to tooth germ, the original 158 bar graphs are listed in Fig S2.

159 Ferroptosis is activated in erastin-treated molar germ

160 Accurate identification of ferroptosis should be determined by series markers, 161 including iron concentration, lipid peroxidation, mitochondria dysmorphia, and 162 overexpression of ferroptotic genes. We estimated the activation of ferroptosis in 163 tooth germ. Indicated by TPE-o-Py detection in Era-10µM of D5 (Fig 3A-a to 3A-b"), 164 more iron is accumulated than that in CTRL and the region of high concentration 165 mainly located in dental mesenchyme (red, Fig 3A-a and 3A-b). Lipid peroxidation 166 was indicated by the expression of 4-HNE; the distribution pattern of 4-HNE was 167 similar to iron accumulation which suggested the activation of iron-dependent lipid

168 peroxidation occurred within Era-10µM of D5 (Fig 3A-c to 3A-d"). Results of D1 169 and D3 showed similar patterns in the accumulation of iron and the expression of 4-170 HNE (Fig S3), also a dose-dependent manner in D5 (Fig S4). Morphological changes 171 of mitochondria revealed severe shrinkage of mitochondria in dental mesenchyme of 172 Era-10µM (Fig 3B). The size of mitochondria in each group is calculated in Fig 3C; 173 results clearly showed the main size distribution of Era-10µM is significantly 174 decreased. Then, the expression of *Gpx4*, *Slc7a11*, and *Ptgs2* in each tooth germ was 175 modulated by q-PCR. Fig 3D showed Ptgs2, a gene representing the peroxidation 176 level, dramatically increased in tooth germ after erastin treatment, while Gpx4 and 177 Slc7a11, anti-ferroptosis related genes, underwent upregulated but much more 178 moderate expression. All these results contributed to the confirmation of the activated 179 ferroptosis occurred in erastin-treated molar germ, and demonstrated that no lethal 180 concentration of erastin could lead to tooth morphogenesis partially through activation 181 of ferroptosis in dental mesenchyme.

182 *Ferroptotic inhibitors partially rescue erastin-impaired tooth organogenesis*

To further prove the function of ferroptosis in the suppression of tooth germ morphogenesis, a rescue assay was applied by using Fer-1 (classical inhibitor of ferroptosis), Lip-1 (inhibitor of lipid peroxidation), and DFO (iron chelator) to coincubated tooth germ with erastin.

As results shown in Fig 4A, although all these three molecules show rescuing efficiency, Fer-1 holds the highest efficiency in recovering the organogenesis of molar germ in gross anatomy (Fig 4A-c) and reducing the number of NLCs at the histological level (Fig 4A-h and 4A-h'). Calculating the height, width, area of tooth germ (Fig 4B, original bar graphs are listed in Fig S5), and the number of NLCs (Fig 4C) showed inhibitors could partially rescue impairment by erastin, while Fer-1 rescued most.

To avoid bias caused by sectioning, sequential HE slides of CTRL and Era-1.5μM
were applied for 3D reconstruction as previously described [24]. In Fig 5D-a to 5D-j,
tooth germ in CTRL showed well-developed morphology in the 3D models. However,
cusp formation in Era-1.5μM (Fig 4D-e to 4D-h) underwent significant suppression

198 comparing to CTRL in different directions of view; original sequential slides are listed

in Fig S6 to CTRL and Fig S7 to Era-1.5µM. 3D reconstruction was also performed in

200 Era-1.5μM+Fer-1 (Fig 4D-i to 4D-l). Compared to Era-1.5μM (Fig 4D-e to 4D-h),

201 Fer-1 reversed abnormal features dramatically in tooth morphology and cusp

formation. Source sequential sections are listed in Fig S8.

Ferroptosis is the dominant cell death type contributes to erastin-impaired tooth
morphogenesis

205 We further estimated key characteristics of ferroptosis to further assess whether Fer-1 206 rescued erastin-impaired tooth germ through inhibiting ferroptosis. Comparing to 207 CTRL group after 5 days ex vivo culture (Fig 5A-a and 5A-d), iron accumulation and 208 lipid peroxidation (indicated by 4-HNE) in Era-1.5µM is much higher (Fig 5A-b and 209 5A-e). Fer-1 treatment vanished iron accumulation and lipid peroxidation caused by 210 erastin (Fig 5A-c and 5A-f). Results of TEM clearly showed decreased mitochondria 211 shrinkage in Era-1.5 μ M+Fer-1 than that of Era-1.5 μ M (Fig 5B and 5C). All these data 212 convinced that Fer-1 rescued erastin-impaired tooth morphogenesis through inhibiting 213 ferroptosis.

214 Apoptosis is the main innate cell death type in physiological process. To exclude 215 possible over activation of apoptosis induced by erastin treatment, we detected CL-216 CASP3 by IF staining, main executor protein of apoptosis, in each group. In Fig 5D, 217 CL-CASP3 is weakly expressed in CTRL group since the activation of apoptosis is 218 physiologically required in tooth development (Fig 5D). The expression of CL-219 CASP3 slightly increased in both Era-1.5µM and Era-1.5µM+Fer-1, but showed no 220 statistical differences comparing to CTRL group (Fig S9A). TUNEL assay is also 221 performed to identify apoptotic cells by detecting DNA damage, results further 222 convinced that apoptosis is not significantly activated in erastin treated tooth germ 223 (Fig S9B). Taken together, ferroptosis is the dominant cell death led by erastin 224 treatment in impaired tooth germ.

225

226 Discussion

227 In the last several decades, the beneficial role of apoptosis in regulating organ

228 development and tissue regeneration has been identified [25]. Apoptosis in tooth 229 development had been characterized by the activation of Caspase 3 and DNA damage, 230 which revealed a spatiotemporal apoptotic cell death pattern due to different stage of 231 tooth morphogenesis [26]. This raises the question of whether other newly determined 232 non-apoptotic cell death pathways also participate in the physiological processes like 233 development, maintaining homeostasis, ageing, etc. Except for NETosis, a neutrophil 234 extracellular trap-related cell death, the possible function and mechanism of the rest 235 non-apoptotic cell death including, pyroptosis, necroptosis, and ferroptosis, are still 236 barely investigated. Characterized by its close relationship with the metabolism of 237 lipid, iron, and ROS [27,28], risk factors inducing ferroptotic stress and/or activating 238 ferroptosis also critically involved in development and ageing, which made 239 ferroptosis and its regulatory mechanism, other than apoptosis, pyroptosis, and 240 necroptosis, a promising undiscovered type of cell death during tooth development.

241 To explore the potential involvement of ferroptosis in organogenesis, we have 242 developed an *ex vivo* culture model of tooth germ morphogenesis. This system allows 243 for the application of erastin to induce significant activation of ferroptosis throughout 244 the developmental process, offers a valuable opportunity to investigate the specific 245 mechanisms underlying the role of ferroptosis in organ development. Our study 246 detected the expression of Gpx4 and the accumulation of iron both in mouse 247 mandibular incisor and developing first molar. Although Gpx4 expressed ubiquitously 248 in both tooth germ and incisor of mouse, its abundance differs in different cell types. 249 Results revealed an increasing iron accumulation both in odontoblast and ameloblast 250 according to the developmental process. The spatiotemporal expression of Gpx4 is 251 also positively linked to odontogenesis and amelogenesis. This phenomenon indicated 252 growing ferroptotic stress (accumulating iron) and strengthening anti-ferroptosis 253 mechanism (increasing expression of Gpx4) physiologically coexisted and may 254 maintain a critical balance along with the differentiation and maturation of 255 odontoblast and ameloblast. However, the discovery of changes in iron and lipid 256 metabolism during tooth morphogenesis is not novel. In the 1930s, pioneer scientists 257 in dental biology had already identified the presence of iron in the tooth of different

258 animals [29-31], then found some defects of enamel in mouse is related to abnormal 259 iron metabolism [32]. Lipid metabolism and lipid peroxidation, the other core risk 260 factors of ferroptosis, were also described in the early stage of dental biology research 261 [20,33,34]. Even when the risk factors of ferroptosis had been reported to participate in tooth development, there is still no reports about the exact ferroptosis related tooth 262 263 developmental defects. Our results provided a new perspective to reconsider the 264 underlying function of all these ancient studies in a comprehensive manner. They 265 illustrated the importance of the Gpx4-dependent anti-ferroptosis pathway in 266 managing all these already existing ferroptotic stress during tooth morphogenesis. 267 Future *in vivo* studies utilizing transgenic mice are needed to systematically analyze 268 the role of ferroptosis/ferroptotic stress during tooth and other organ development. 269 To further investigate the meaning of this precarious balance between ferroptotic 270 stress and expression of Gpx4, we use erastin to inhibit the internalization of GSH, 271 which is the critical substrate for Gpx4 to protect cells from lipid peroxidation. The 272 developmental role of Gpx4 had been studied even before the ferroptosis was 273 formally described (before 2012). In situ hybridization indicated expression of Gpx4 274 in all developing germ layers during gastrulation and in the somite stage in the 275 developing central nervous system and in the heart [35], which made Gpx4 (-/-) mice 276 die embryonically in utero by midgestation (E7.5) and are associated with a lack of 277 normal structural compartmentalization [36]. Specific deletion of Gpx4 during 278 developmental process were found to participate in the maturation and survival of cerebral and photoreceptor cell [37,38]. Recent years, more ferroptosis related 279 280 function of Gpx4 were discovered in neutrophil [39] and chondrocyte [40] of adult 281 mice, in which specific deletion will lead to ferroptosis-induced organ dysregulation 282 and degeneration. Thus, it is essential to assess the biological function Gpx4-283 dependent ferroptotic suppressing system. In our study, erastin significantly impaired 284 tooth morphogenesis in a dose-dependent manner within the ex vivo tooth germ 285 culture model. The ex vivo organ culture of tooth germ is a well-established classical 286 model for the study of tooth morphogenesis. Compared to in vivo study, the ex vivo 287 culture of tooth germ is convenient for manipulating culture conditions and 288 investigate factors affecting tooth germ in a high throughput way [41]. Although 289 lacking of circulation and immune system, the ex vivo culture of the tooth germ, other 290 than the traditional 2D culture of dental cells in vitro, can retain most properties of 291 tooth development, like interactions among oral epithelia, mesenchyme, and stromal 292 cells. We successfully established this model and the tooth germs from D0 to D7 are 293 well developed (Fig S1B). To induce ferroptosis, erastin is the most used agent 294 inhibiting GSH transport but is not stable *in vivo*, which makes *ex vivo* organ culture 295 of tooth germ the ideal way to study the possible function of ferroptosis/ferroptotic 296 stress in tooth morphogenesis. Moreover, according to our results, erastin treatment 297 will not induce overactivation of apoptosis in tooth germ (Fig 5D).

298 The histologic analysis by HE staining showed an increased number of NLCs 299 located in dental mesenchyme of erastin-treated tooth germ. 3D reconstruction of all 300 the slides convinced that necrotic mainly occurred in the region of the odontoblast 301 layer. Unlike CL-CASP3 to apoptosis, membrane localization of GSDM family 302 proteins and MLKL to pyroptosis and necroptosis respectively, ferroptosis is a special 303 type of cell death which have unique inducer but no special proteins reflecting its 304 activation. Thus, ferroptosis in erastin-treated tooth germ is determined by the 305 accumulation of iron, upregulation of 4-HNE, shrunken and condensed mitochondria, 306 dramatically upregulated expression of *Psg2* (risk marker to ferroptosis), and mildly 307 increased expression of Gpx4, Slc7a11 (protective factor to ferroptosis). These results 308 indicated that erastin could lead to abnormal tooth morphology via activating 309 ferroptosis. Moreover, characterized by more apparent NLCs, stronger iron and 4-310 HNE fluorescence signal, and severer mitochondria degeneration, dental mesenchyme 311 cell/odontoblast seems more sensitive to erastin-induced ferroptosis than that of dental 312 epithelium cell/ameloblast.

According to our results, ferroptotic stress is physiologically increased during tooth development. Thus, rather than "initially activate ferroptosis" in tooth morphogenesis, erastin, a system x_c^- inhibitor, inhibits GSH production, and induced ferroptosis in tooth germ is more likely lead by "overburdened ferroptotic stress". As a multi-step process of cell death, the different inhibitor has the different target to suppress 318 ferroptosis. Here in our study, regular ferroptosis inhibitors were applied in the rescue 319 assay of impaired tooth morphogenesis. Tooth germ was treated by Fer-1, Lip-1 320 (radical trapping agents which inhibit the propagation of lipid peroxidation), and DFO 321 (an iron chelator). Surprisingly, although all these inhibitors could reverse erastin-322 induced impairment of tooth morphogenesis, Fer-1 reduced the number of NLCs 323 much more efficiently than lip-1 and DFO. Different from lip-1 or DFO, which only 324 targets lipid peroxidation or labile iron accumulation, Fer-1 could both inhibit lipid 325 peroxidation and reduce the labile iron pool in cells, notably is not consumed while 326 inhibiting iron-dependent lipid peroxidation [42]. Sustaining double effects of Fer-1 in 327 inhibiting ferroptosis possibly enables it effectively rescue the impairment of tooth germ than other agents. 328

In conclusion, by using *ex vivo* culture model of tooth germ, we identified continuing accumulation of ferroptotic stress in both odontogenesis and amelogenesis during tooth development. Activation of ferroptosis impaired tooth morphogenesis, especially within dental mesenchyme, and could be partially rescued by ferroptotic inhibitor. This study provides a promising model to effectively investigate the developmental role of ferroptosis and will broaden our knowledge about possible involvement of non-apoptotic RCD during organogenesis.

336

337 Materials and methods

338 Animals and organ culture

339 ICR mice were purchased from Chengdu Dossy Experimental Animals Co., Ltd (Sichuan, China). All animal work was done according to the National Institutes of 340 341 Health guidebook and approved by the Committee on the Ethics of Animal 342 Experiments of Sichuan University (WCHSIRB-D-2021-12544). The presence of a 343 vaginal plug was used as an indication of embryonic day 0 (E0). The first mandibular 344 molar tooth germs of E15.5 were dissected under stereomicroscope (Carl Zeiss, 345 Germany). All the dissection steps were performed on ice. The dissected tooth germs 346 were placed in the upper chamber of the Trans-well (3450, Corning, USA) and four

347 tooth germs were placed on one dish. The culture medium and drugs were mixed and 348 placed in the lower chamber. Tooth germs were cultured in DMEM/F12 supplemented 349 with 10% fetal bovine serum, 50 U/ml penicillin/streptomycin, and 100 µg/ml 350 ascorbic acid and incubated at 37°C and 5% CO₂. Medium was changed every other 351 day. Tooth germs were cultured for 5 days in the presence of different concentration 352 of Erastin (0µM, 1.5µM, 5µM, 10µM, 20µM). Culture medium added with Fer-353 1(Ferrostatin-1 1µM), Lip-1(Liproxstatin-1, 200nM), DFO (Deferasirox, 100µM) 354 were used to pretreat molar germ for 4 hours. All these small molecules were 355 purchased from Selleck. After 5 days of cultivation, the tooth germs were fixed in 4% 356 PFA, embedded in paraffin. Tooth germs were photographed at 0-day, 1 day, 3 day, 5 357 day and each tooth size (width, height and area) were measured (n=9 for each group) 358 by using the modified method reported in the literature previously. Each experiment 359 was repeated 3 times [43].

360 Immunohistochemistry

361 Sections were dewaxed and rehydrated before antigen repair, and then cooled to room

temperature. Sections were incubated overnight at 4°C with Gpx4 (1:200, Invitrogen,

PA5-102521) in 2% bovine serum albumin (BSA)/phosphate buffered saline (PBS),
pH 7.4. Negative control sections were incubated with 2% BSA/PBS. After washed,
sections were then incubated with SignalStain® Boost IHC Detection Reagent (8114S,
Cell Signal Technology, USA) and then washed and incubated with diaminobenzidine
(DAB) to detect any reactions and were examined by light microscopy after
counterstaining with hematoxylin.

369 Immunofluorescence and iron probe staining

Tissue sections were treated with phosphate-buffered saline (PBS) with 0.5% Triton
X-100 for permeabilization. After washing and blocking, sections were incubated with
4-HNE (1:200, Abcam, ab48506), cleaved Caspase 3 (1:200, Cell Signaling
Technology, 9664) in blocking buffer (PBS and 2% bovine serum albumin) overnight
at 4□. After washed, second antibody Alexa Fluor® 488 (1:200, Abcam, ab150077),
Alexa Fluor® 594 (1:200, Abcam, ab150116) mixed with 4',6-diamidino-2-

phenylindole (DAPI) was applied. Incubate at room temperature for one to two hoursand then wash and seal the slices.

378 To accurately detect the changes of iron accumulation within differently cultured 379 tooth germs, we use an aggregation-induced emission featured iron (III) probe from 380 ortho-substituted pyridinyl-functionalized tetraphenylethylene (TPE-o-Py) [22], 381 which is kindly gifted by Prof. Youhong Tang. This probe displays high sensitivity 382 and selectivity toward iron (III) detection. The recognition arises from the position 383 isomer of ortho-substitution, and the fact that TPE-o-Py has a low acid dissociation constant (pKa) that is close to that of hydrolyzed Fe^{3+} . The iron probe staining was 384 385 performed as described. Briefly, TPE-o-Py was dissolved in THF (Tetrahydrofuran) 386 before being added to PBS and diluted to working concentration of 20μ M. The 387 working solution was then placed on sections, and incubated at room temperature for 388 half an hour and then aspirated and sealed, then a laser confocal microscope 389 (Olympus FV3000, Japan) was used to detect the fluorescence signal. Since the TPE-390 o-Py probe pronounced red-shift in fluorescence emission which is positively related to the concentration of iron, low concentration of Fe³⁺ was detected under 391 392 fluorescence channel of Alexa Fluor 405 (Excitation wave length: 402nm, Emission 393 wave length: 421nm), while Alexa Fluor 594 (Excitation wave length: 590nm, 394 Emission wave length: 617nm) for high concentration.

395 *Tissue preparation for transmission electron microscope*

Prefixed with a 3% glutaraldehyde, then the tissue was postfixed in 1% osmium tetroxide, dehydrated in series acetone, infiltrated in Epox 812 for a longer, and embedded. The semithin sections were stained with methylene blue and Ultrathin sections were cut with diamond knife, stained with uranyl acetate and lead citrate.
Sections were examined with JEM-1400-FLASH Transmission Electron Microscope.

401 **RNA extraction and qPCR**

402 Total RNA of tooth germs (E15.5 cultured for 5 days) were extracted by using 403 TRIzol[™] Reagent (Invitrogen, USA) according to instruction of manufacturer. The 404 amount and the integrity of RNA were assessed by measurement of absorbance at 260 405 and 280 nm. First-strand cDNA synthesis was performed with a HiScript III Q RT

- 406 SuperMix for qPCR (Vazyme, China). The levels of Gpx4, Ptgs2, Slc7all were
- 407 measured by quantitive real-time PCR (Bio-Rad, USA) with ChamQ Universal SYBR
- 408 qPCR Master Mix (Vazyme, China) and normalized to the level of β -Actin mRNA.
- 409 These experiments were performed in triplicate. Primer sequences used in qPCR are
- 410 listed below.

Gene	Sequence
Gpx4	F: CCTCCCCAGTACTGCAACAG
	R: GGCTGAGAATTCGTGCATGG
Ptgs2	F: CTGCGCCTTTTCAAGGATGG
	R: GGGGATACACCTCTCCACCA
Slc7a11	F: GATGGTCCTAAATAGCACGAGTG
	R: GGGCAACCCCATTAGACTTGT
β -Actin	F: AGATGTGGATCAGCAAGCAG
	R: GCGCAAGTTAGGTTTTGTCA

411 3D reconstruction of tooth germ sections

412 Sequential section and subsequent HE staining of each tooth germ were performed as
413 described before. Digital pathological system (Olympus vs200) was used to scan all
414 the stained sections and reconstructed by following previously described protocol [24].

415 TUNEL assay

We used a One Step TUNEL Apoptosis Assay Kit (Beyotime, C1090) to detect the
possible DNA damage in apoptotic cell of tooth germs. The TUNEL assay was
performed following instructions.

419 *Statistical analysis*

420 Analysis of Gpx4 relative expression levels were carried out by Image-pro plus7.0 421 (Media Cybernetics, USA). Statistical analyses were carried out using the GraphPad 422 Prism version 8.00 (GraphPad Software, San Diego, CA, USA). All statics were 423 shown as the arithmetic mean \pm the standard error of the mean. The significance of 424 differences between groups was tested by using the one-sample t-test. Differences 425 were considered significant when p < 0.05.

426

427

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437 Contributed to conception and design critically revised manuscript; Fanyuan Yu: 438 Contributed to conception and design, critically revised manuscript; Ling Ye: 439 Contributed to conception and design, critically revised manuscript. All authors gave 440 their final approval and agree to be accountable for all aspects of the work. The 441 authors declare no potential conflicts of interest with respect to the authorship and/or 442 publication of this article.

443

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571 Figures and Figure Legends

572 Figure 1 Spatiotemporal characterization of Gpx4 expression and iron accumulation 573 in tooth morphogenesis. (A) (i~v) HE staining for tooth germ in E13.5 to PN3, scar 574 bars=200µm; (a~e) Gpx4 expression detected by IHC, scar bars=200µm; (a'~e') Enlarged view of each Gpx4 staining, scar bars=50µm; epithelia versus mesenchyme 575 ***P < 0.001 (B) (i~v) Iron probe staining in for tooth germ in E13.5 to PN3, scar 576 bars=200µm; (a~e) Enlarged view of selected region, scar bars=100µm, low 577 concentration (a'~e', blue) and high concentration of iron (a''~e'', red) are present, 578 scar bars=100µm. Epi=Epithelia, Mes=Mesenchyme. 579 580 **Figure 2** Erastin impairs tooth morphogenesis especially within dental mesenchyme. 581 (A) Gross anatomy of tooth germ cultured *ex vivo* for five days, scar bars= 500μ m; (B) 582 (a, c) HE staining for tooth germ in D5, scar bars= 100μ m; (b, d) High resolution of 583 epi-mes junction papilla, scar bars=50µm; Black stars pointe out NLCs; (b', d') NLCs 584 indicated by black stars. (C) (a~e) Gross anatomy of tooth germs in different 585 concentrations of erastin in D5, scar bars= $500\mu m$; (f~j) HE staining of different 586 concentration treated tooth germ in D5; $(f' \sim j')$ High-resolution view of epi-mes 587 junction region of each tooth germ, scar bars= 50μ m; (**D**) Rada graph for calculation 588 of height, width, and area in each tooth germ. Black dotted line outlines ameloblasts,

589 AM=ameloblast, DP=dental papilla.

590 Figure 3 Ferroptosis is activated in erastin-treated molar germ. (A) (a, b) High density of Fe³⁺ (red) in CTRL and Era-10µM of D5, white star pointed out strong 591 fluoresce signal of Fe^{3+} , scar bars=50µm; (a', b') Low density of Fe^{3+} (grey) and (a'', 592 b") merged view of iron probe staining; (c, d) Merged view of IF straining of 4-HNE 593 594 (Magenta) and DAPI (blue), white star pointed out strong fluoresce signal of 4-HNE, 595 scar bars=5050µm; (c', d') for DAPI and (c", d") for 4-HNE; AM=ameloblast, 596 DP=dental papilla, HD=high density, LD=low density; (B) TEM scanning for CTRL 597 and Era-10 μ M in D5; (a, d) Epi-Mes junction area of CTRL and Era-10 μ M in D5 are 598 detected, scar bars= 5μ m; (b, c) representative view of cells in epithelia and 599 mesenchyme for CTRL, scar bars=2µm; Black arrow pointed out typical 600 mitochondria in each region (b') for epithelia and (c') of mesenchyme, outlined by 601 white dotted line; (e, f) representative view of cells in epithelia and mesenchyme for 602 Era-10 μ M, scar bars=2 μ m; Black arrow pointed out typical mitochondria in each 603 region (e') for epithelia and (f') of mesenchyme, outlined by white dotted line; (C) 604 Relative frequency of the mitochondrial size in both groups; (D) Fold changes of gene 605 expression in CTRL and Era-10 μ M in D5, versus CTRL ****P*<0.001.

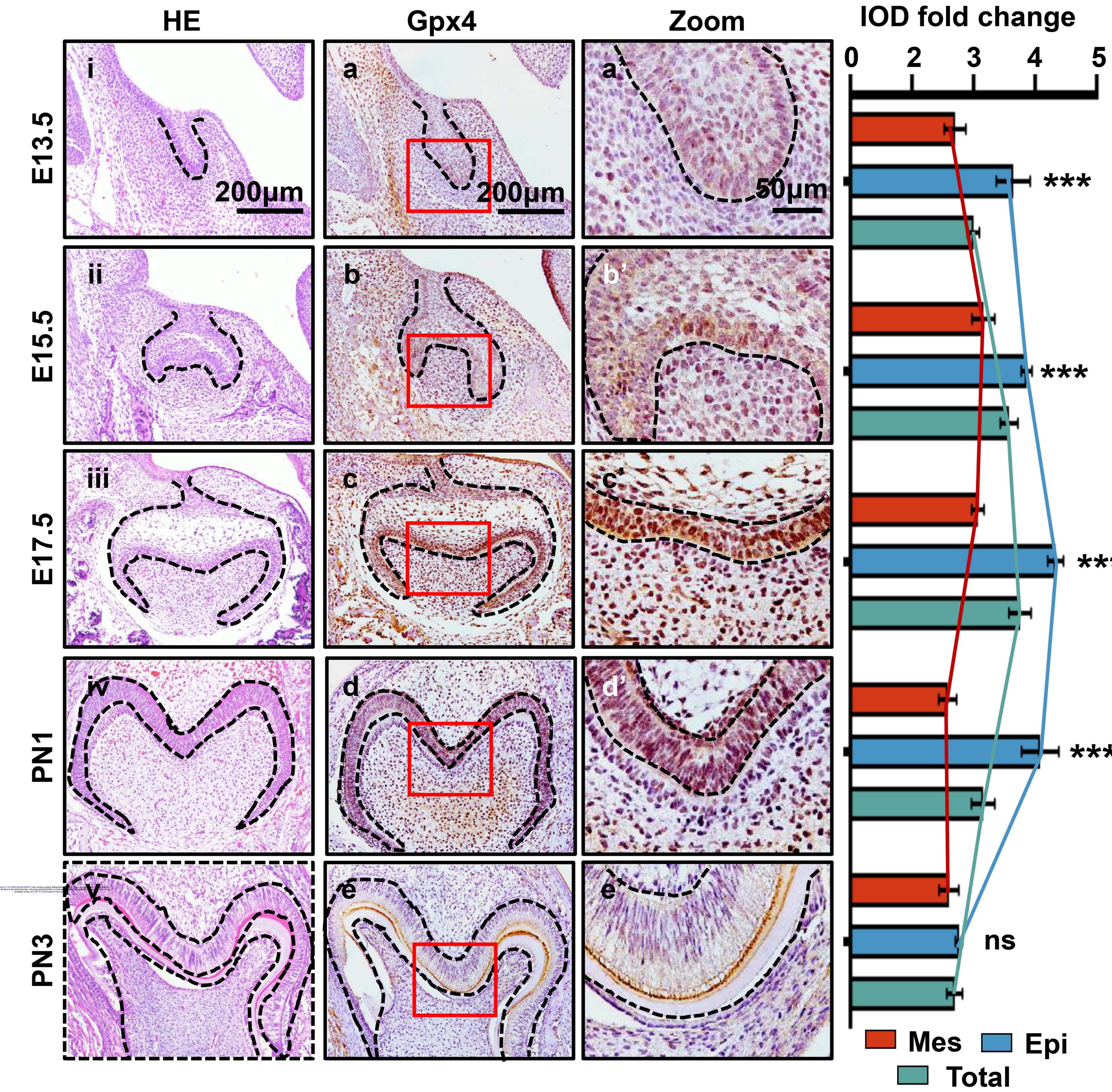
606 **Figure 4** Ferroptotic inhibitors partially rescue erastin-impaired tooth organogenesis. 607 (A) (a~e) Gross anatomy of tooth germs in differently treated group, scar bars= $500 \mu m$; 608 (f~j) HE staining of differently treated tooth germ, scar bars=100 μ m; (f'~j') High 609 resolution view of epi-mes junction region of each tooth germ; Black dotted line 610 outlines ameloblasts, black star pointed out NLCs, AM=ameloblast, DP=dental 611 papilla, scar bars= $50\mu m$; (B) Rada graph for calculation of height, width, and area in 612 each tooth germ; (C) Average number of NLCs in each group, versus Era- 1.5μ M 613 ***P < 0.001, versus Era-1.5 μ M+Lip-1 ##P < 0.01; (**D**) 3D reconstructed view of tooth 614 germ in D5. (a~d) CTRL from the front view, coronal plane, sagittal plane, and 45° 615 side view; (e~h) to Era-1.5 μ M and (i~l) to Era-1.5 μ M+Fer-1 were viewed by the same

616 way. Scar bars= $100\mu m$.

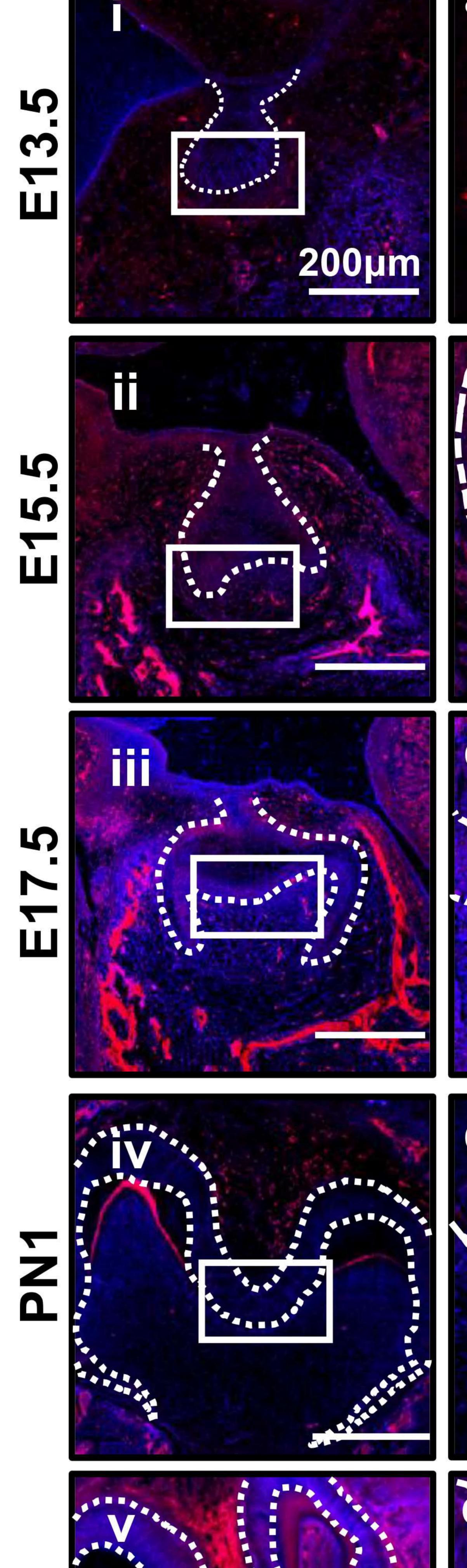
617 Figure 5 Ferroptosis is the dominant cell death type contributes to erastin-impaired tooth morphogenesis. (A) (a~c) High density of Fe³⁺ (red) in CTRL, Era-1.5µM and 618 Era-1.5 μ M+Fer-1 of D5, white star pointed out strong fluoresce signal of Fe³⁺, scar 619 bars=50µm; (a'~c') Low density of Fe³⁺ (grey) and (a''~c'') merged view of iron 620 621 probe staining; (d~f) Merged view of IF straining of 4-HNE (Magenta) and DAPI 622 (blue), white star pointed out strong fluoresce signal of 4-HNE, scar bars= $50\mu m$; (d'~ 623 f') for DAPI and (d"~ f") for 4-HNE; AM=ameloblast, DP=dental papilla, HD=high 624 density, LD=low density; (B) TEM scanning for Era-1.5µM and Era-1.5µM+Fer-1 in 625 D5; (a, d) Epi-Mes junction area of Era-1.5 μ M and Era-1.5 μ M+Fer-1 in D5 are 626 detected, scar bars= $5\mu m$; (b, c) representative view of cells in epithelia and 627 mesenchyme for Era-1.5µM, scar bars=2µm; Black arrow pointed out typical 628 mitochondria in each region (b') for epithelia and (c') of mesenchyme, outlined by white dotted line; (e, f) representative view of cells in epithelia and mesenchyme for 629

630 Era-1.5μM+Fer-1, scar bars=2μm; Black arrow pointed out typical mitochondria in

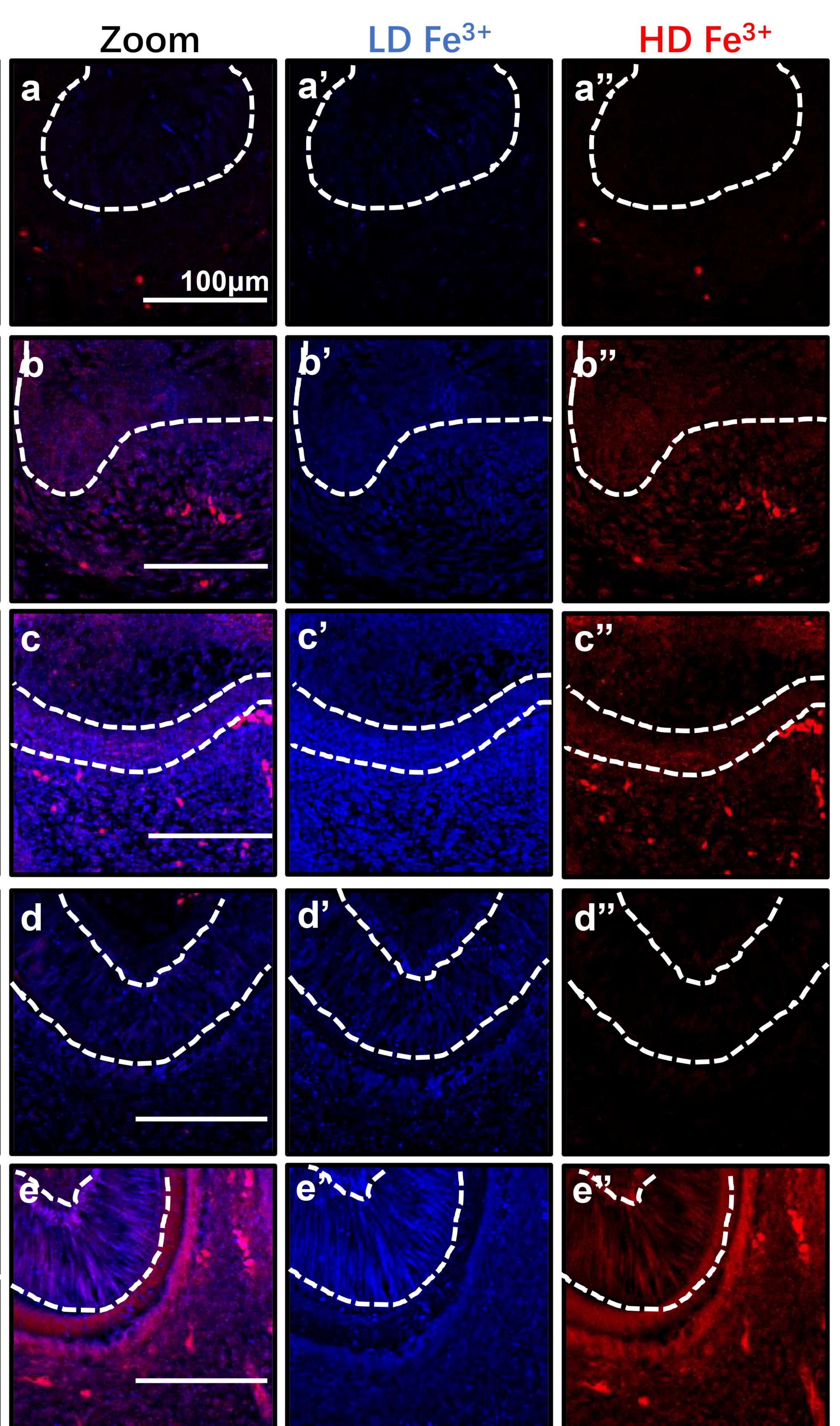
- each region (e') for epithelia and (f') of mesenchyme, outlined by white dotted line;
- 632 (C) Relative frequency of the mitochondrial size in both groups. (D) (a~c) Expression
- 633 of CL-CASP3 (green) in CTRL, Era-1.5 μ M and Era-1.5 μ M+Fer-1 in D5, scar
- 634 bars=200μm; (a'~c') Enlarged view of CL-CASP3 in each group, scar bars=50μm. (E)
- 635 Schematic model illustrates the overburdened ferroptotic stress impaired tooth636 morphogenesis in *ex vivo* organ culture model.
- **Figure supplement 1** (A) Negative control of different developmental stage of molar
- 638 germ in Gpx4 IHC staining. (B) *Ex vivo* culture of tooth molar germ from D0 to D7.
- 639 Scar bars= $20\mu m$.
- Figure supplement 2 Original bar graphs of height, width, and area of erastin-treated tooth germ. *P < 0.05, **P < 0.01 * **P < 0.001
- 642 Figure supplement 3 Results for iron accumulation and 4-HNE expression in Era-
- 643 10μ M of D1 and D3. Scar bars=50 μ m
- 644 Figure supplement 4 Results for iron accumulation and 4-HNE expression in Era-
- 645 1.5 μ M, Era-5 μ M, Era-10 μ M, and Era-20 μ M of D5. Scar bars=50 μ m
- **Figure supplement 5** Original bar graphs of height, width, and area of tooth germ in
- 647 rescue assay. **P*<0.05, ***P*<0.01****P*<0.001
- **Figure supplement 6** Source sequential HE slides for 3D reconstruction of CTRL
- Figure supplement 7 Source sequential HE slides for 3D reconstruction of Era1.5μM
- Figure supplement 8 Source sequential HE slides for 3D reconstruction of Era1.5μM+Fer
- **Figure supplement 9** (A) Mean influence intensity of ROI of CL-CASP3 in CTRL,
- 654 Era-1.5μM, and Era-1.5μM+Fer-1. Both Era-1.5μM and Era-1.5μM+Fer-1 showed
- slightly increased CL-CASP3 activation than that in CTRL but showed no statistical
- 656 differences. (B) Results of TUNEL assay in each experimental group. DNA damage
- 657 detected by TUNEL assay is similar to that of CL-CASP3, which indicated that
- apoptosis is not significantly activated in erastin treated tooth germ. Scar bars=200µm
- 659

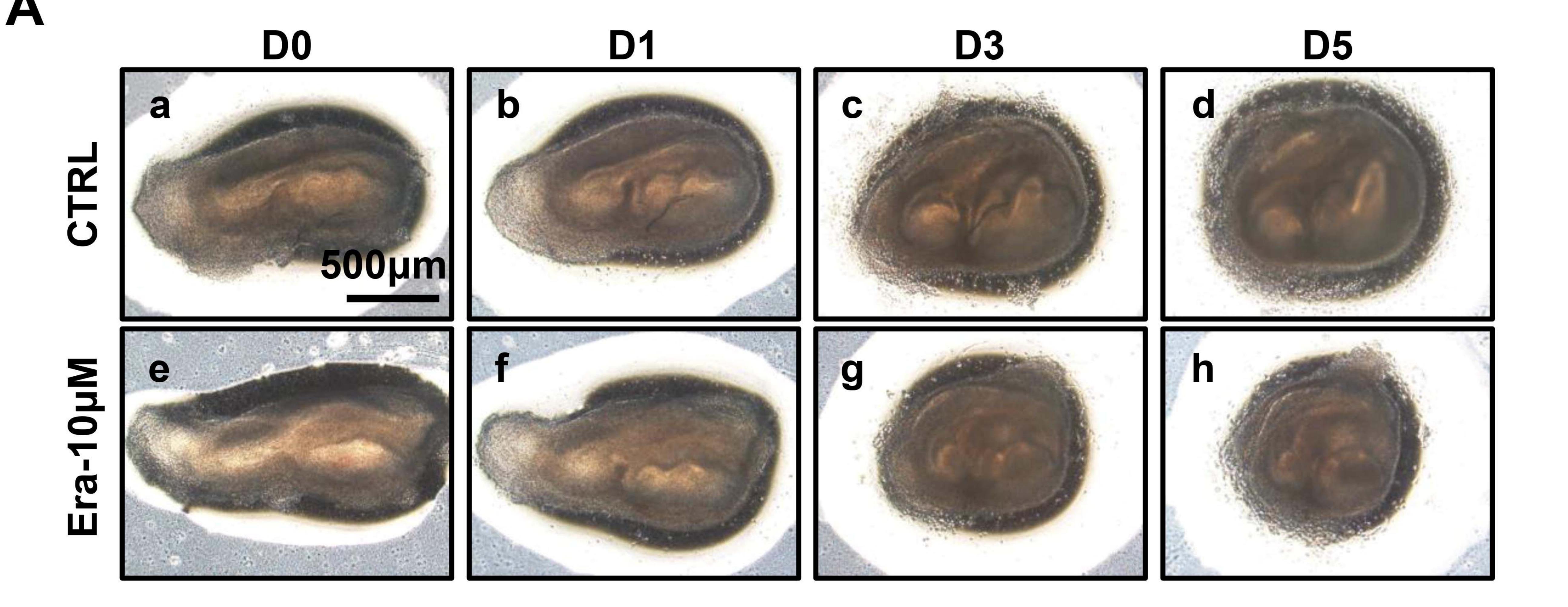






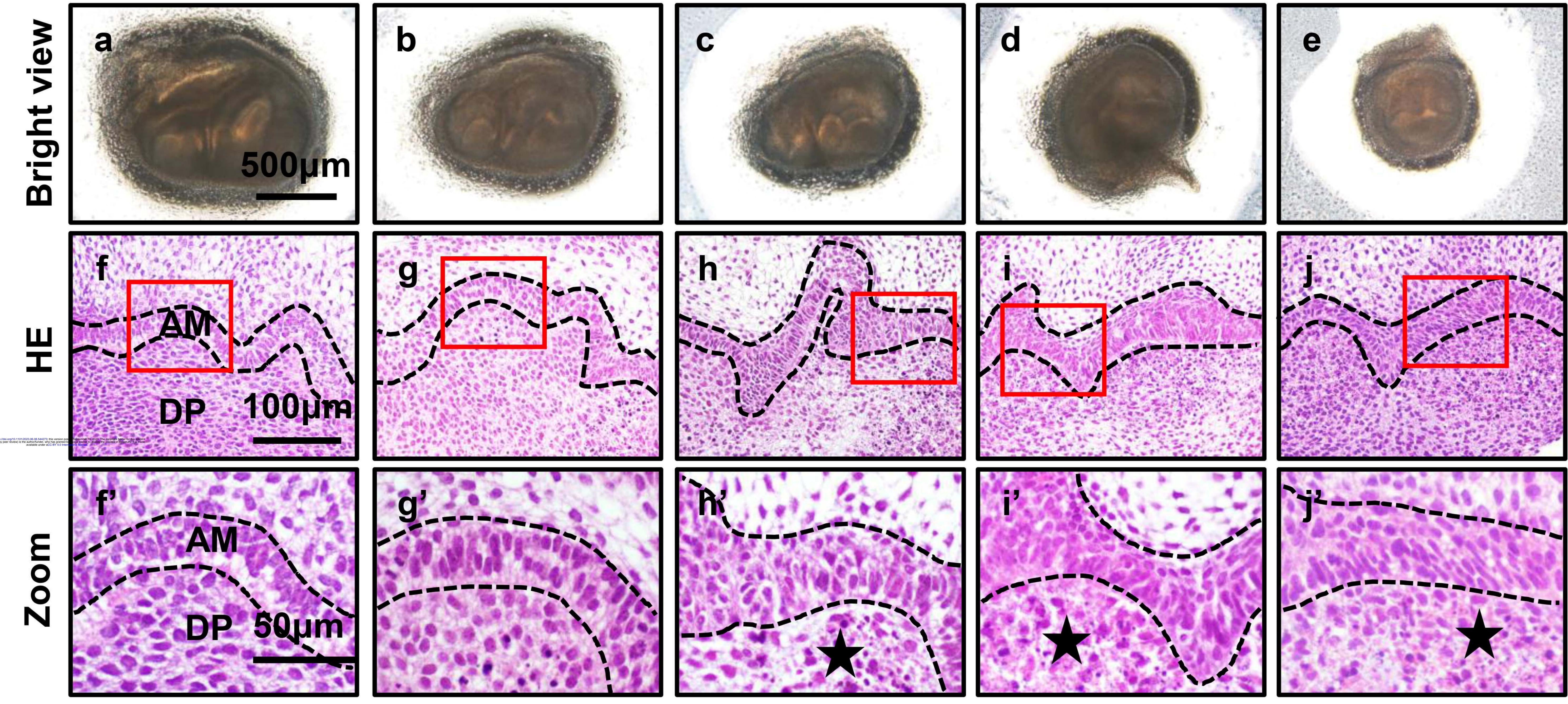
Overview





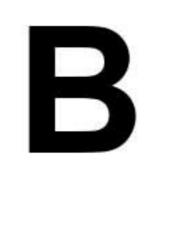


Era-1.5µM

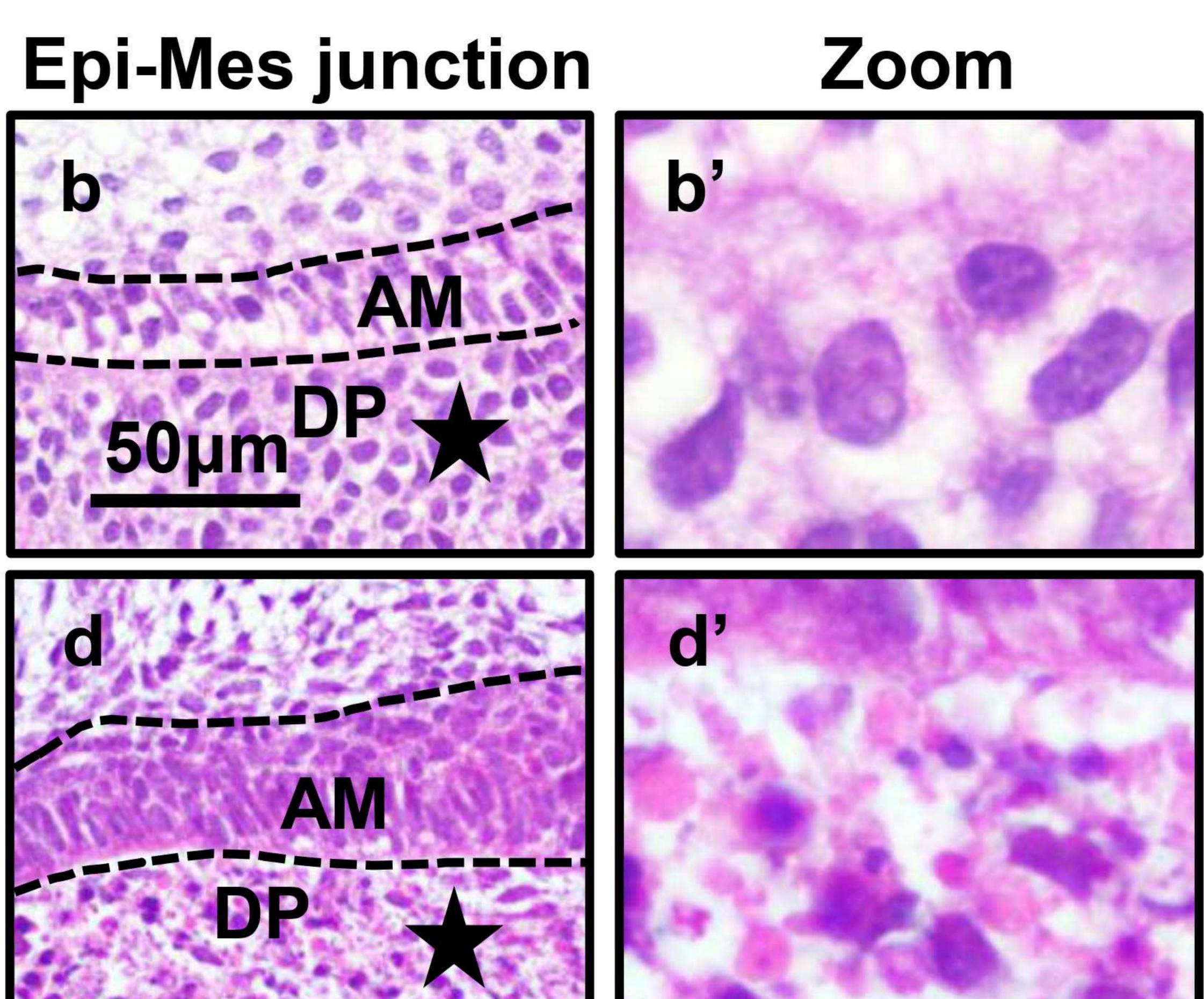


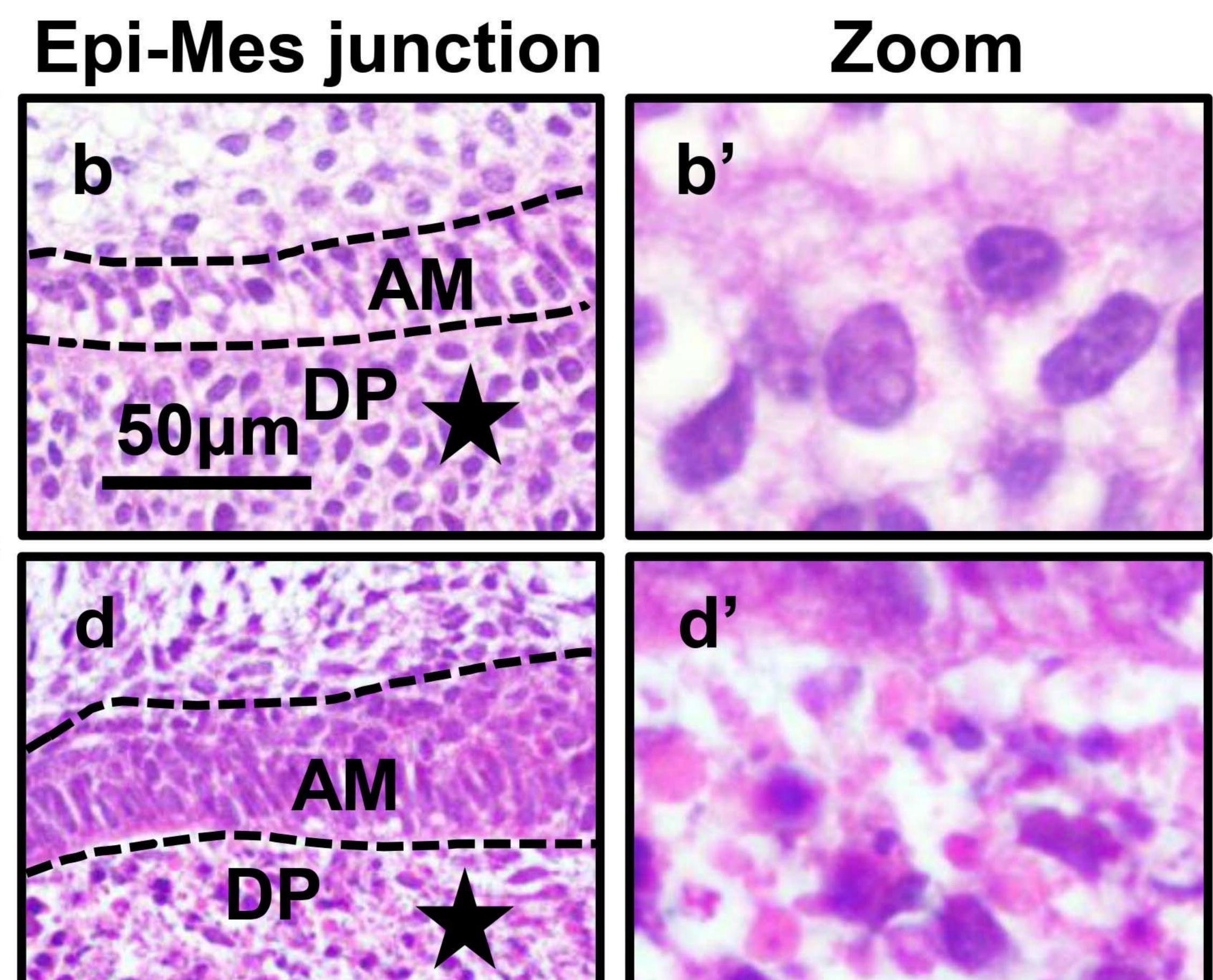
Era-5µM

Era-10µM



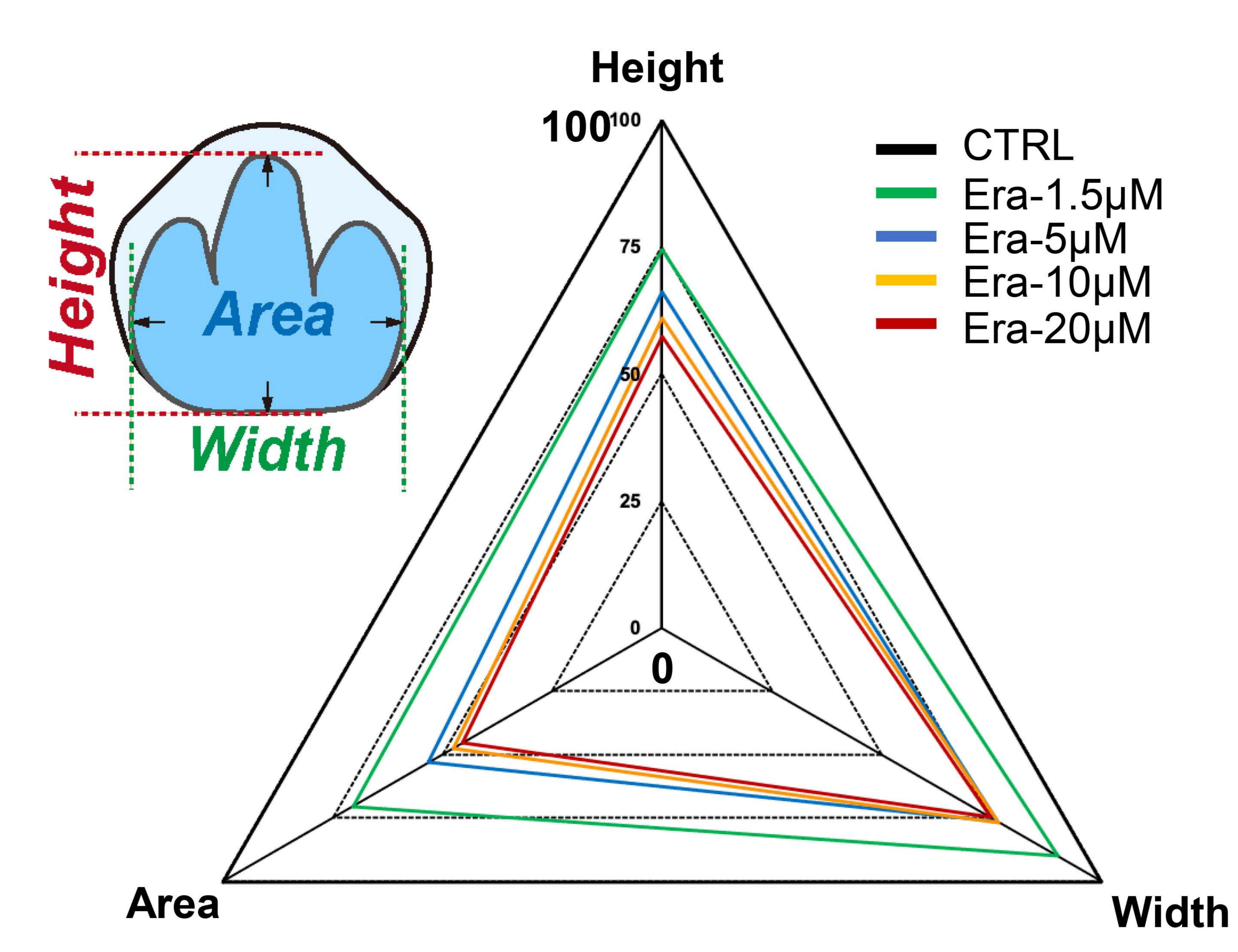
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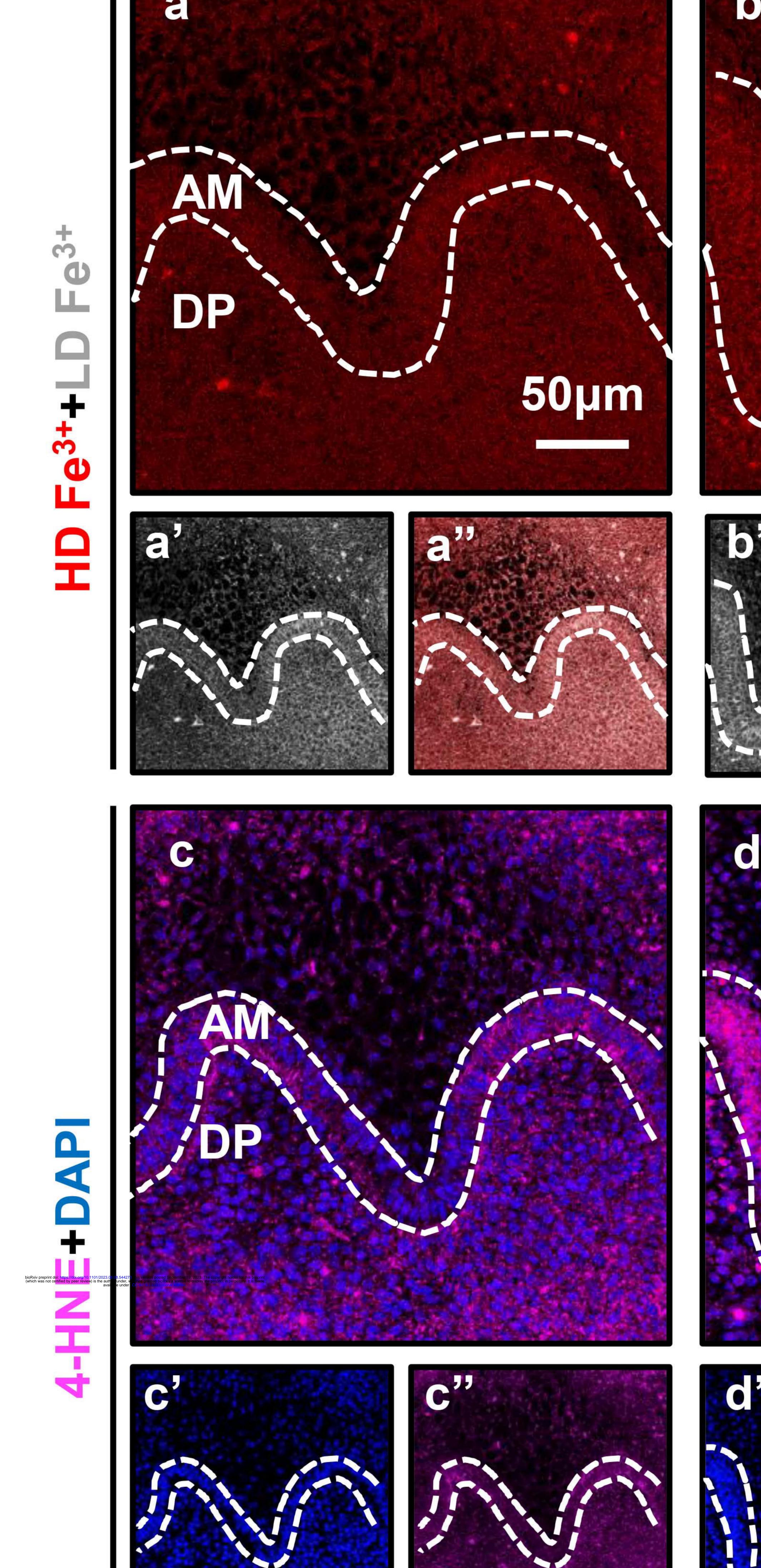




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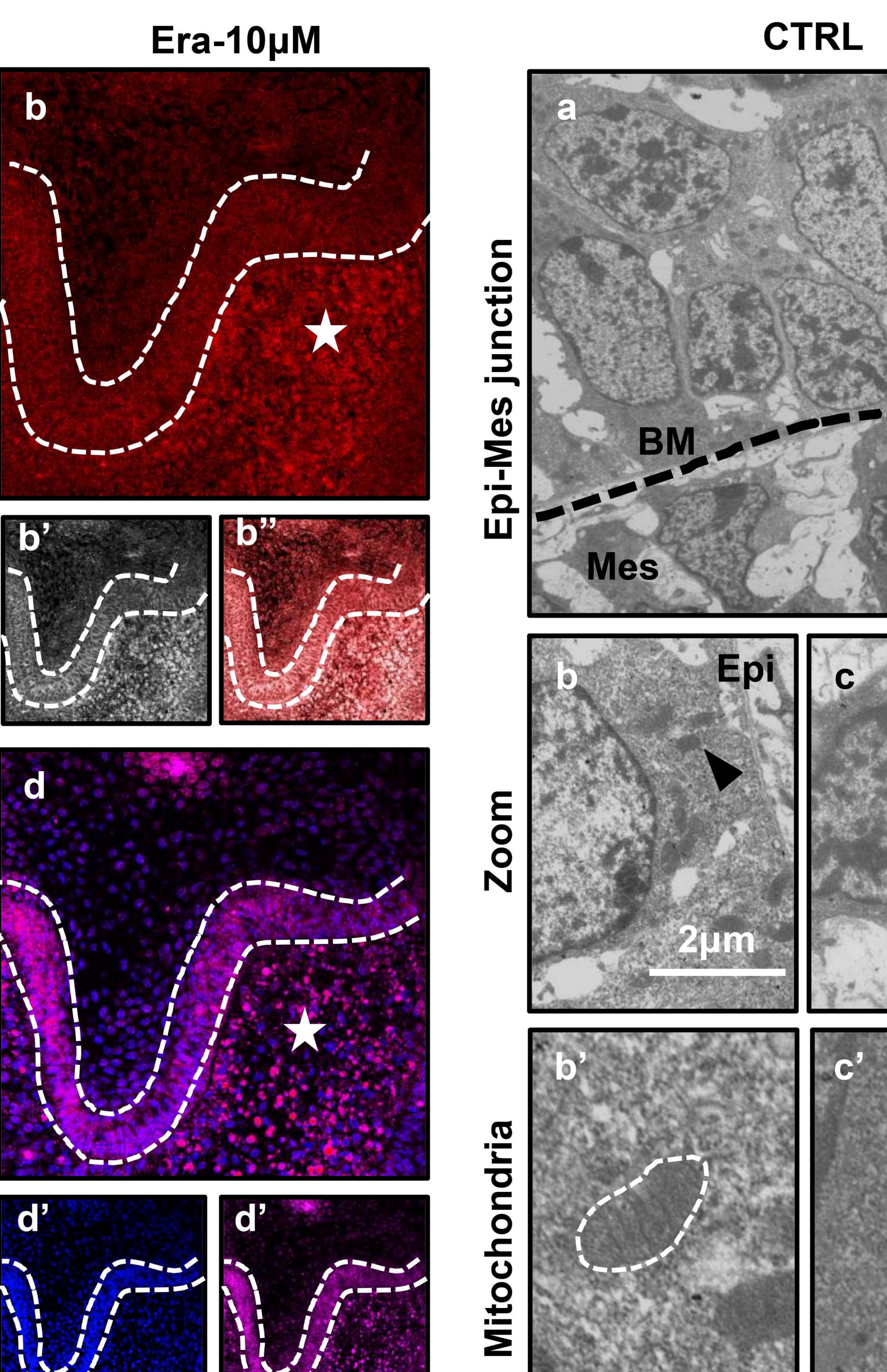




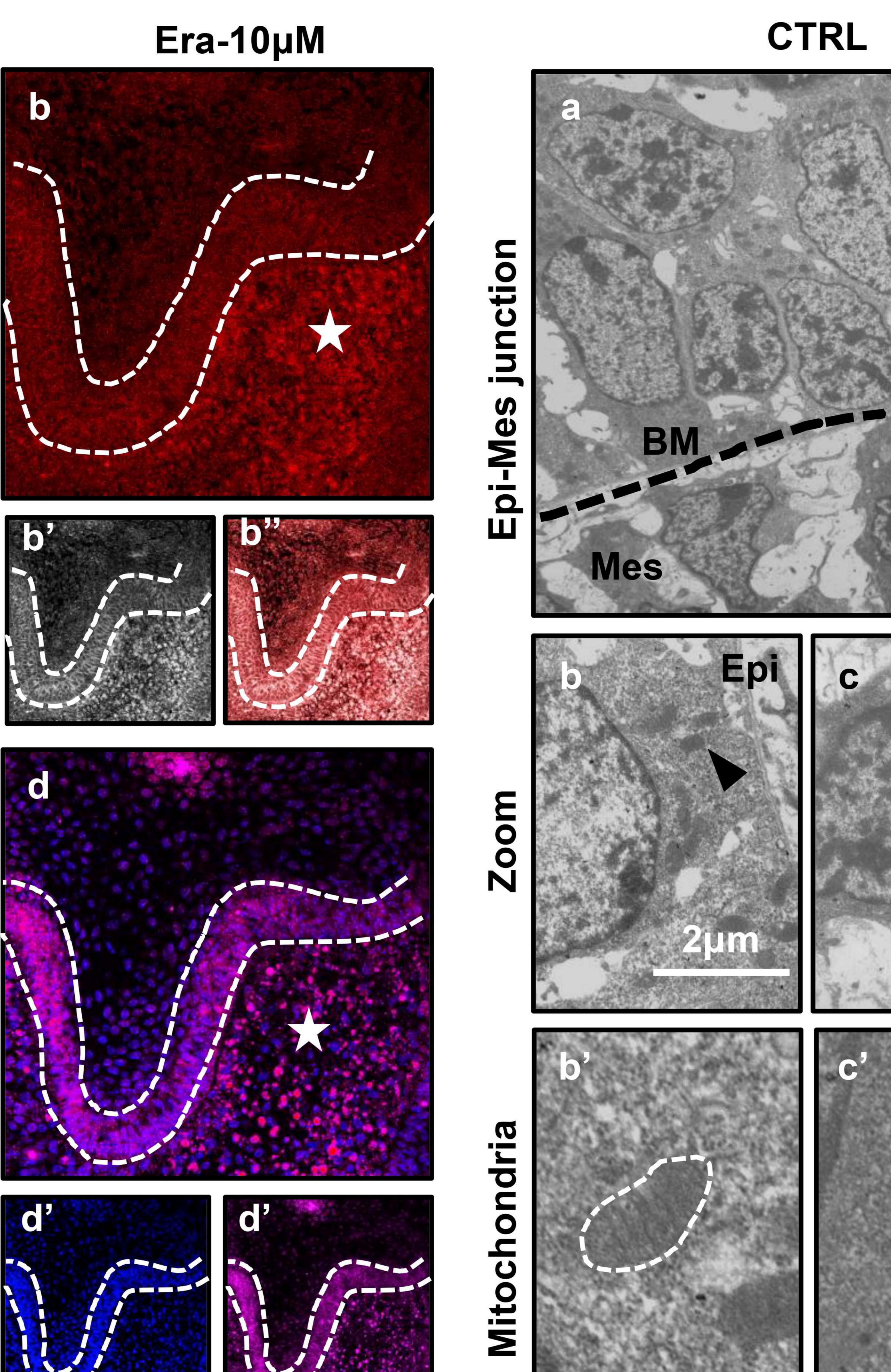


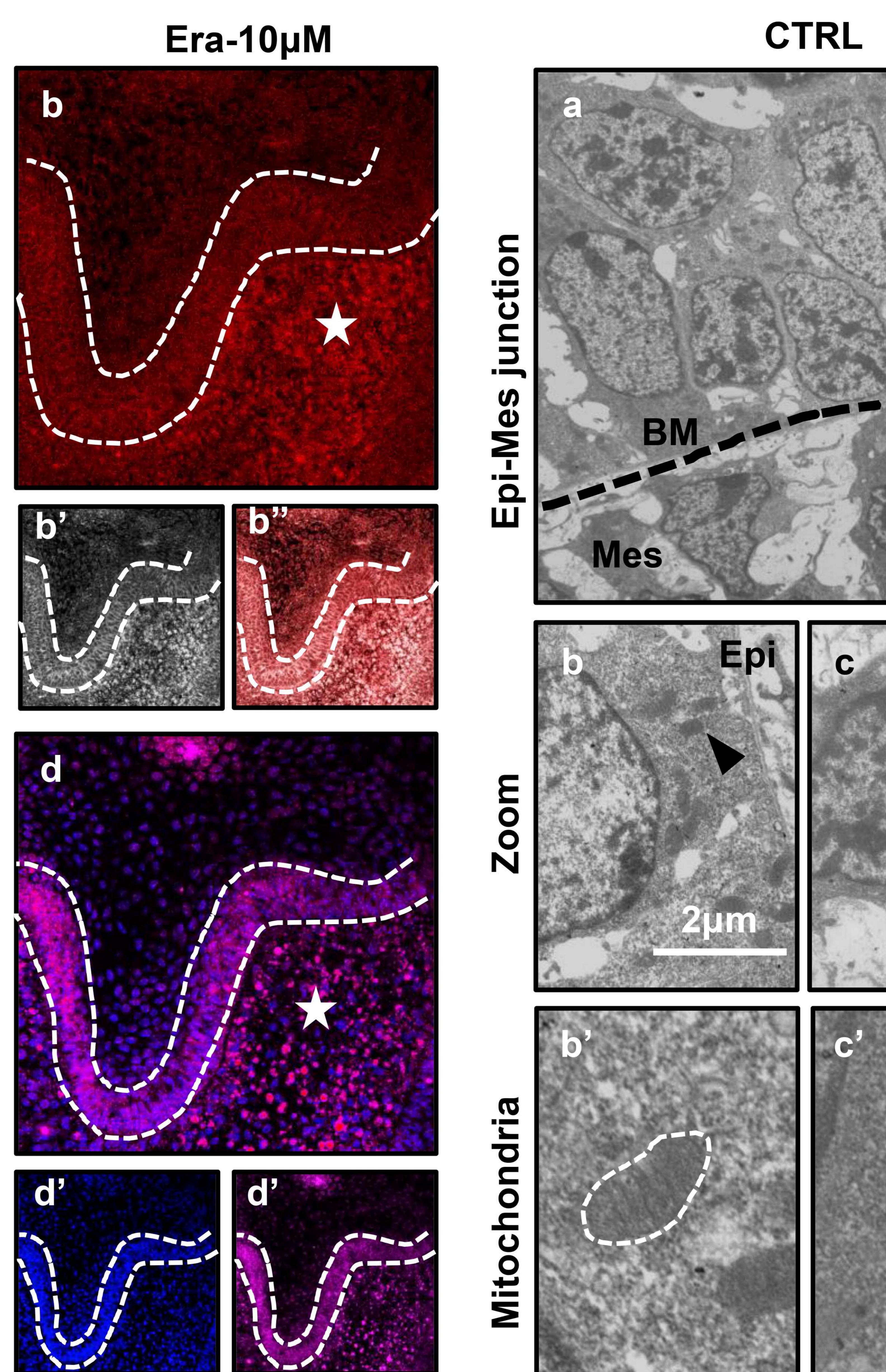
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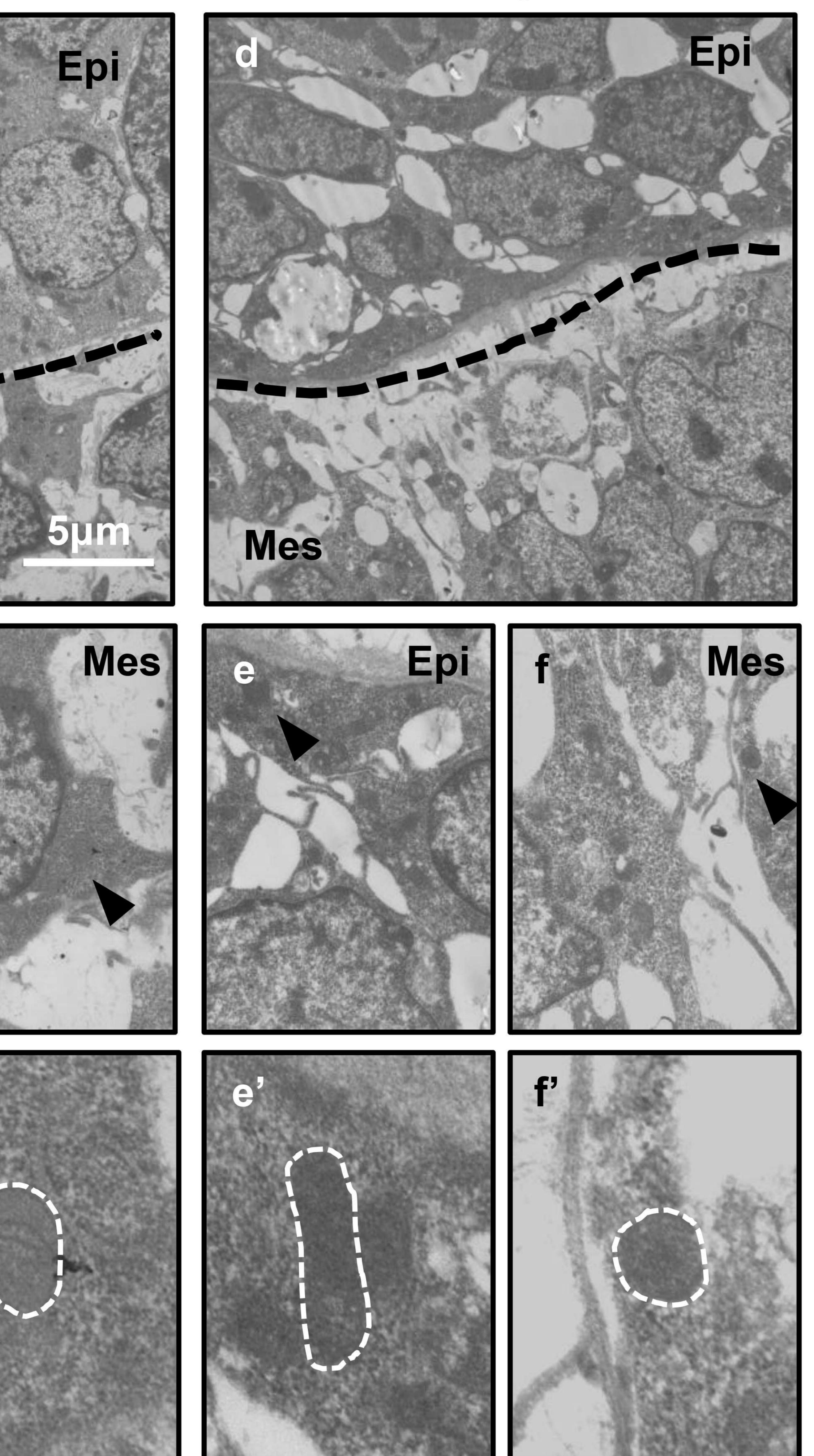


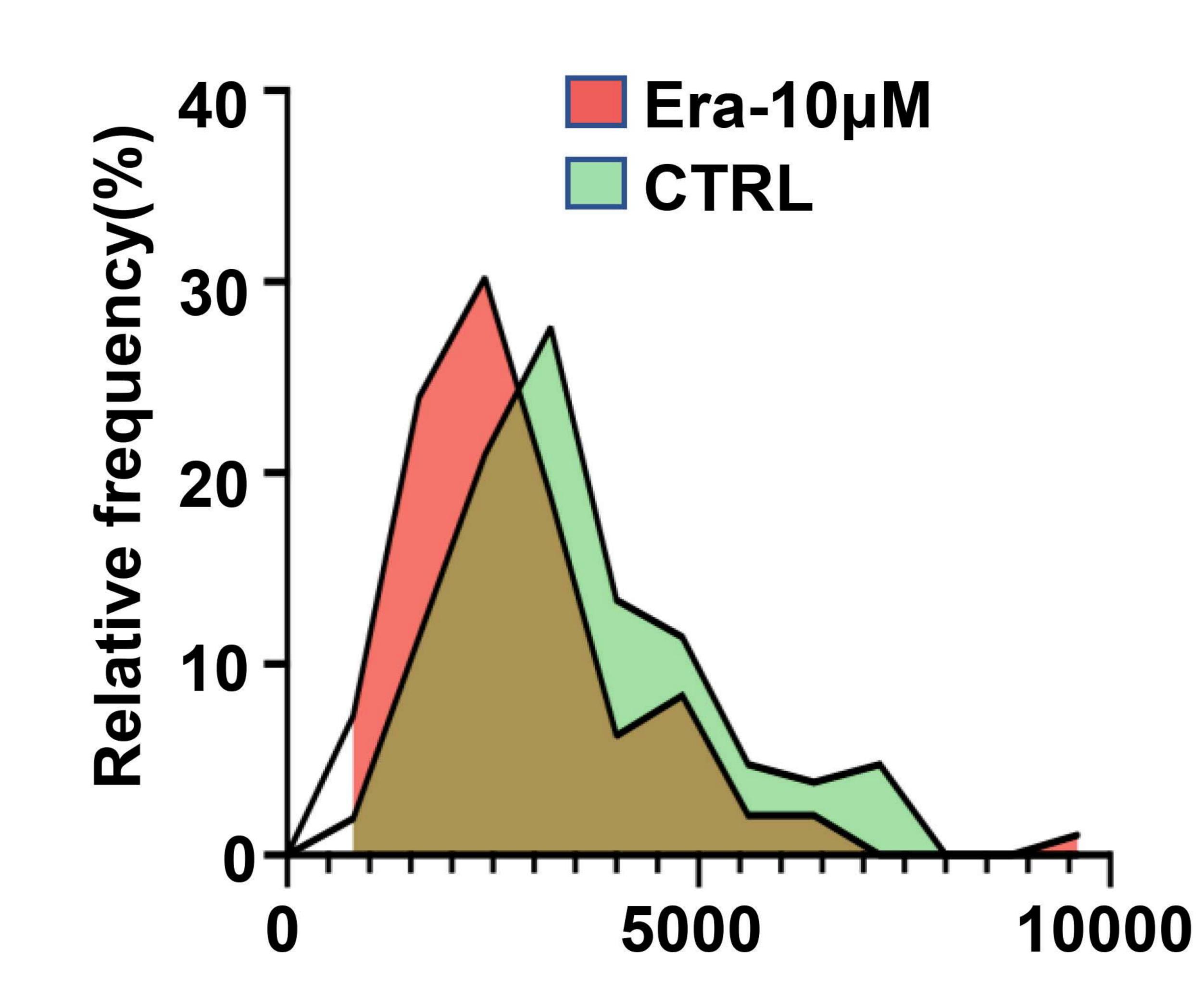
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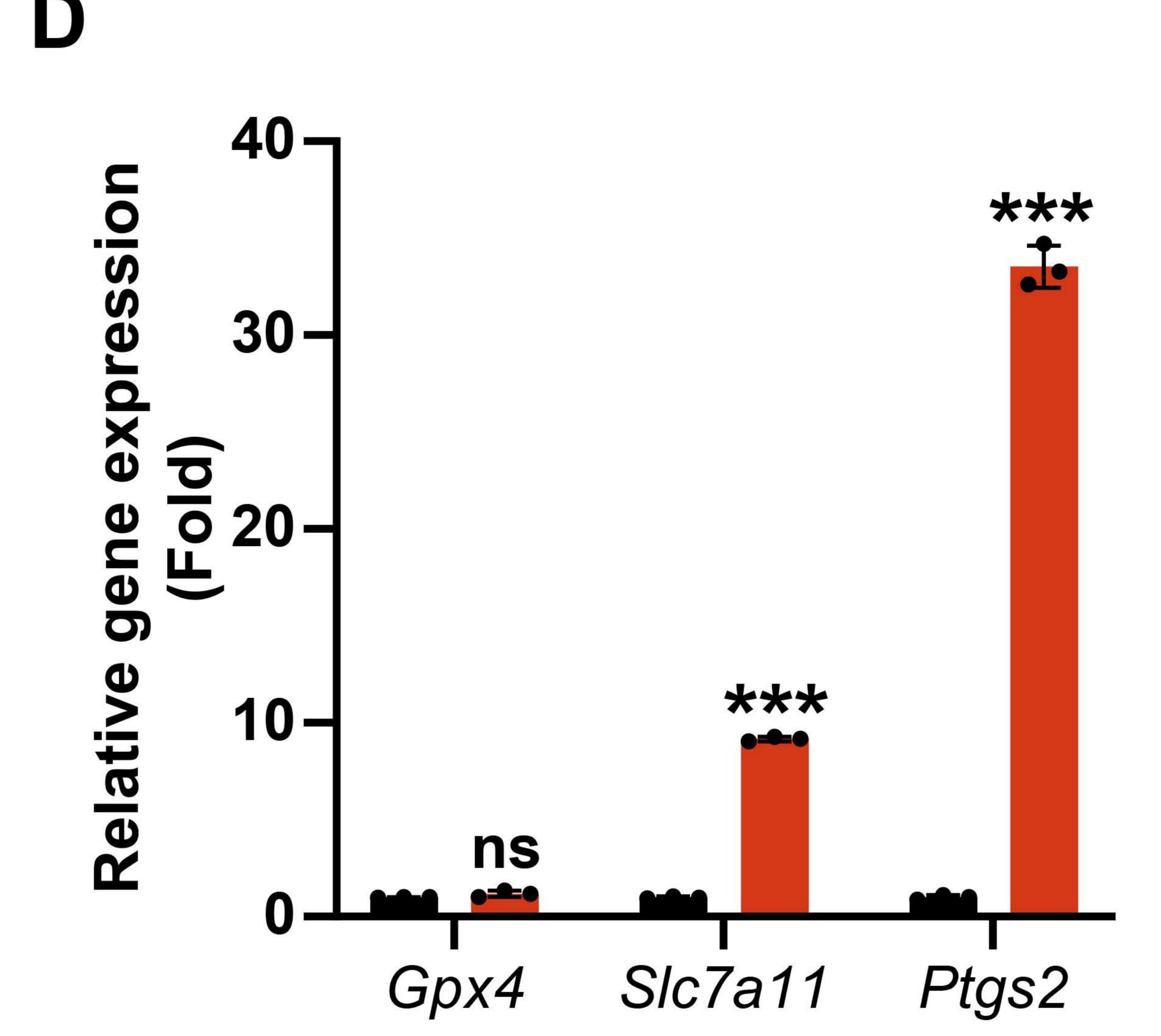


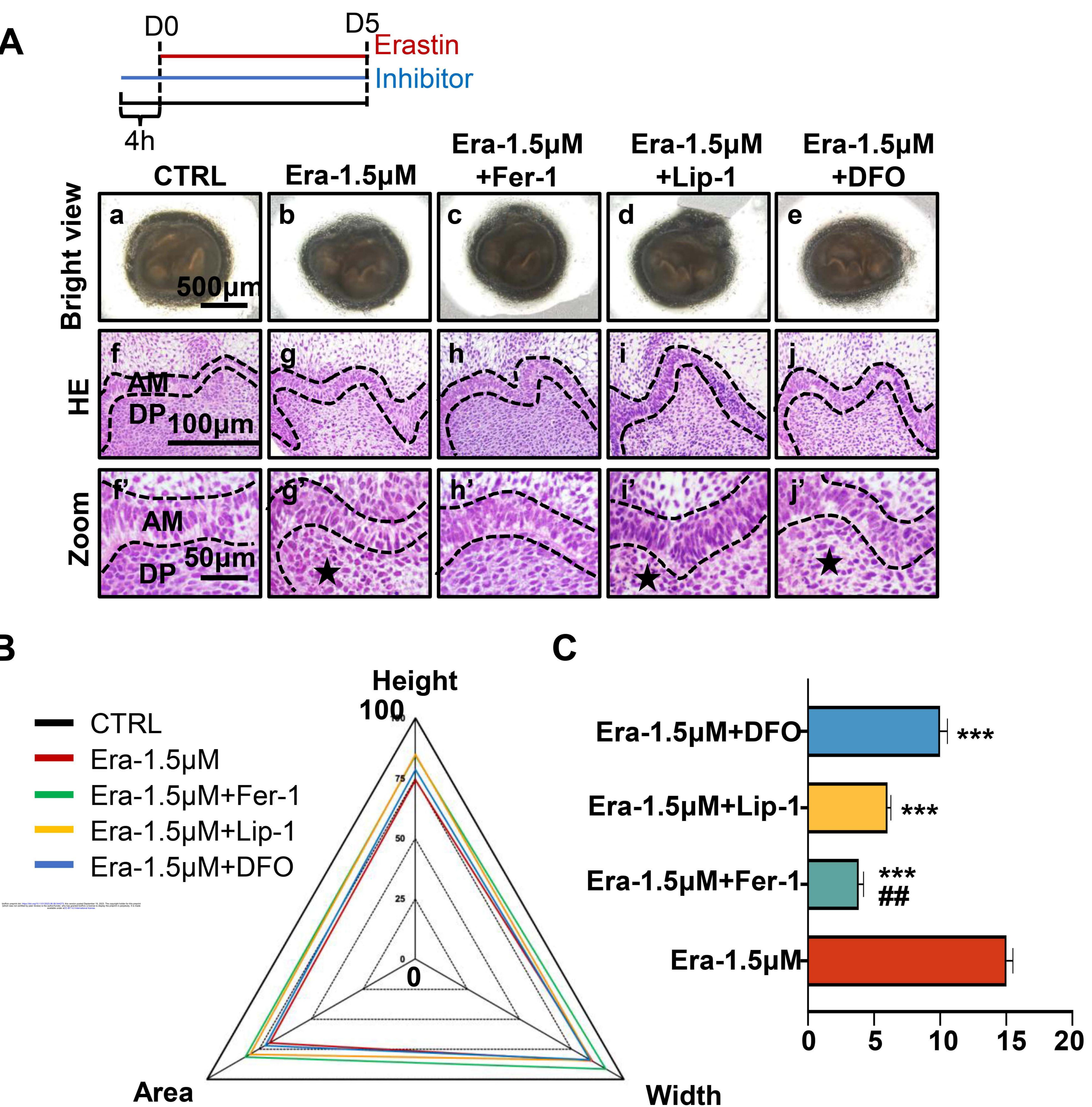


Era-10µM



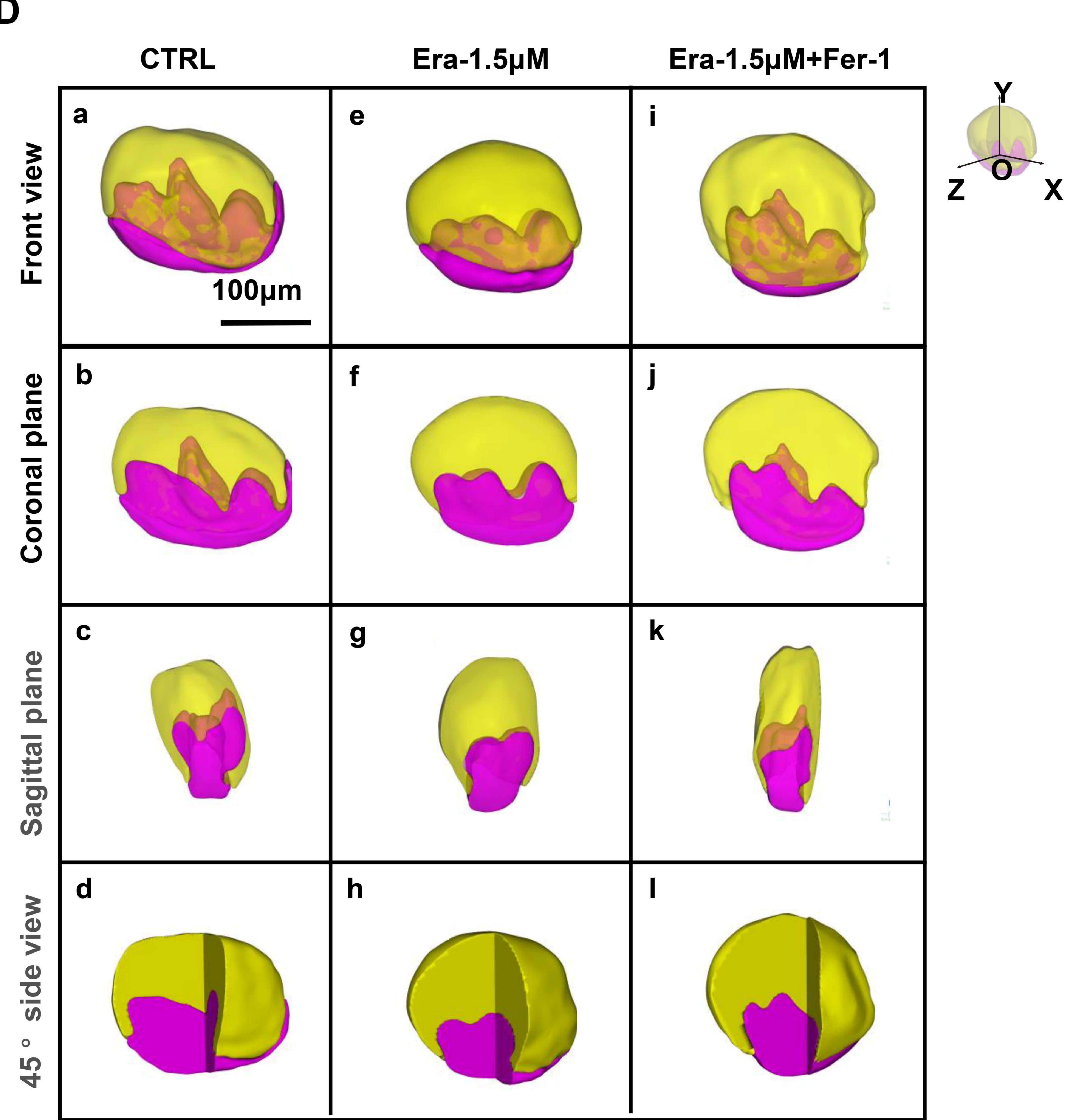


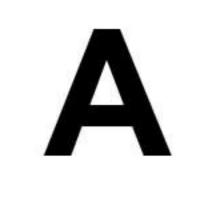




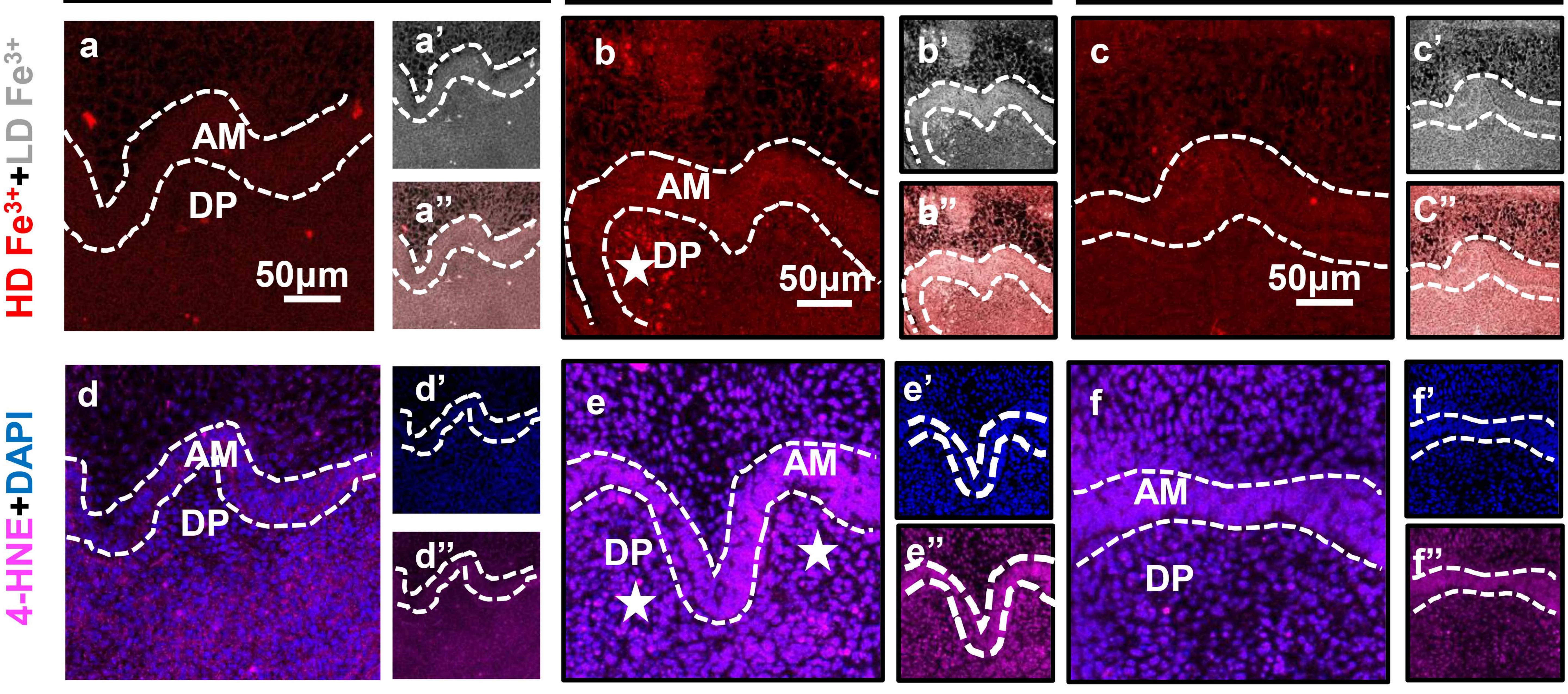
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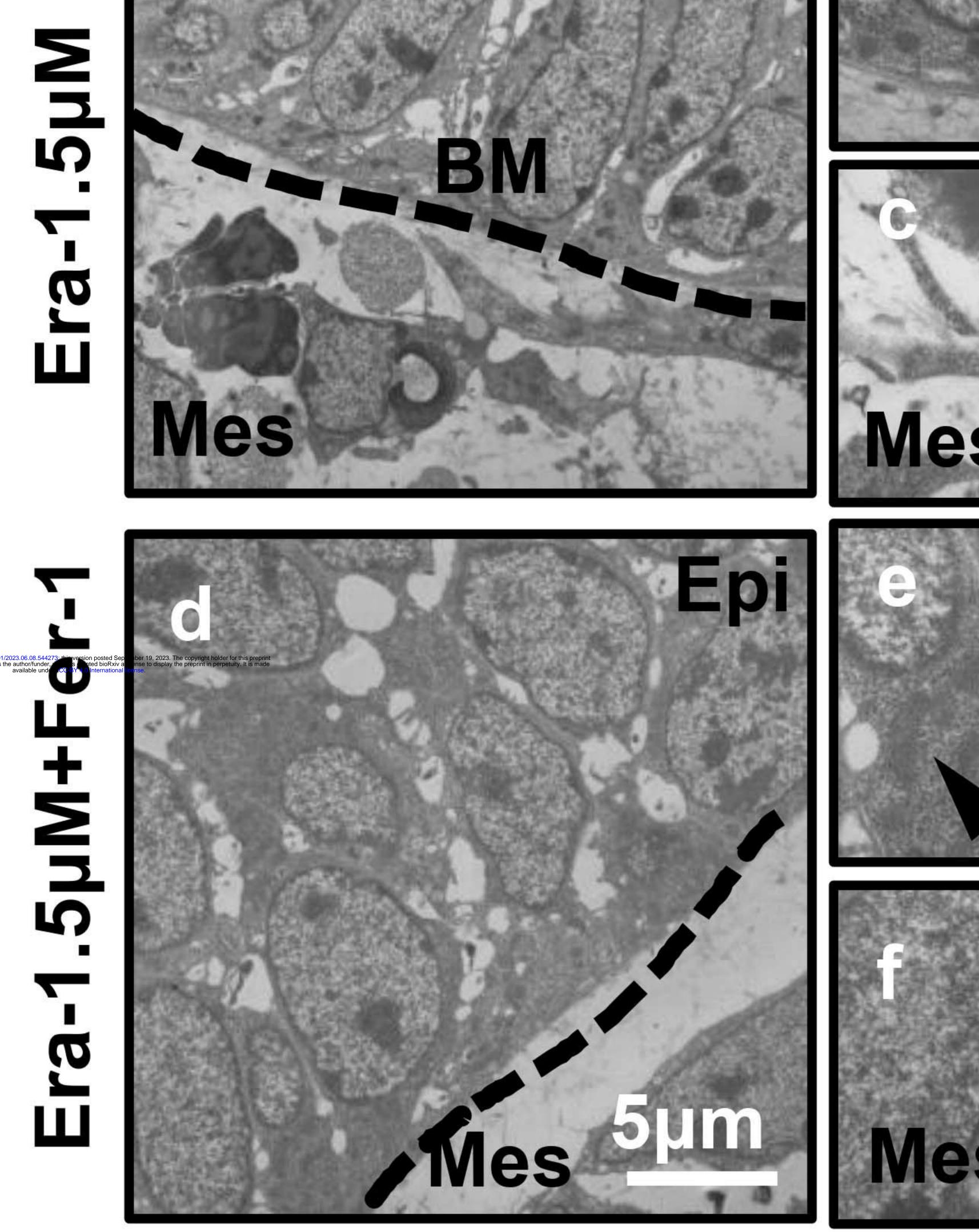
Epi-Mes junction

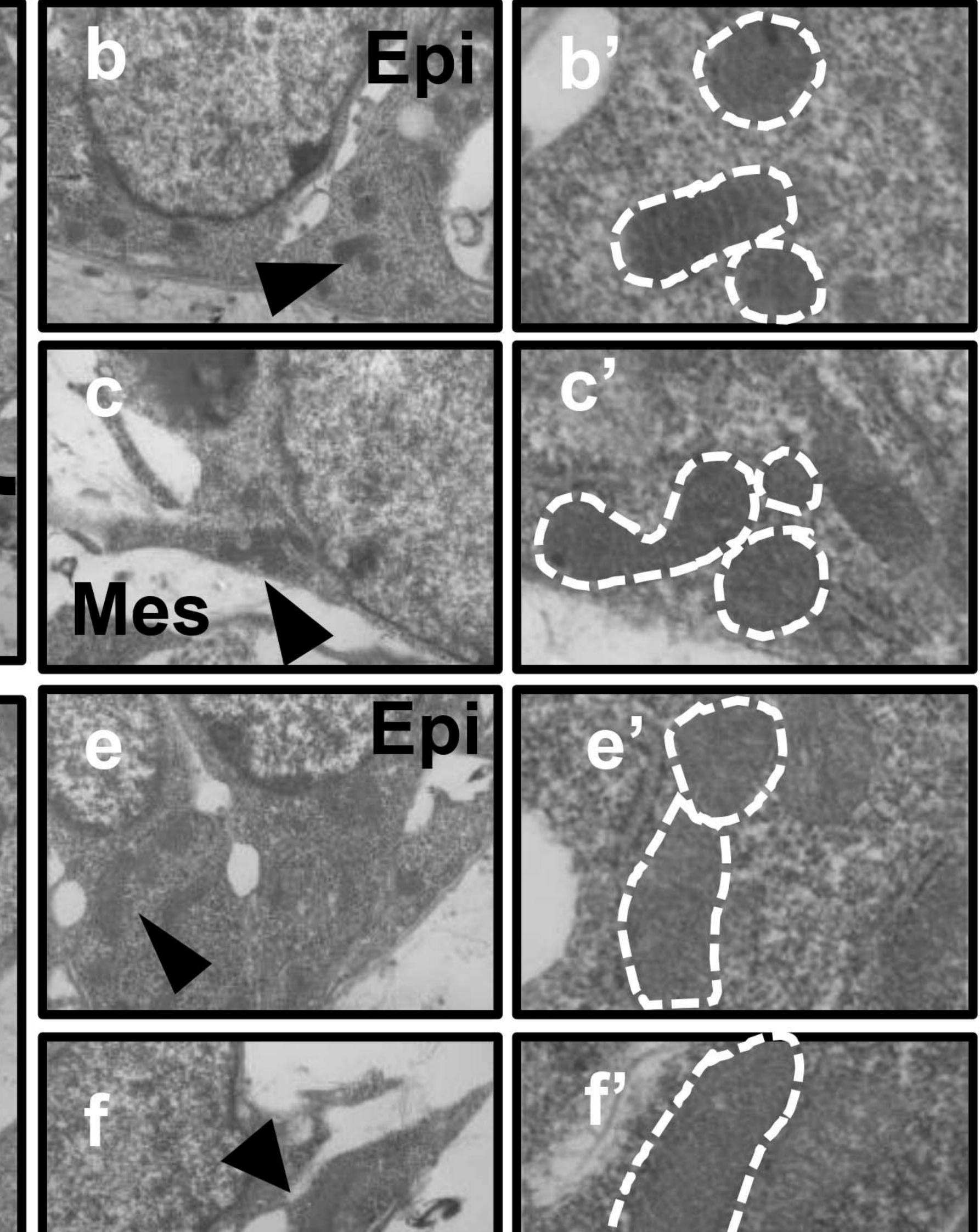
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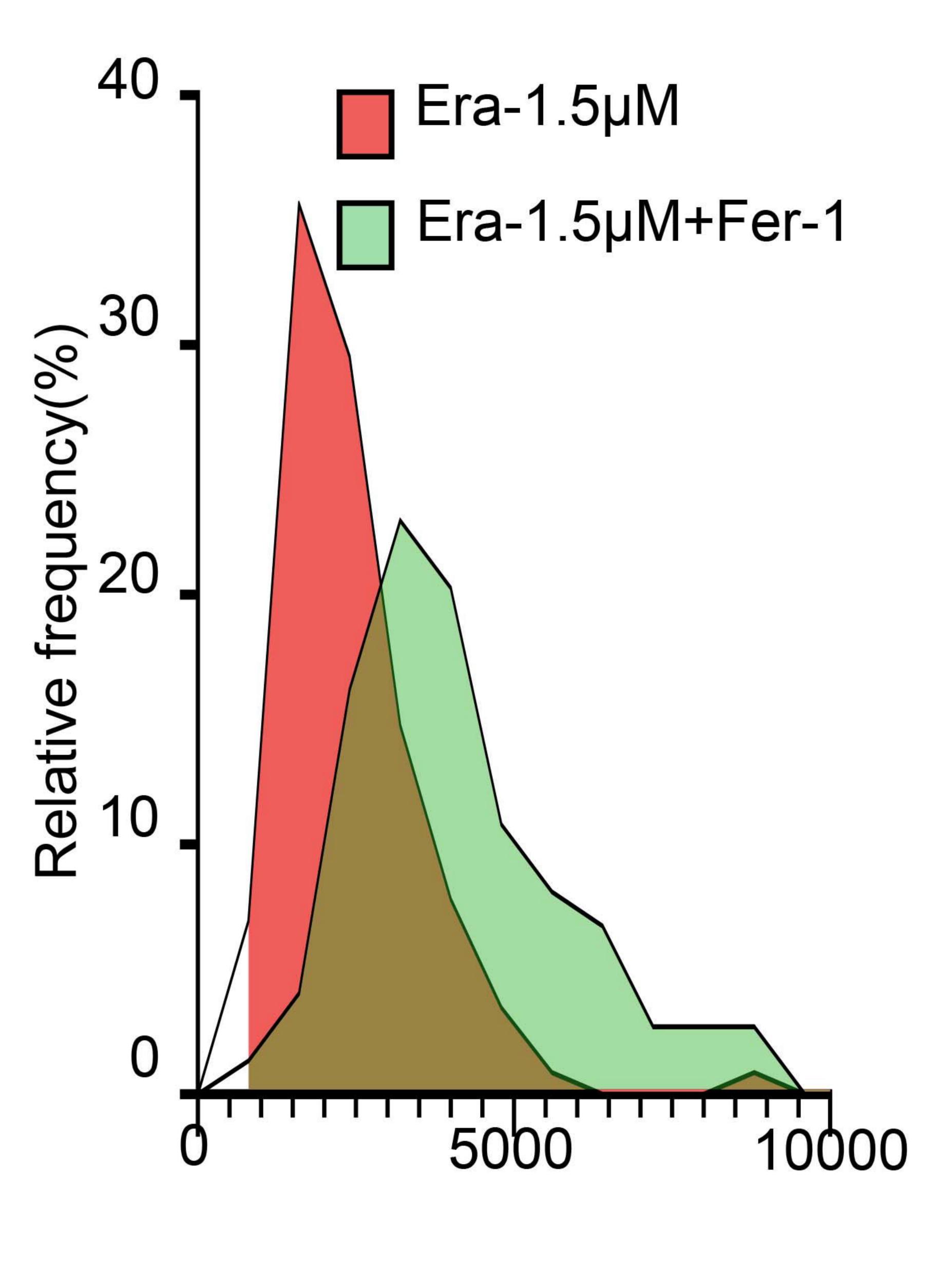




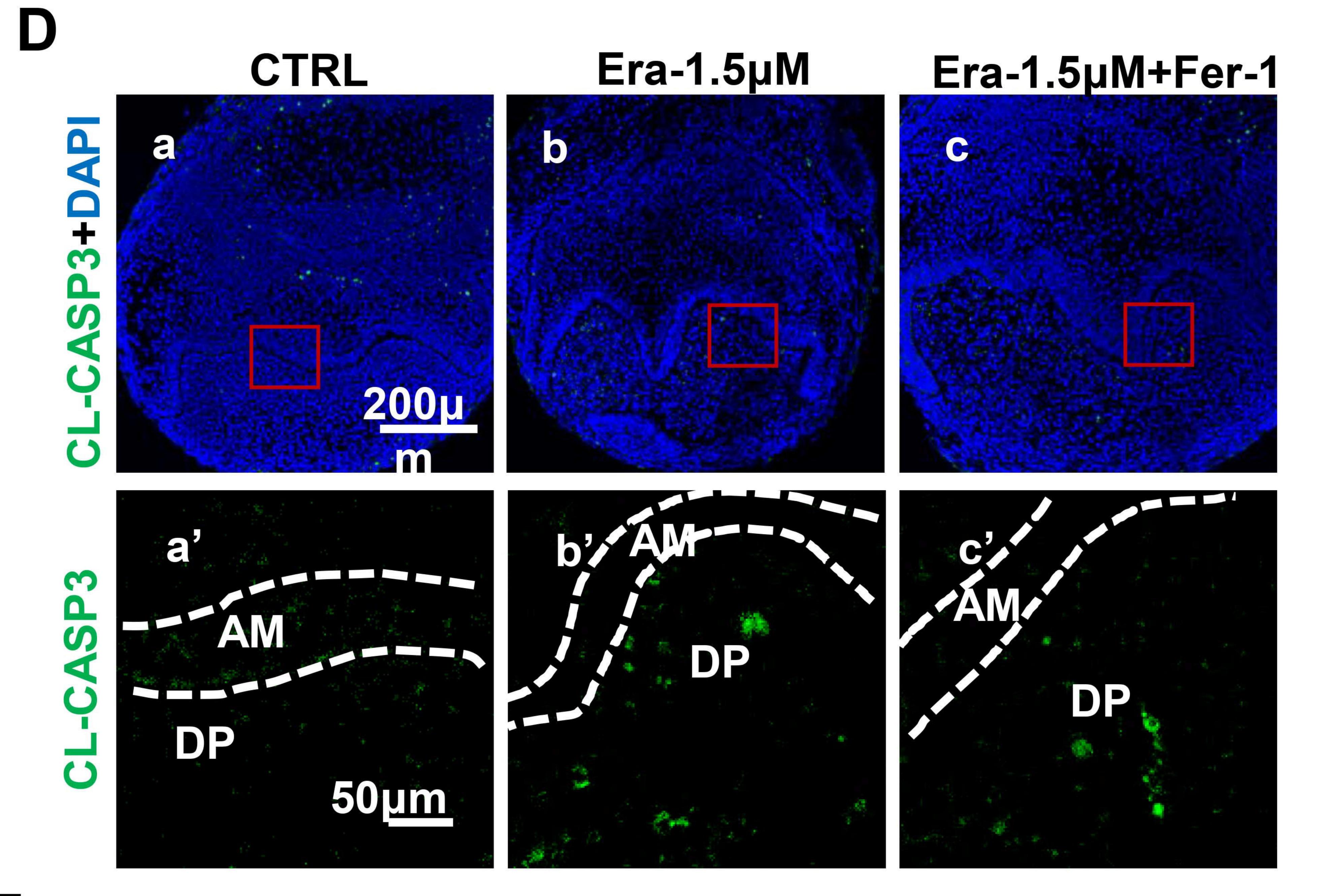
Era-1.5µM

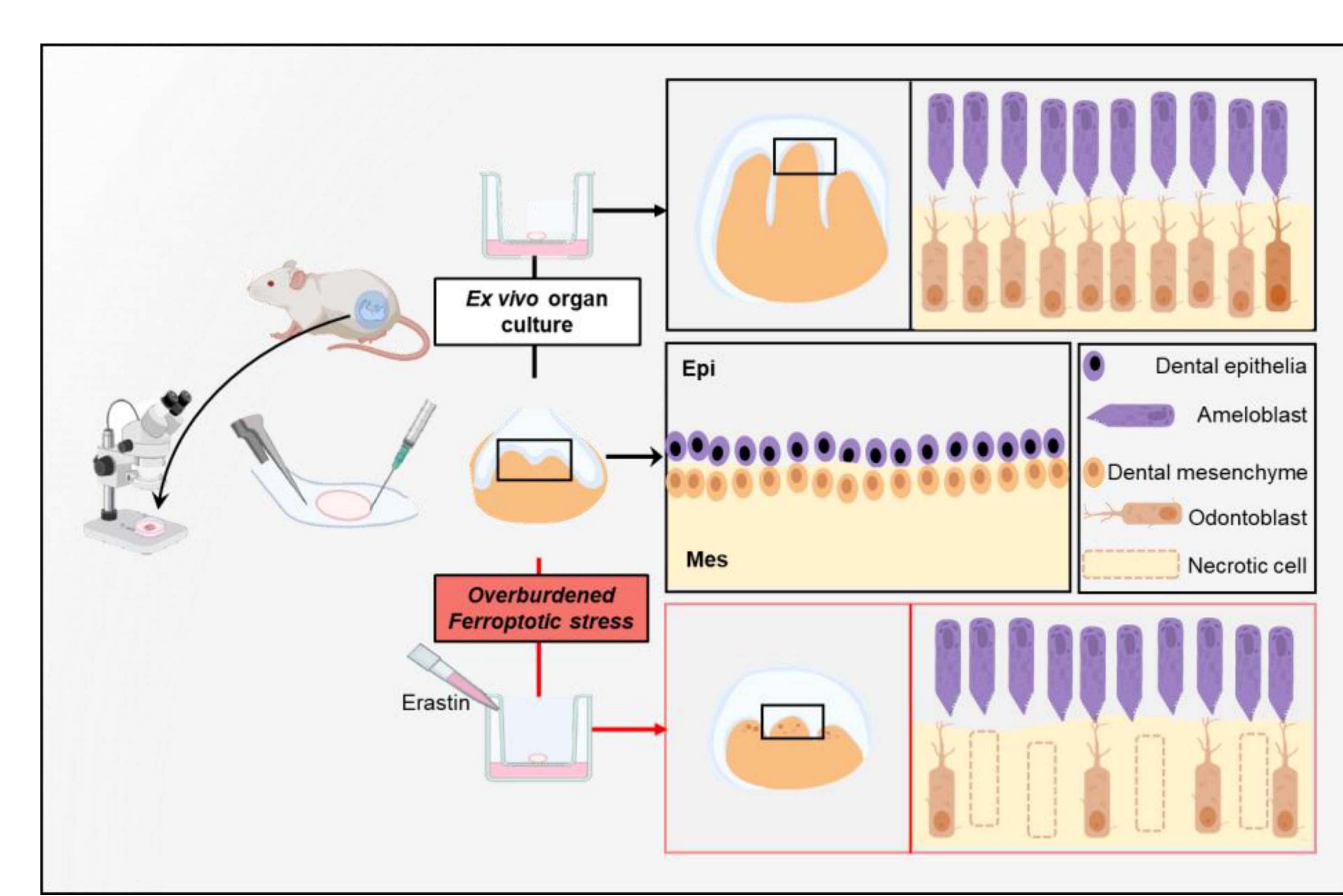
Era-1.5µM+Fer-1

Mitochondria





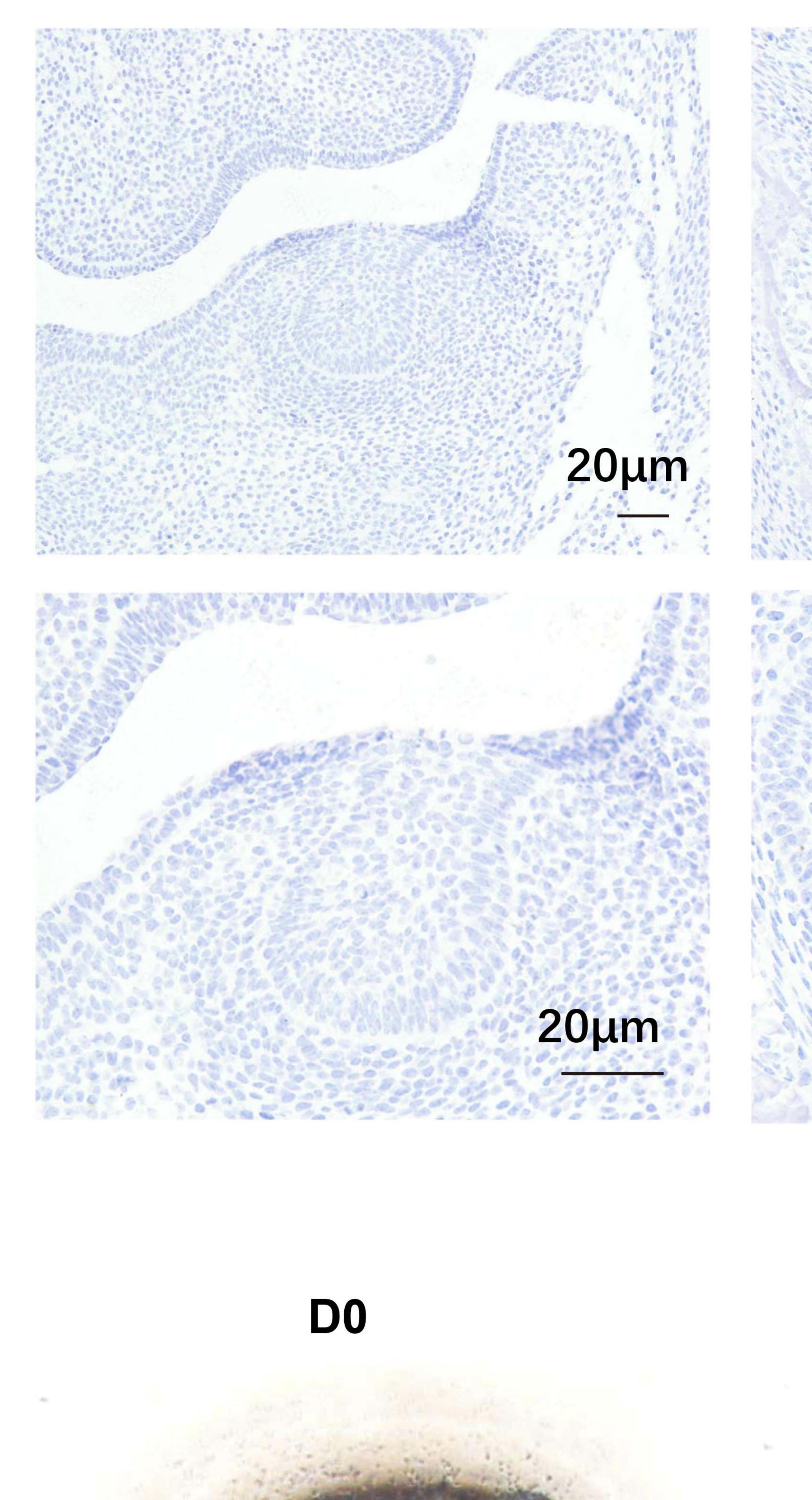




E13.5

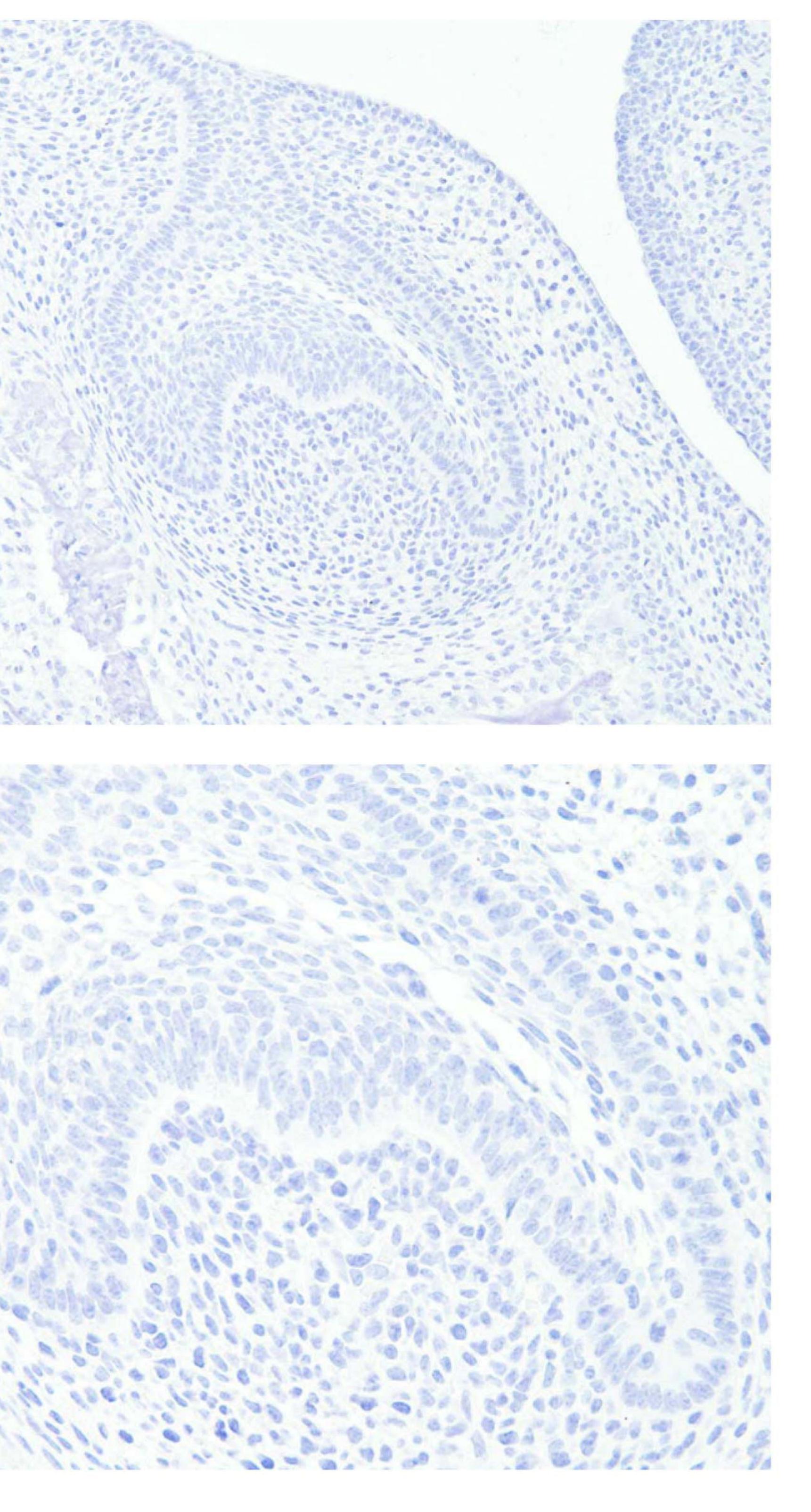
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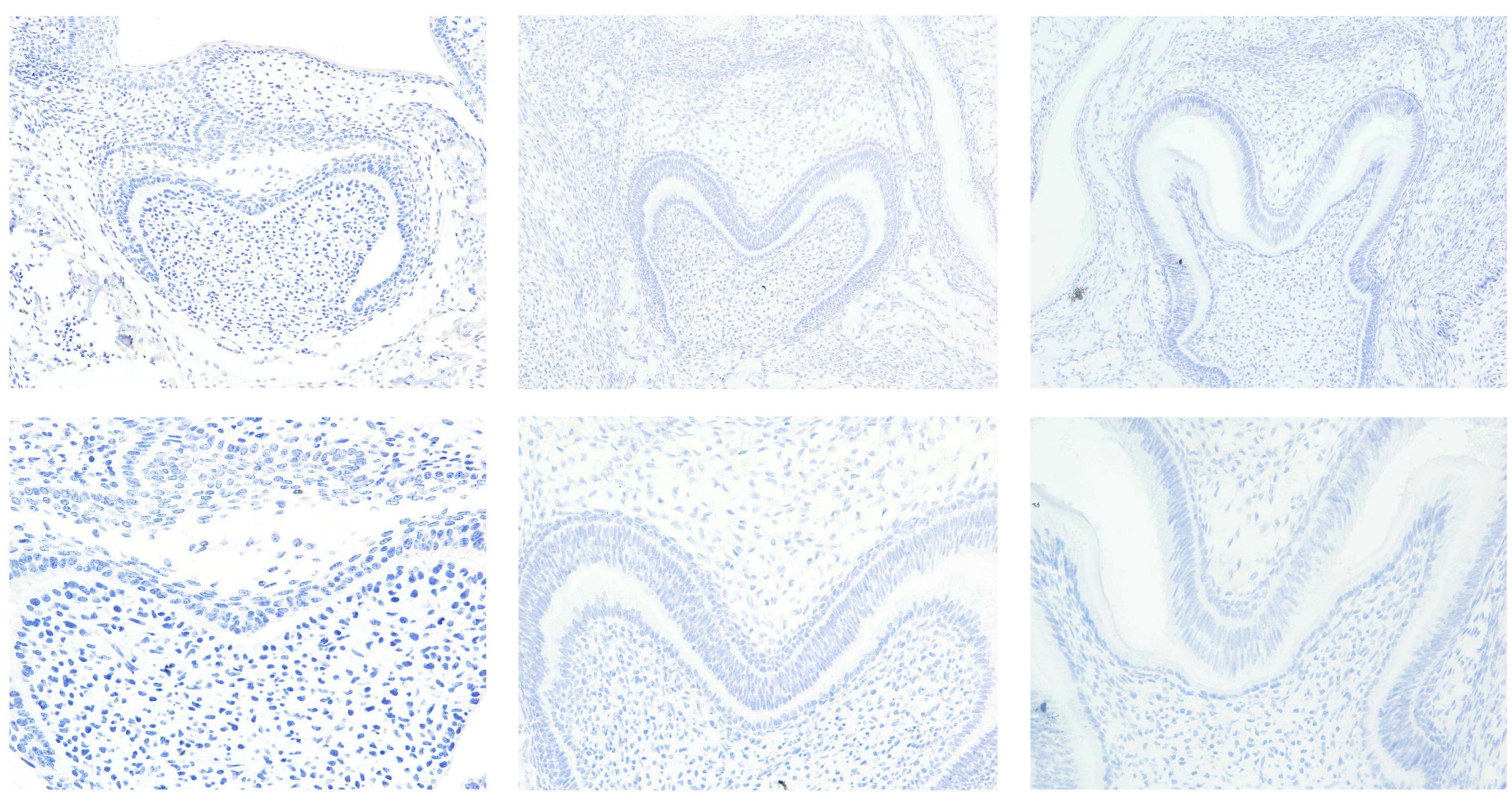
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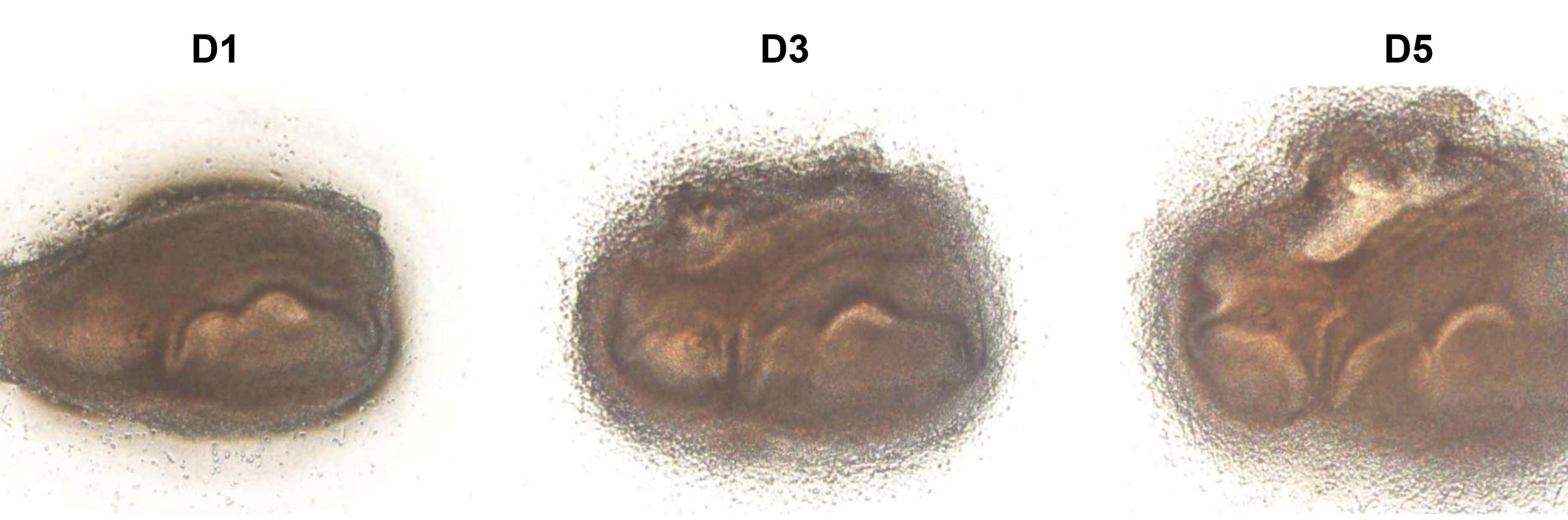






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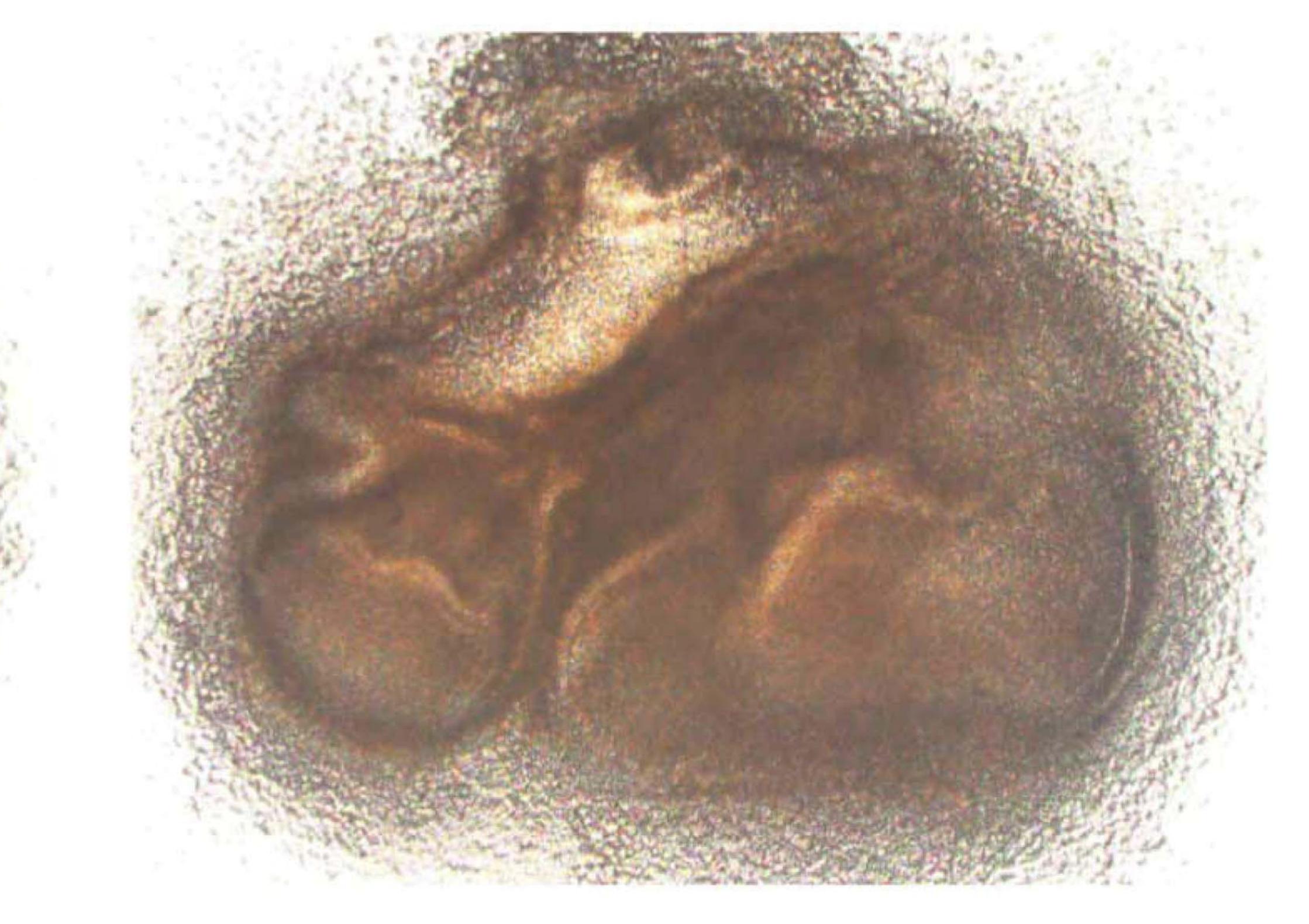


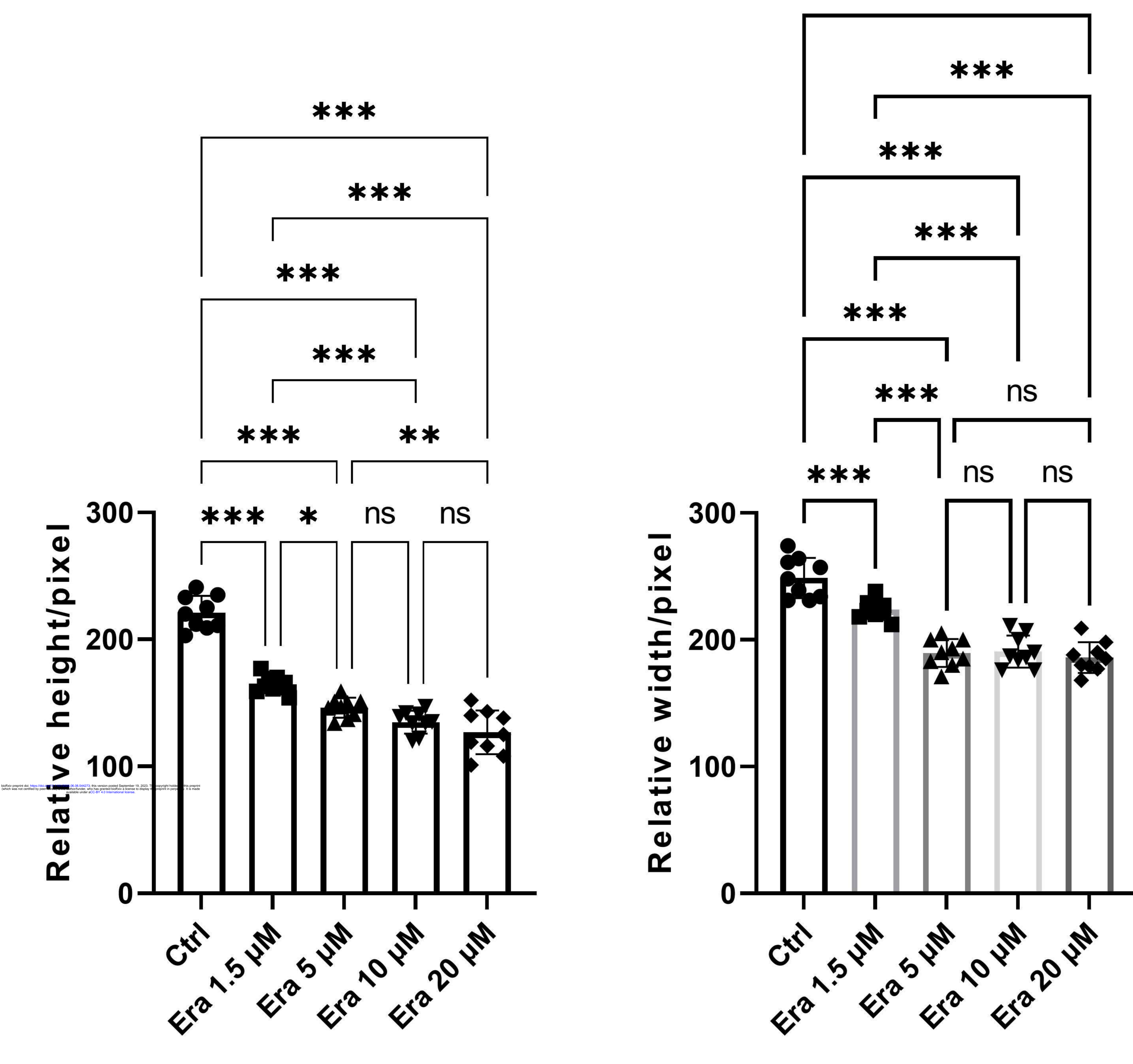


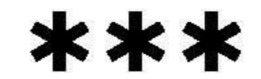
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P3



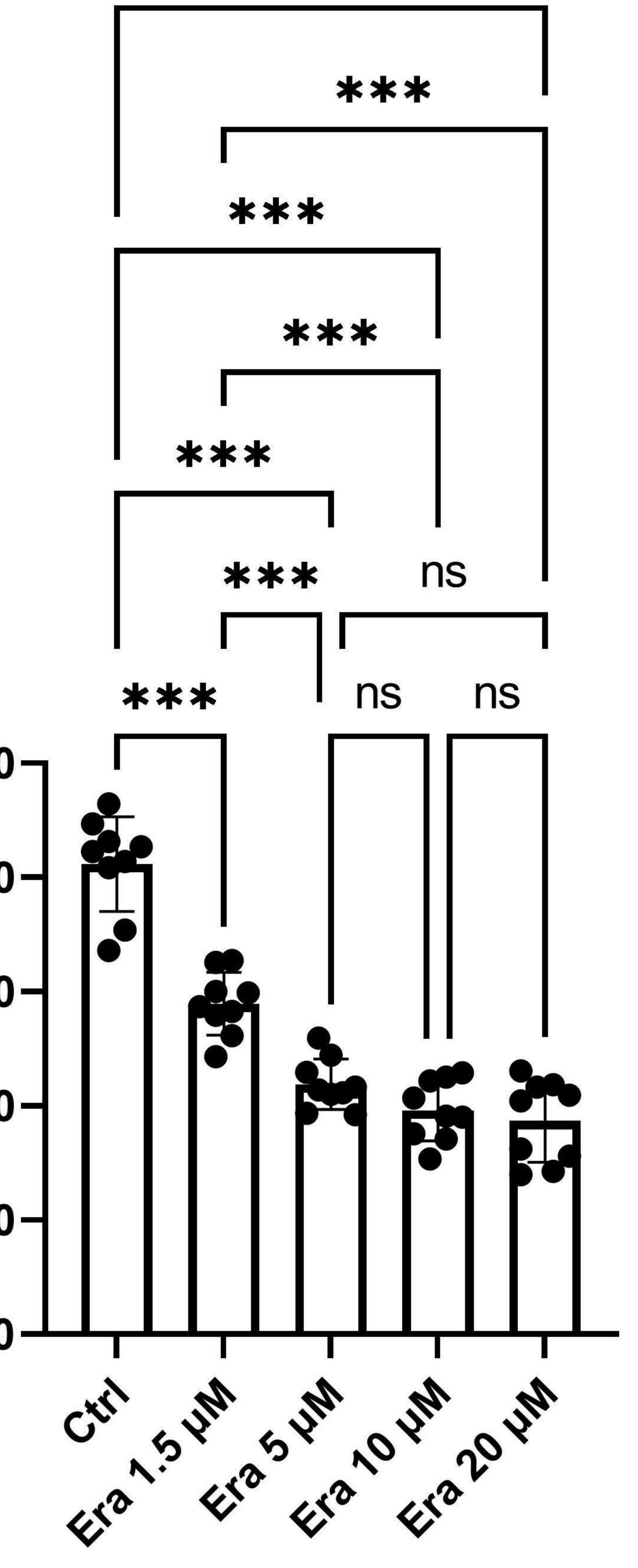




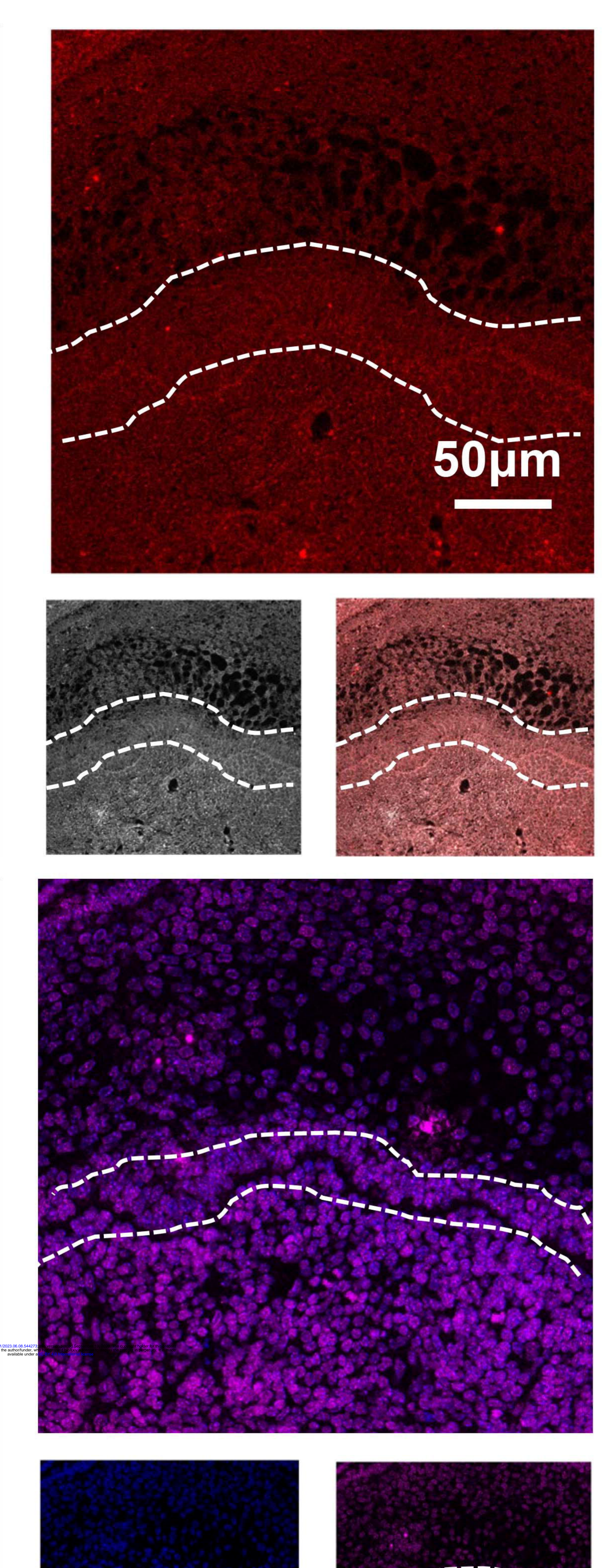


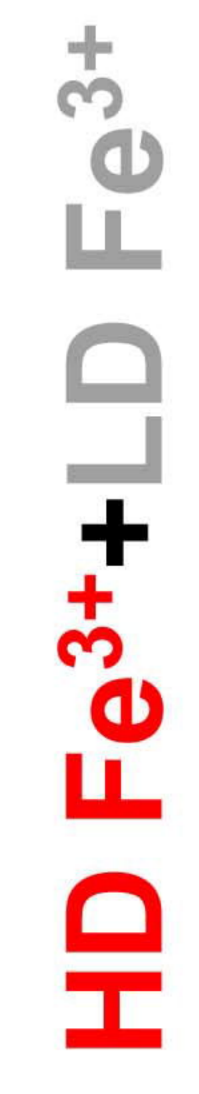
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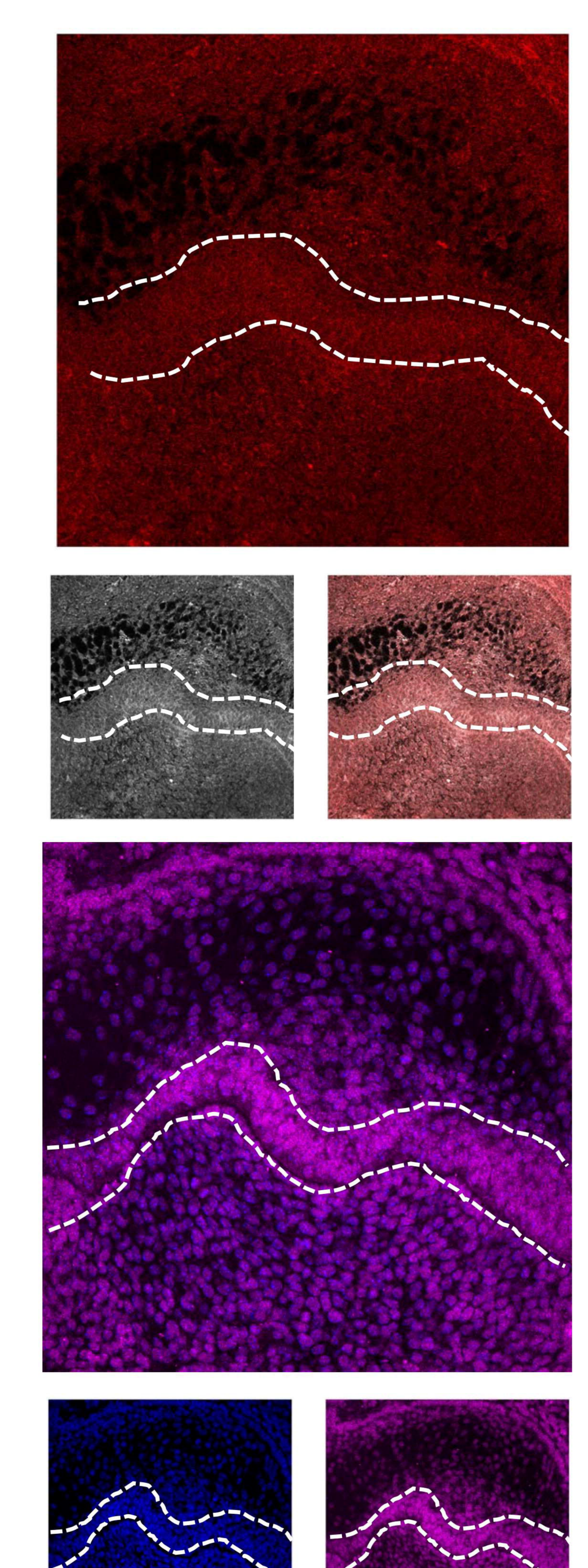


CTRL D1

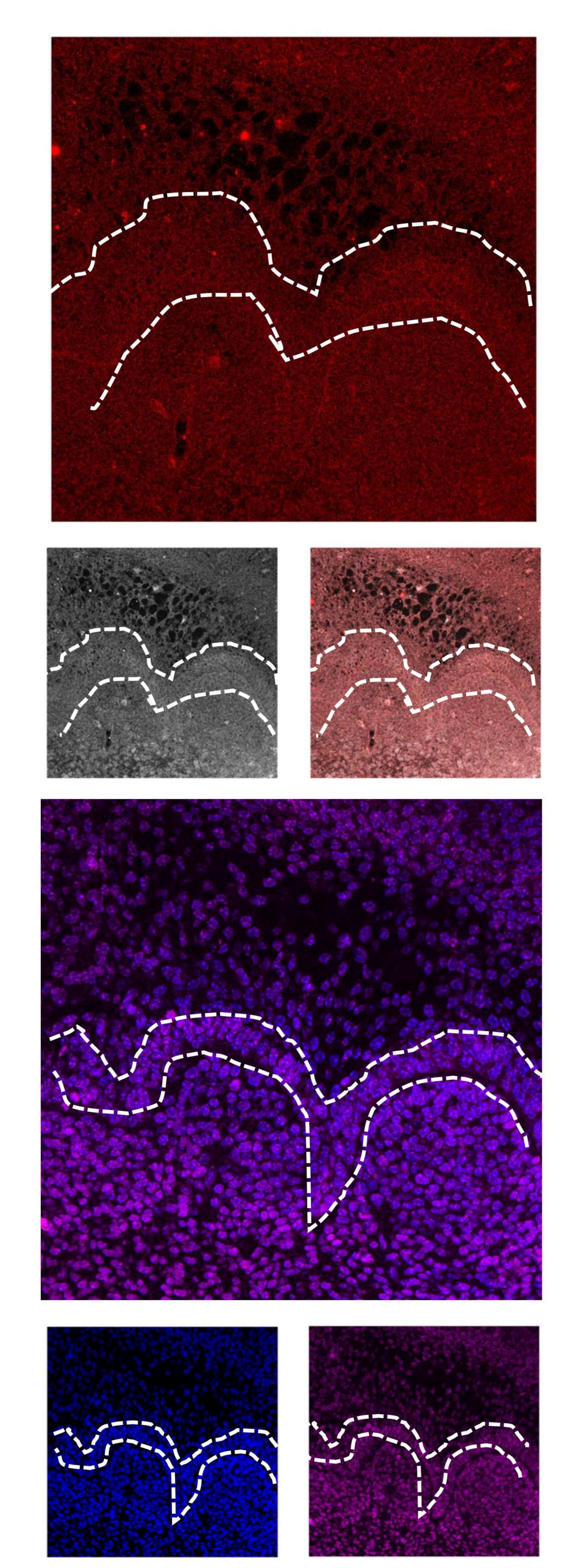




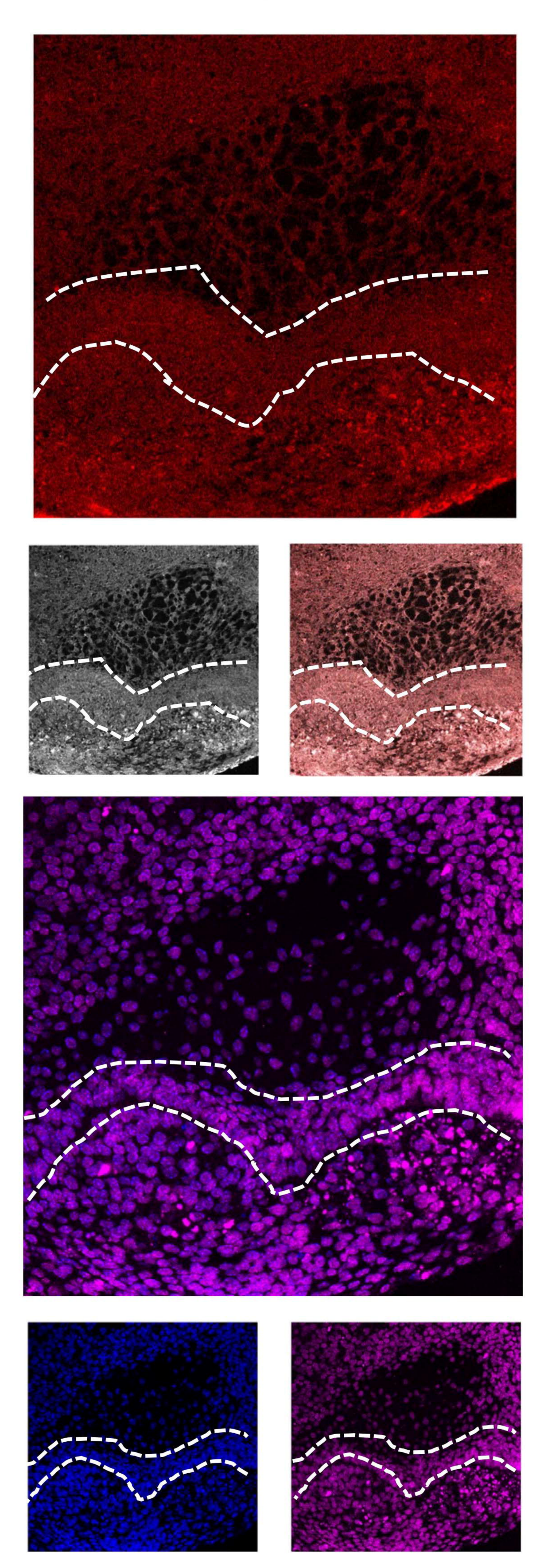
Era-10µM D1



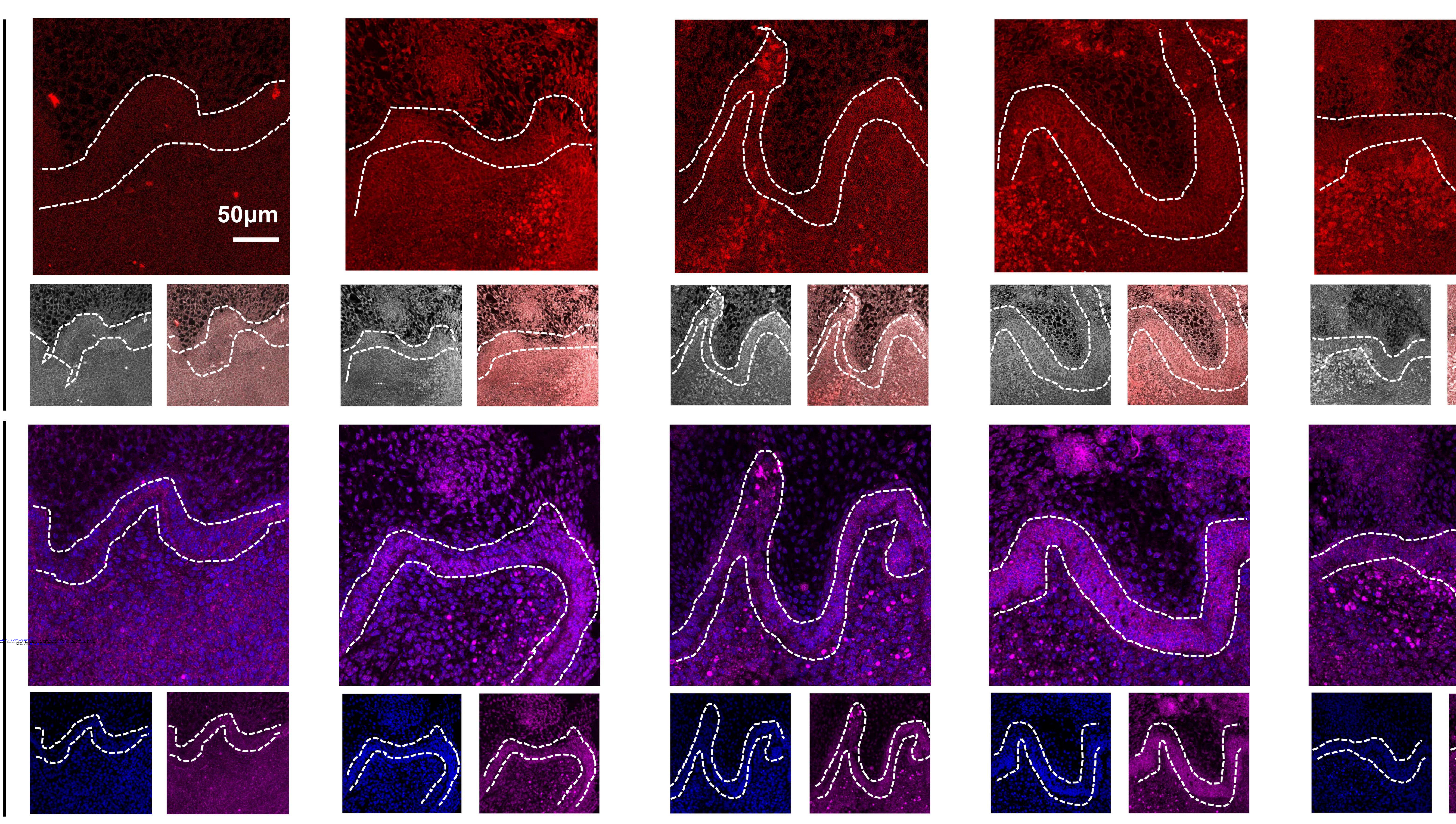
CTRL D3



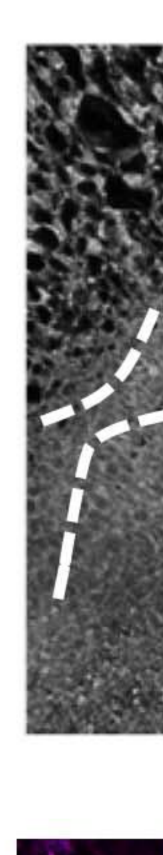
Era-10µM D3

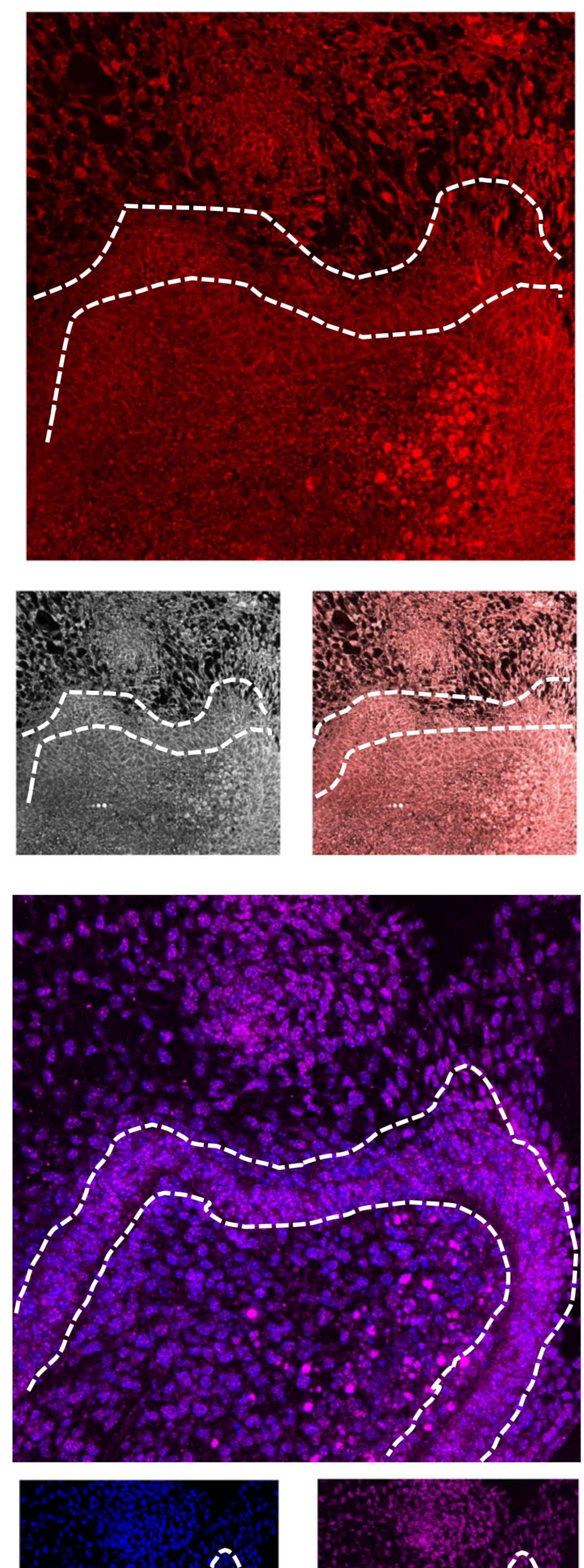


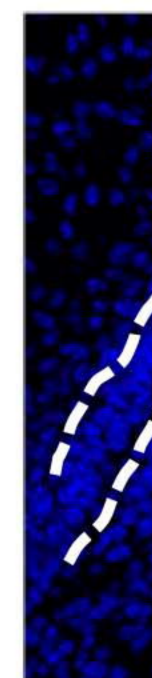
CTRL



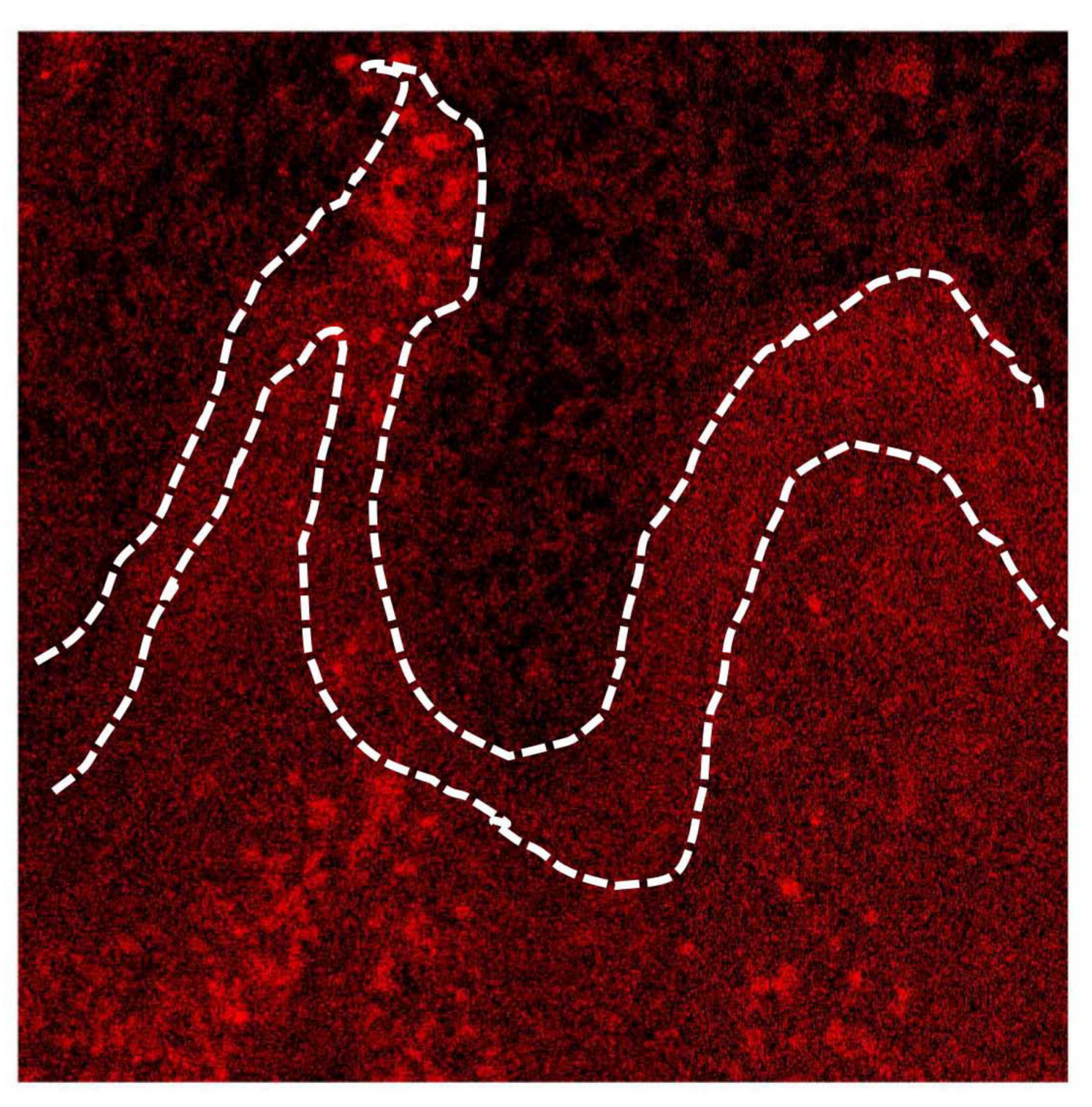


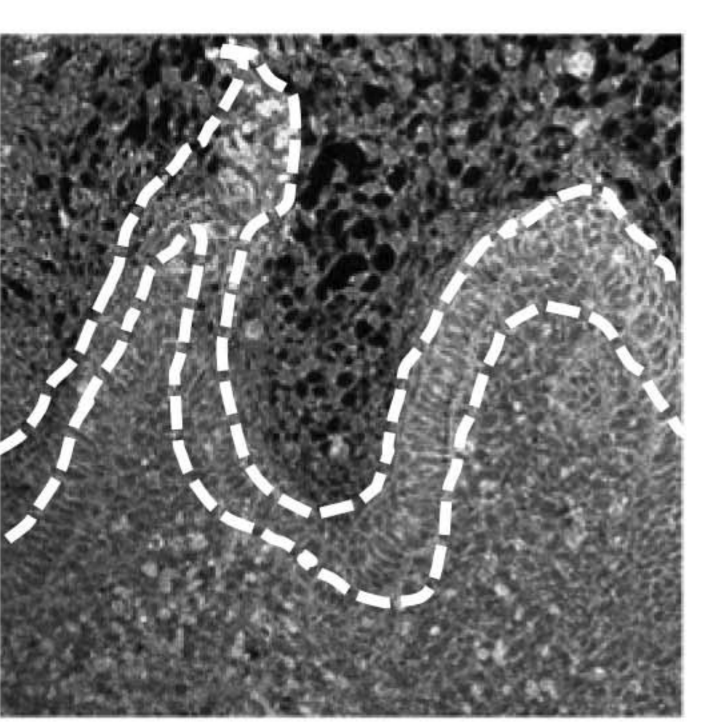


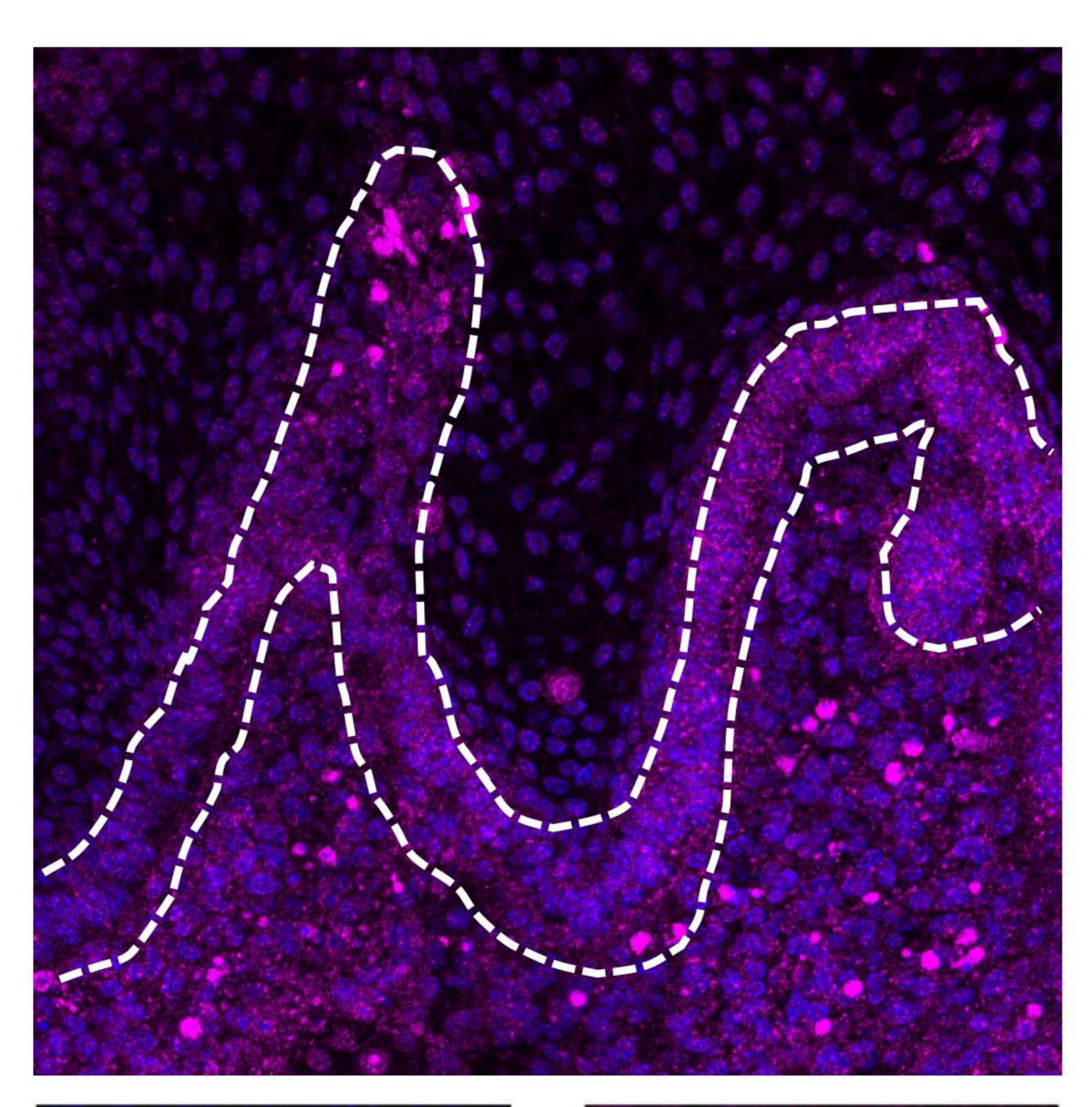


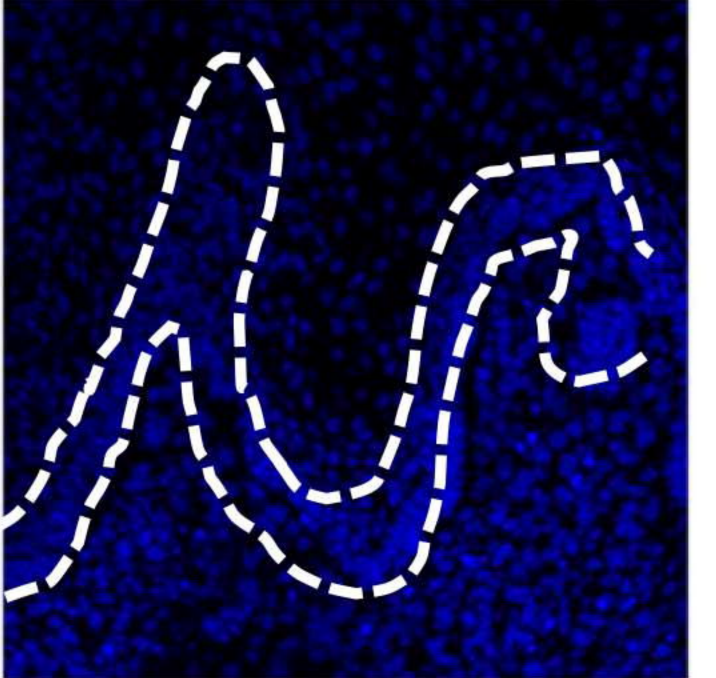


Era-1.5µM



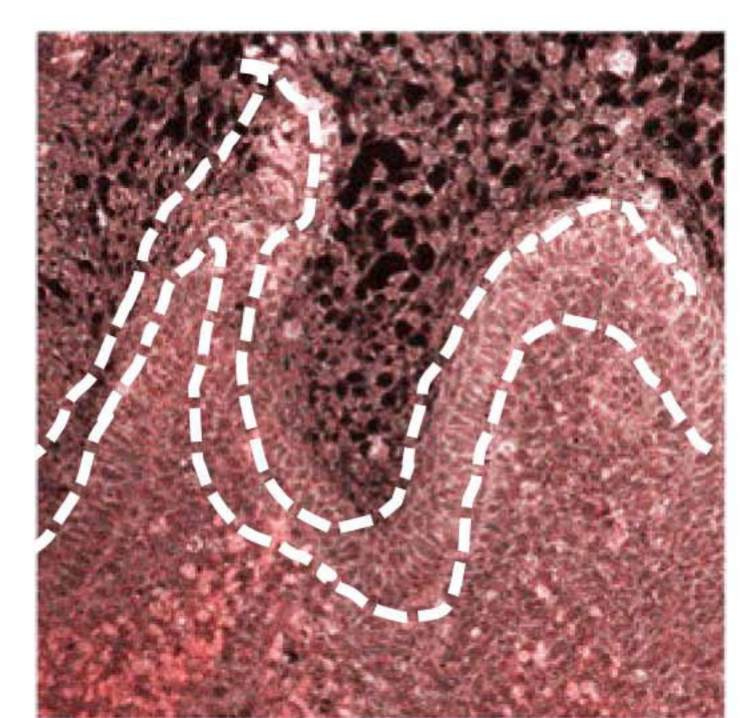


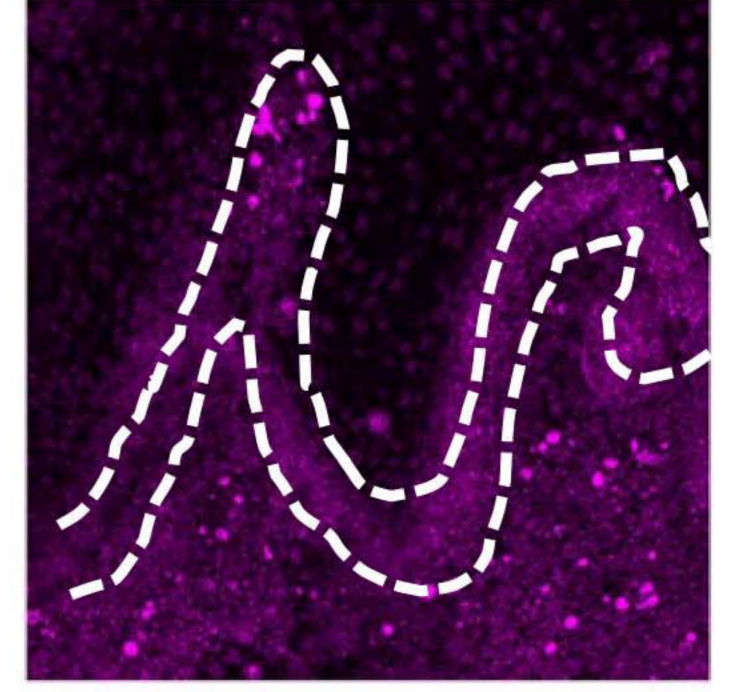


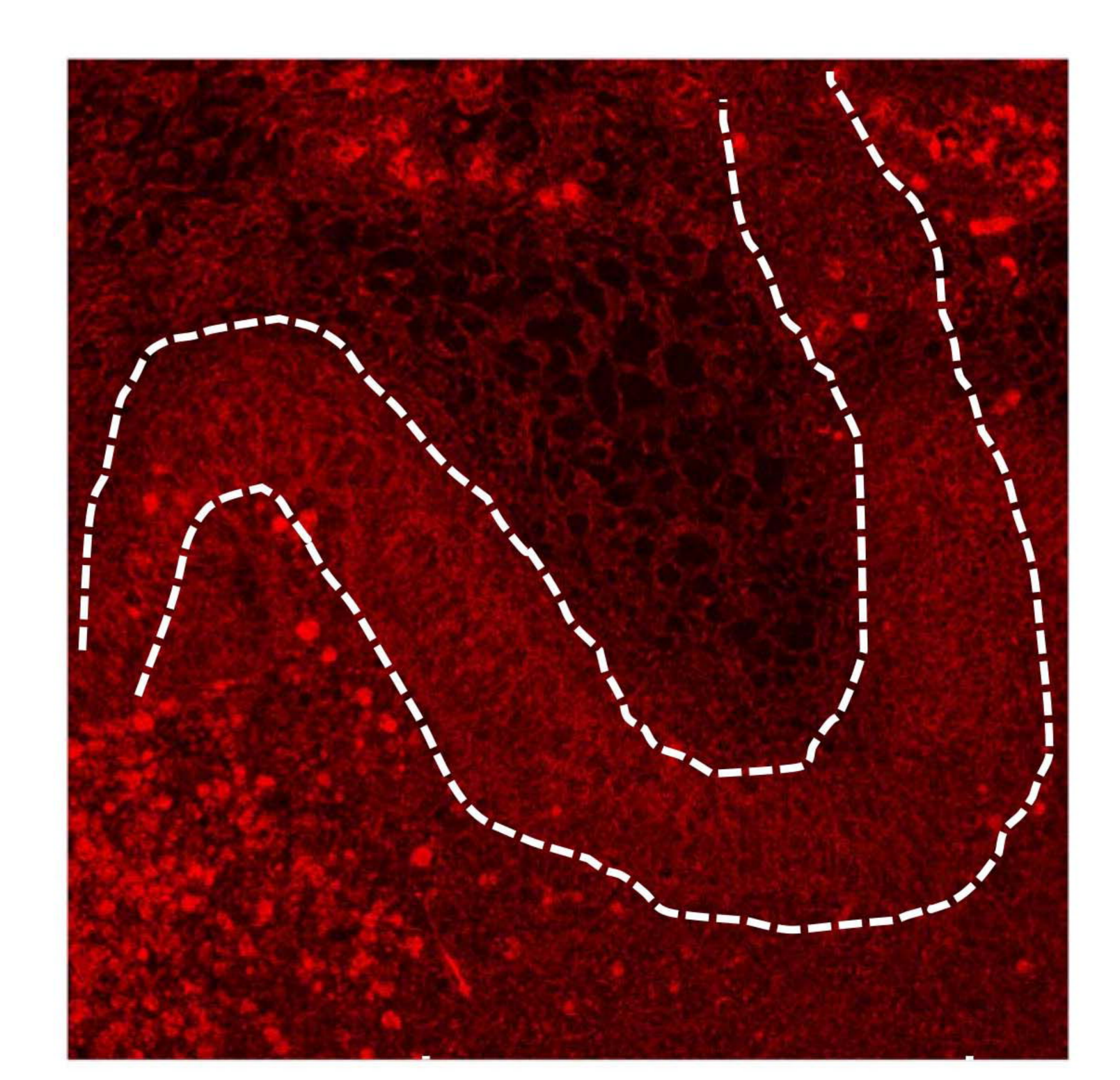


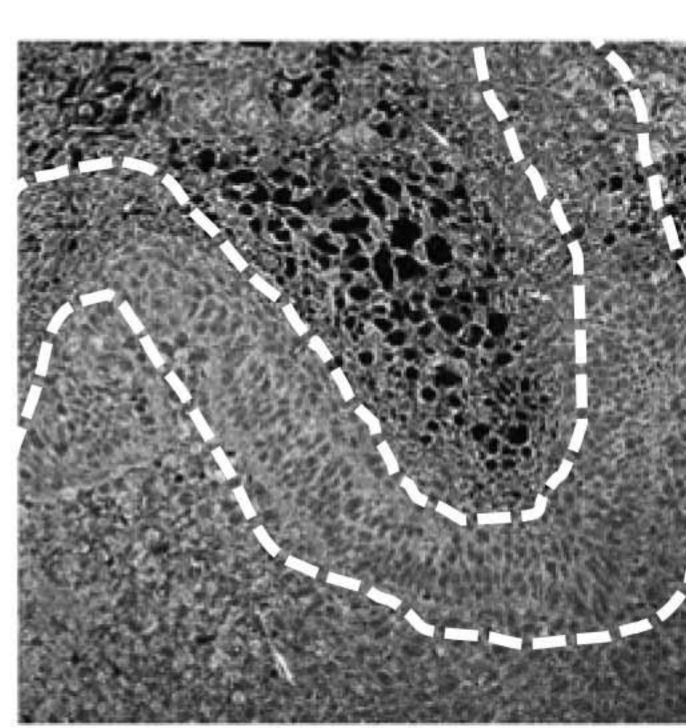
Era-5µM

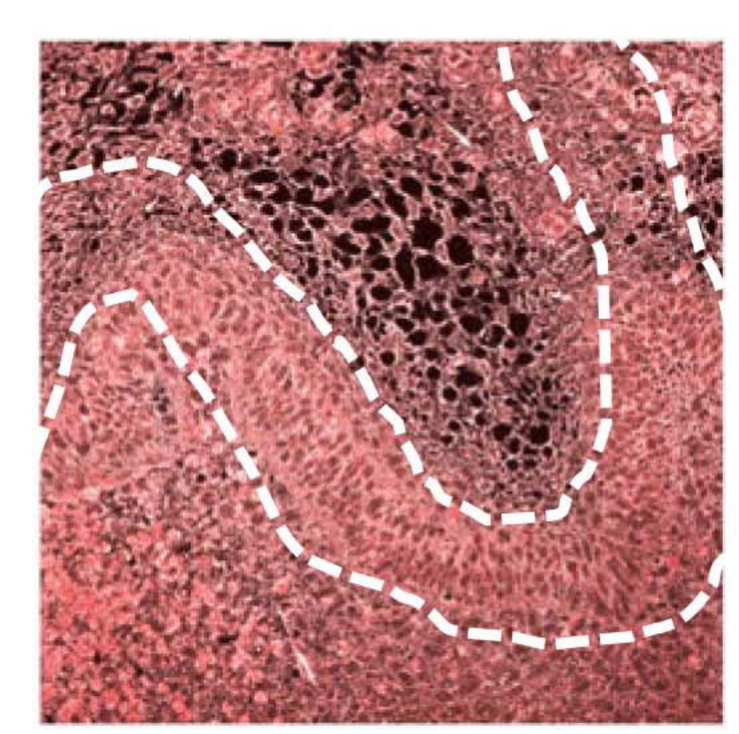
Era-10µM

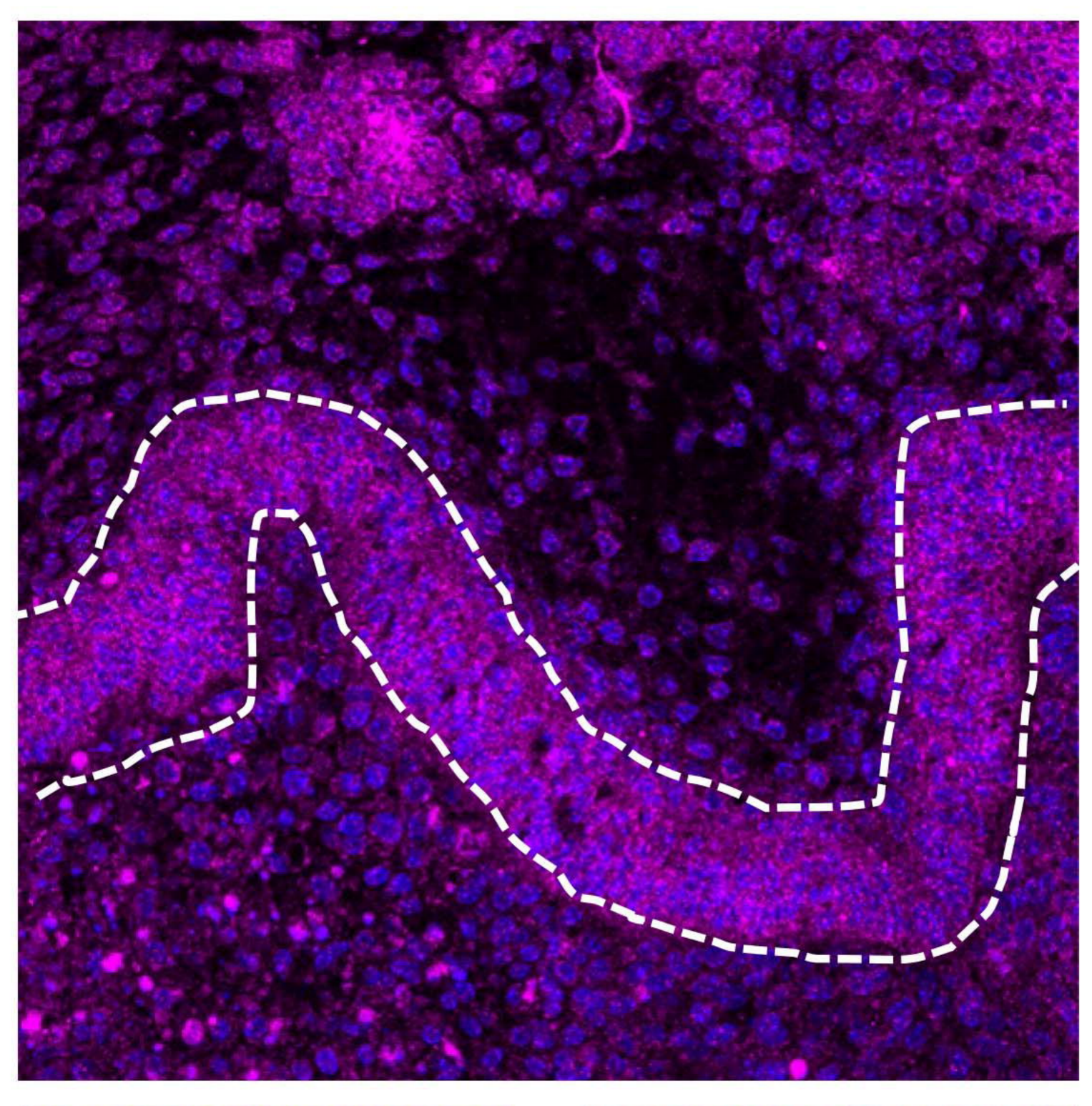


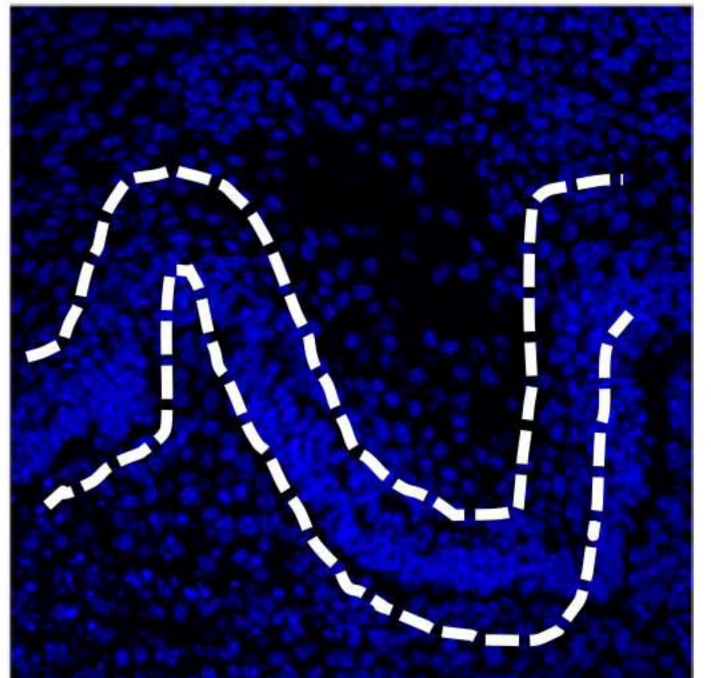


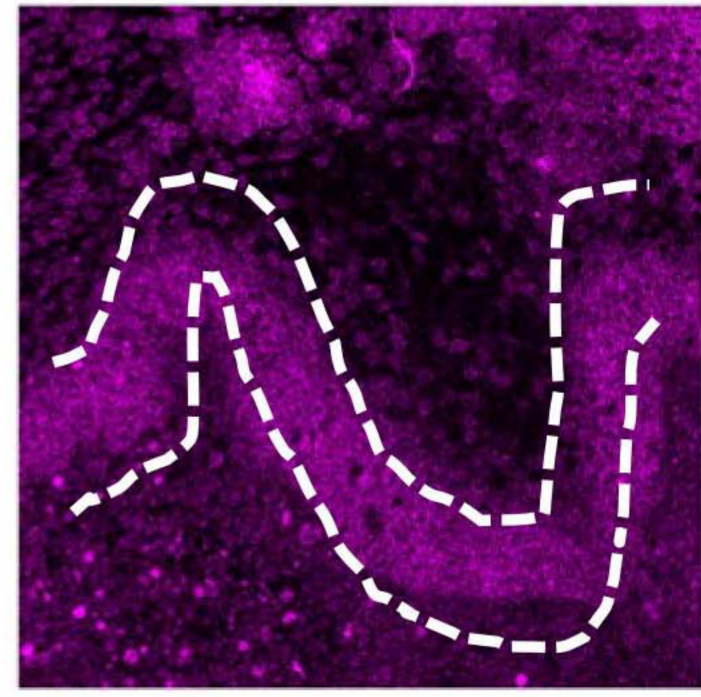




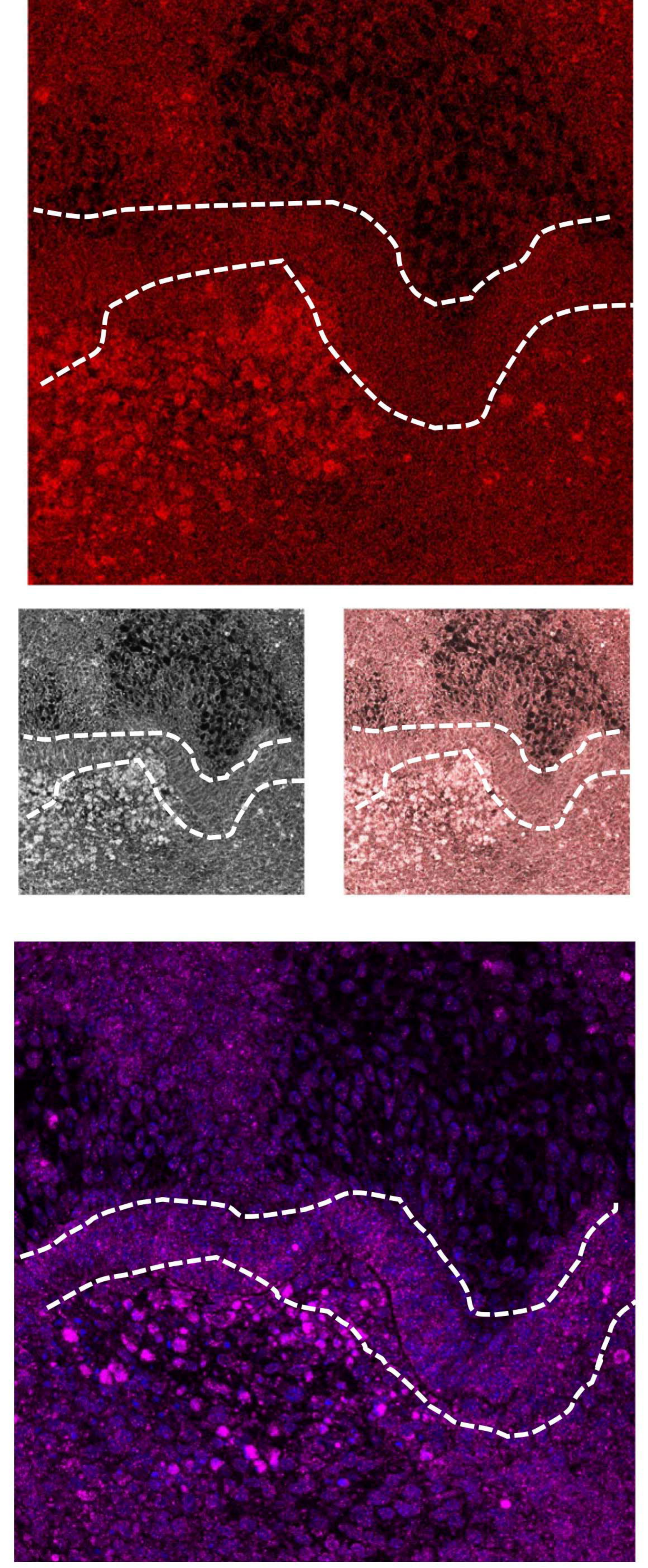


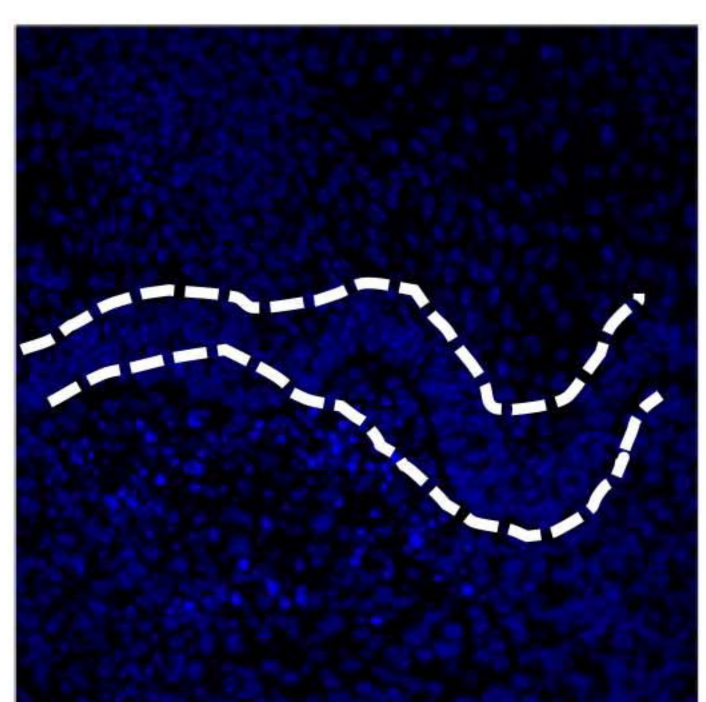


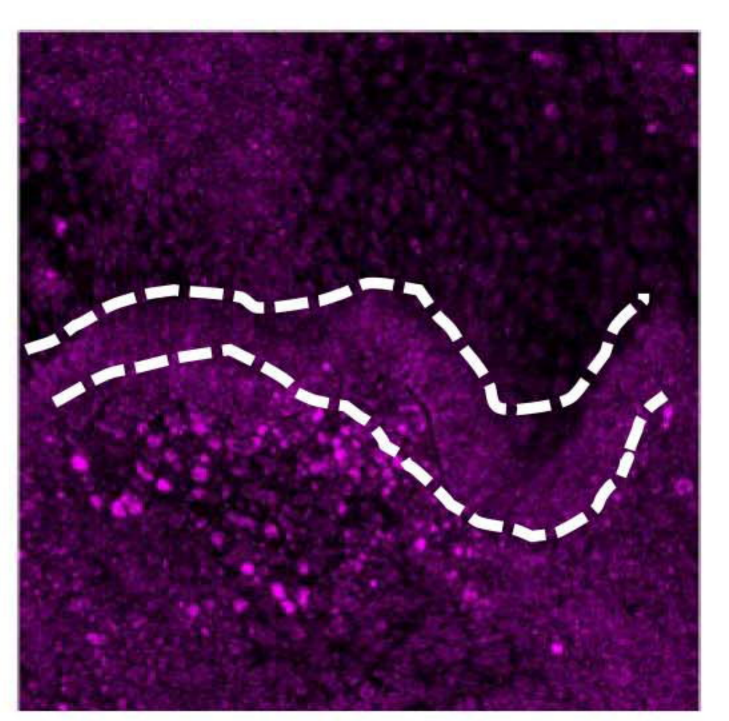


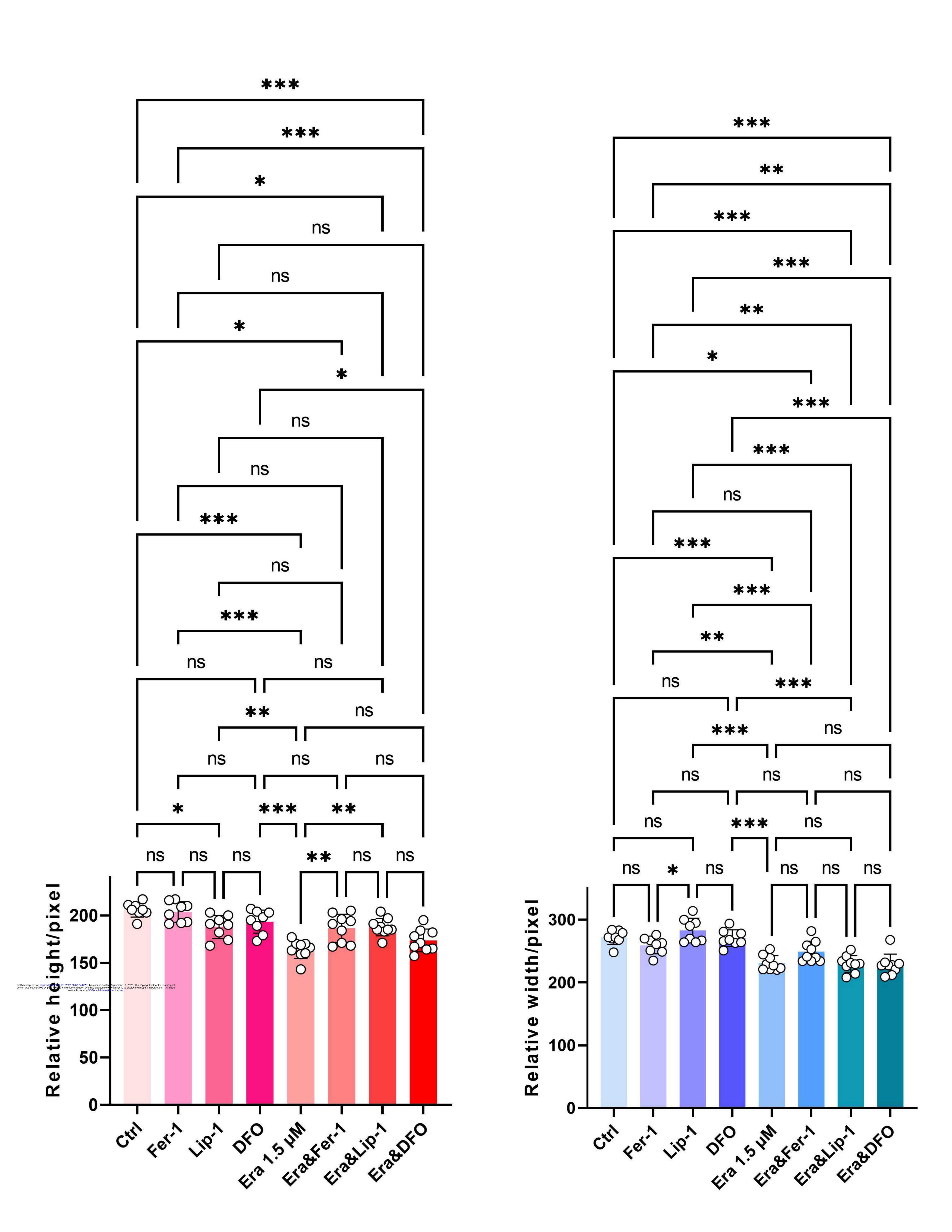


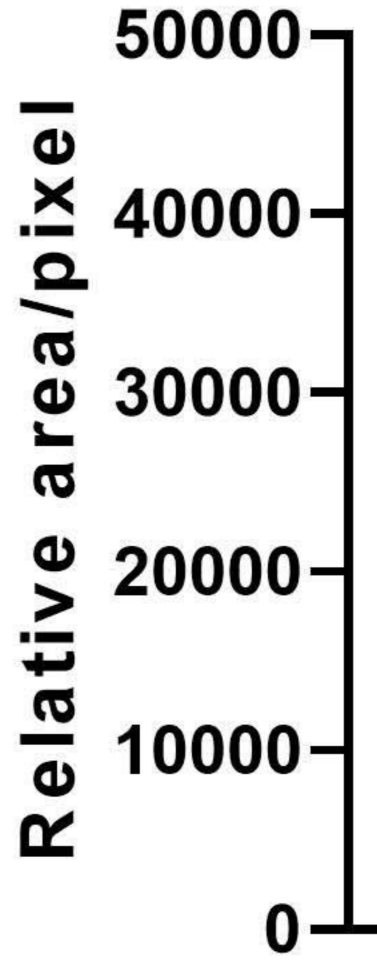
Era-20µM

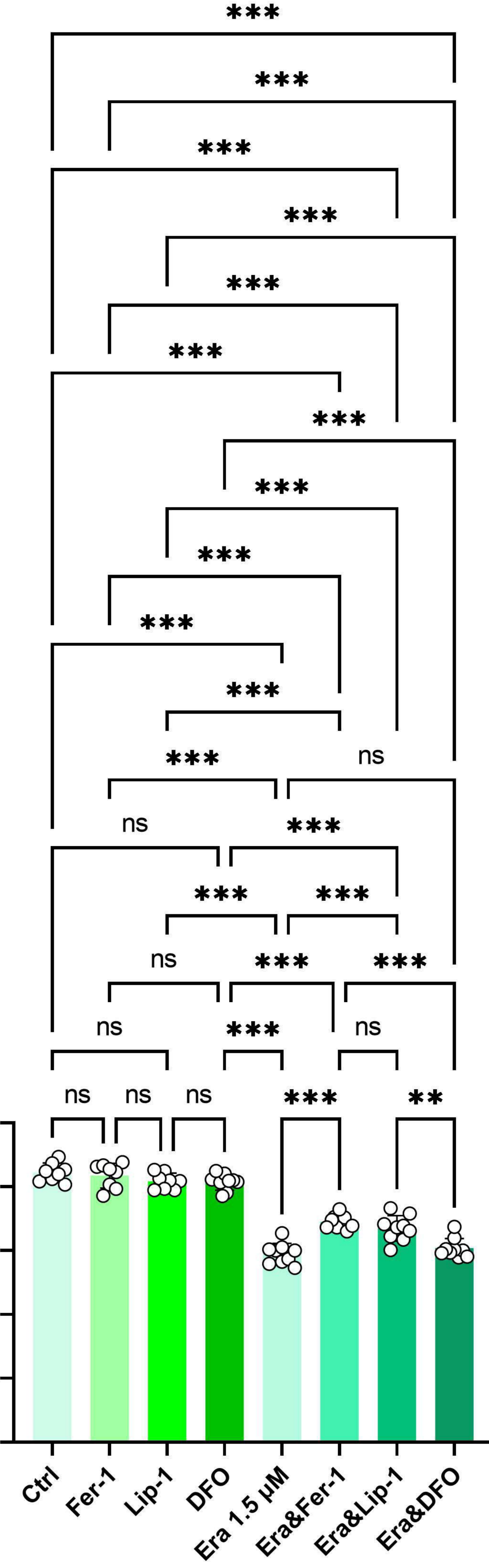




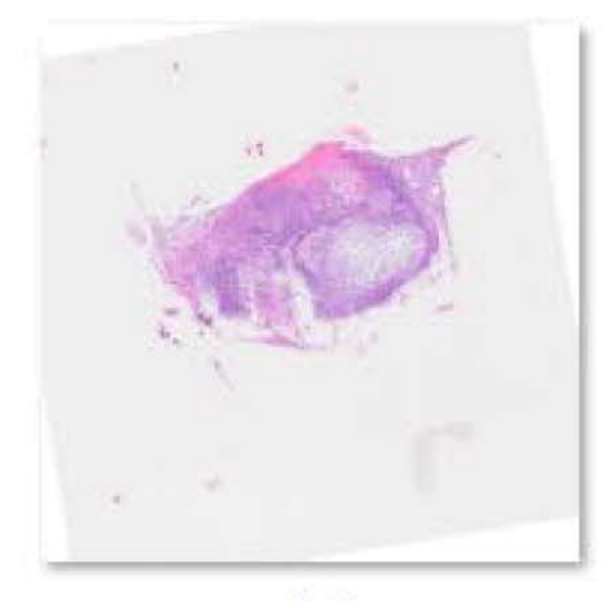










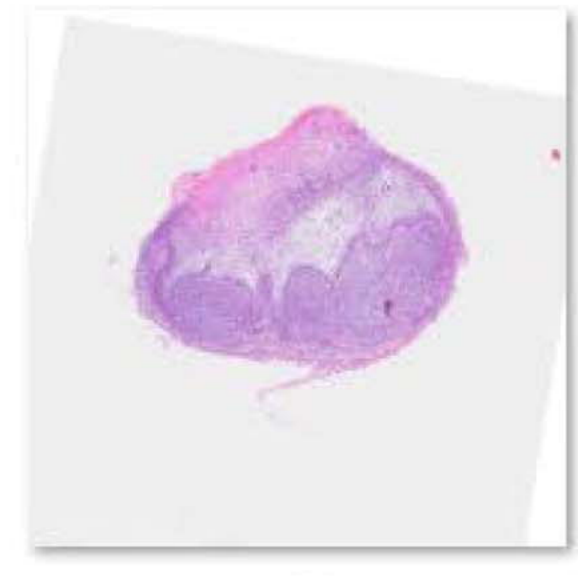


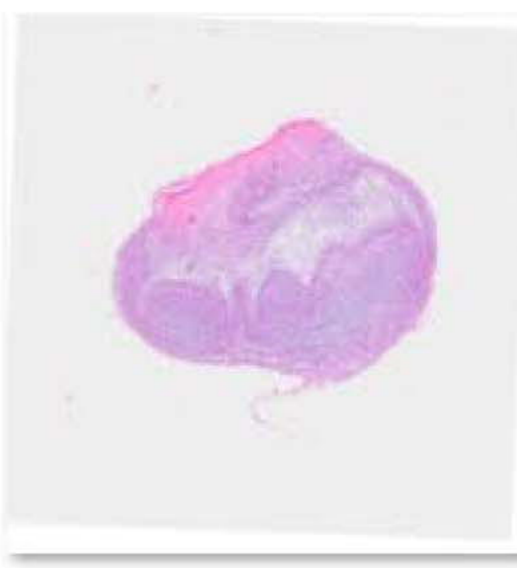






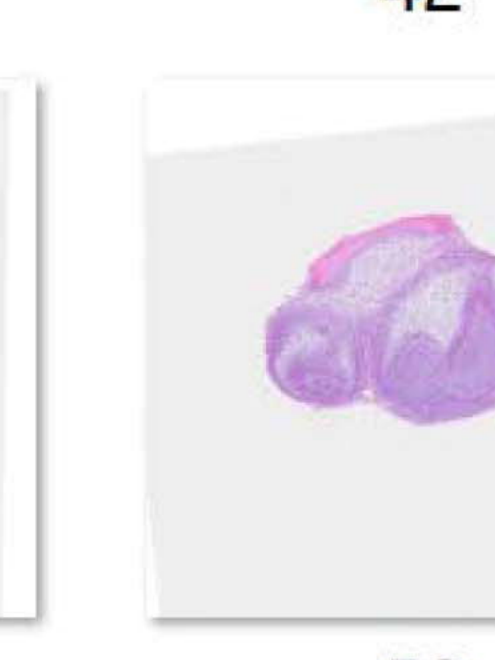


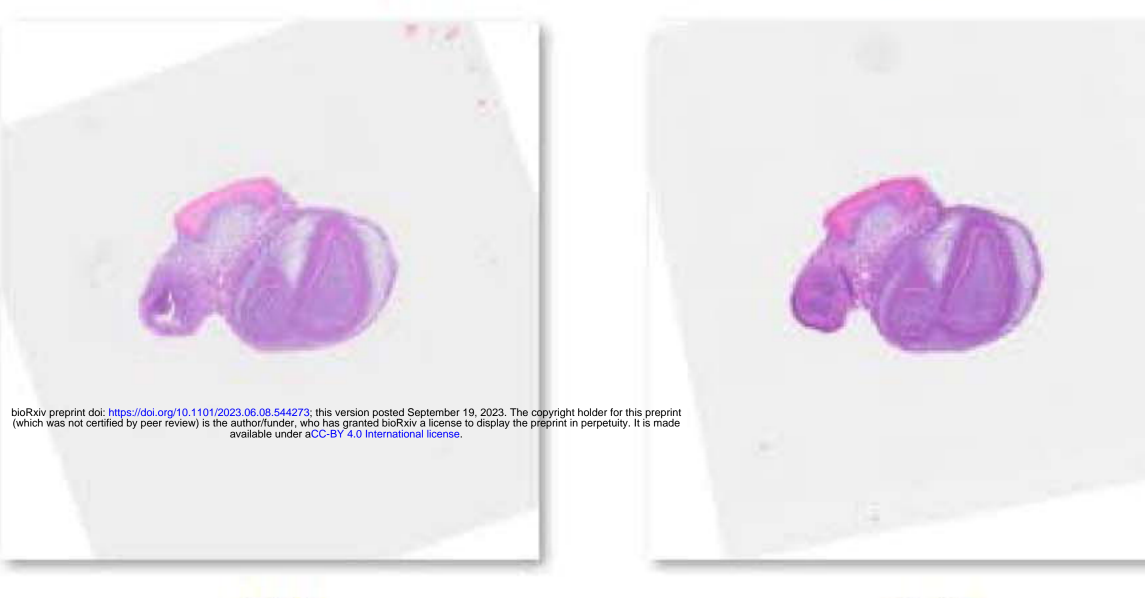




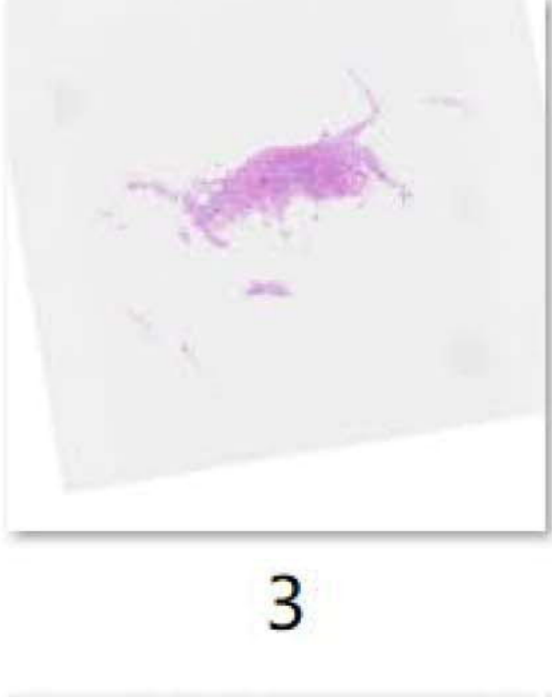


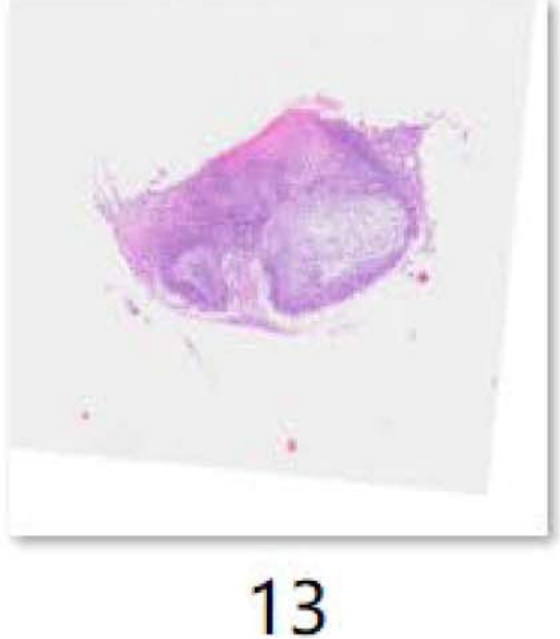


















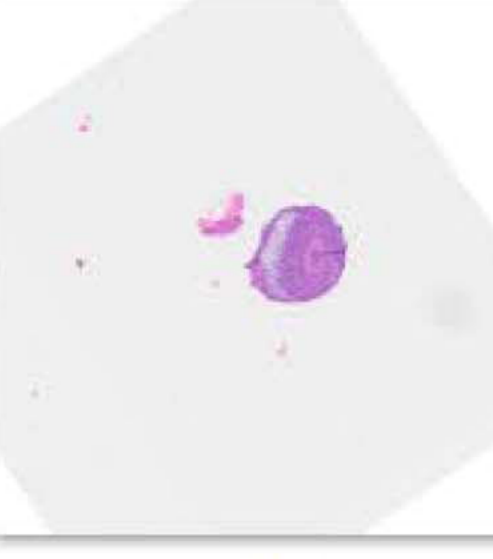


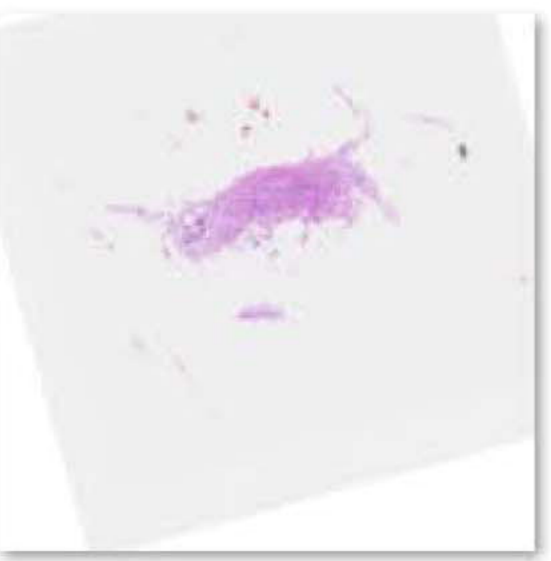




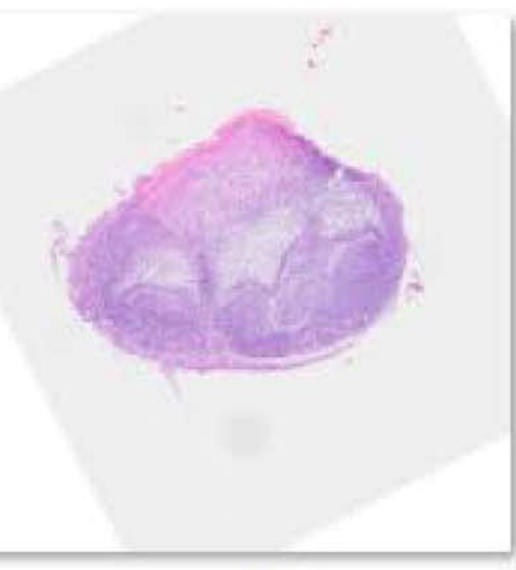


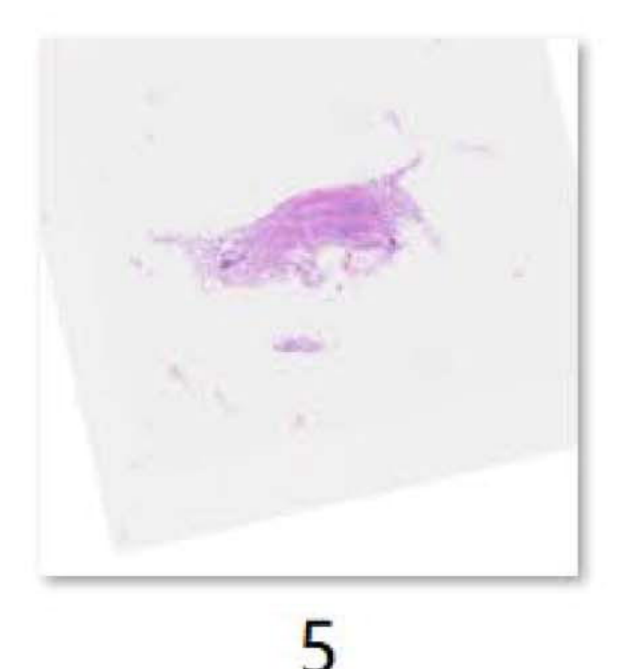


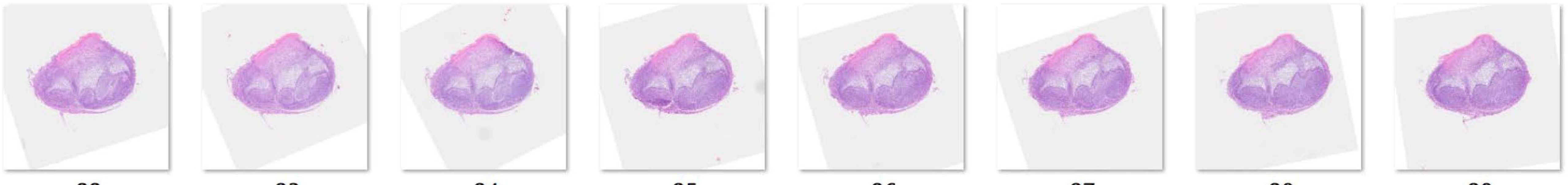












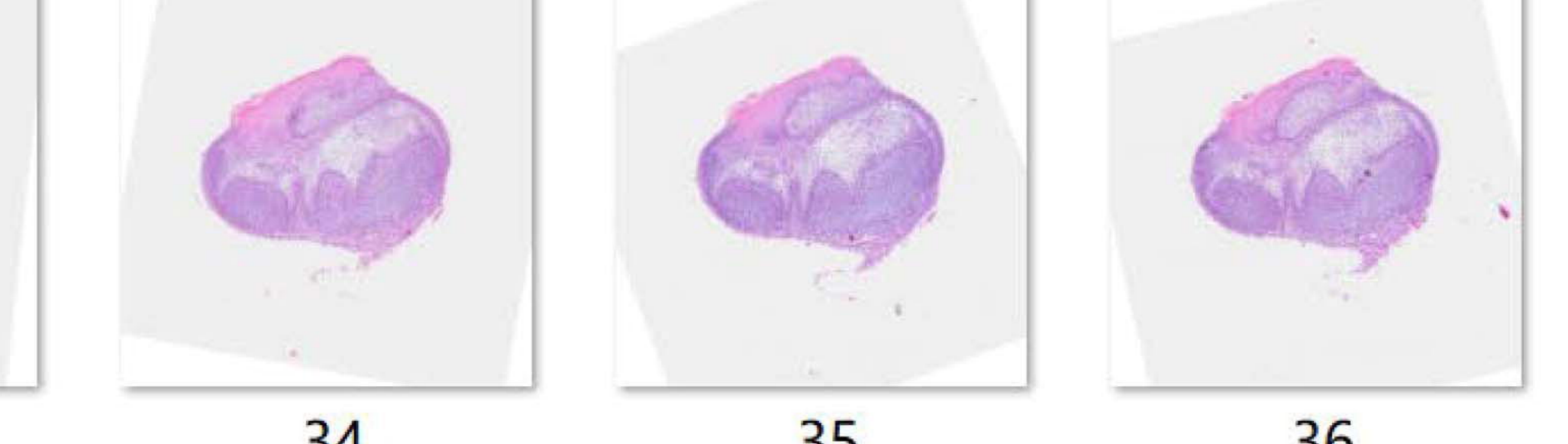
















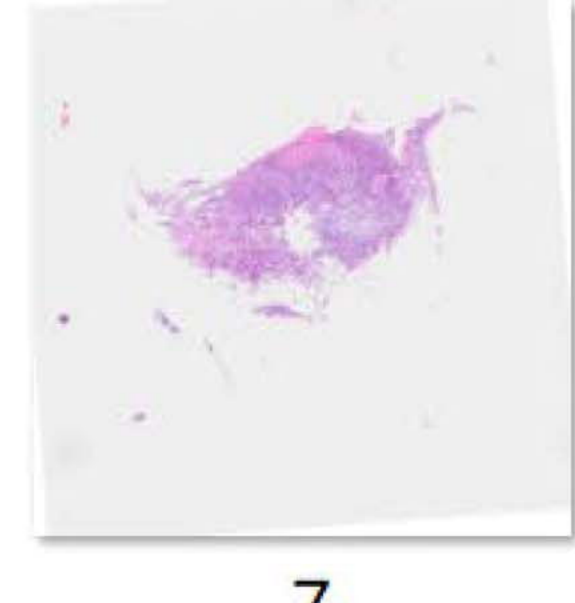


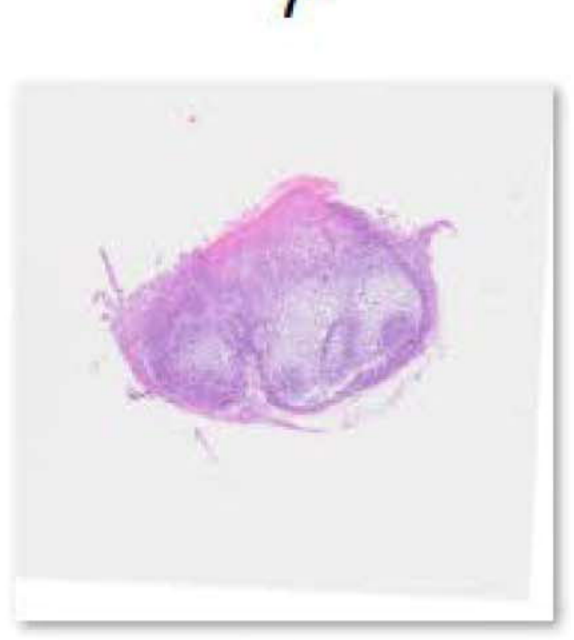




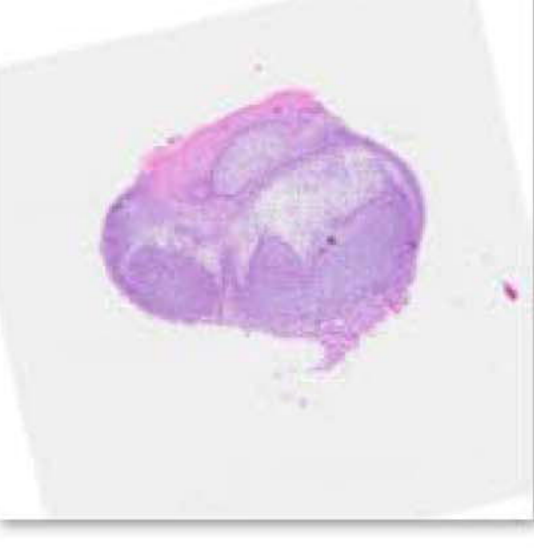












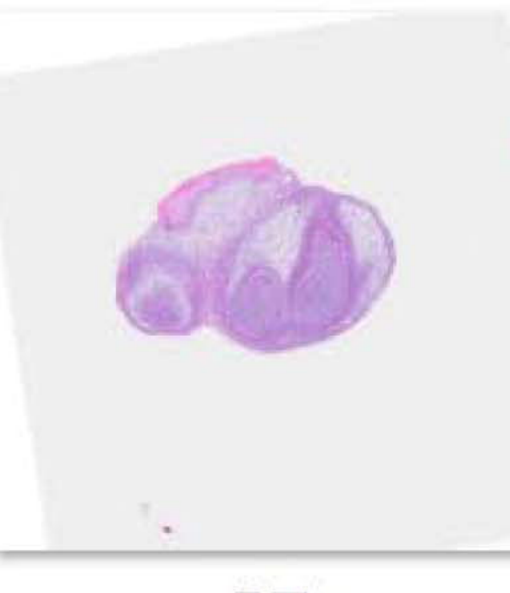








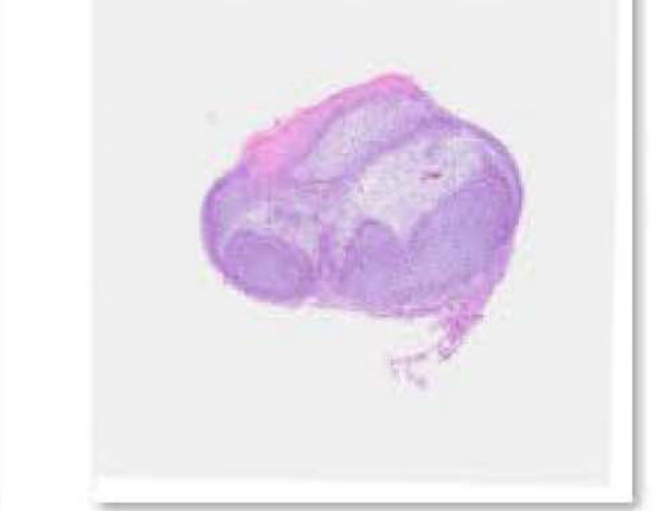




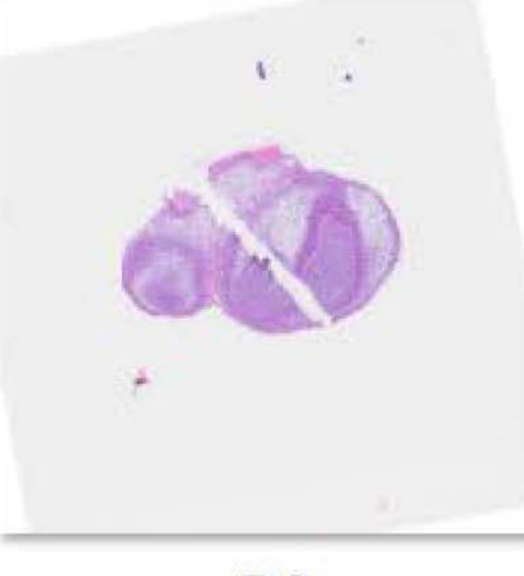






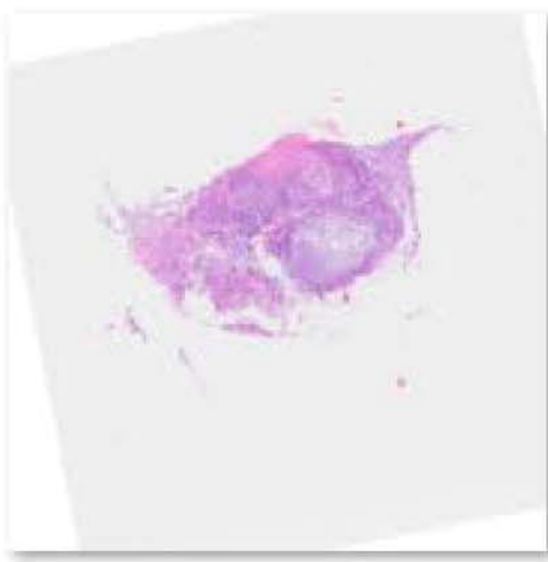




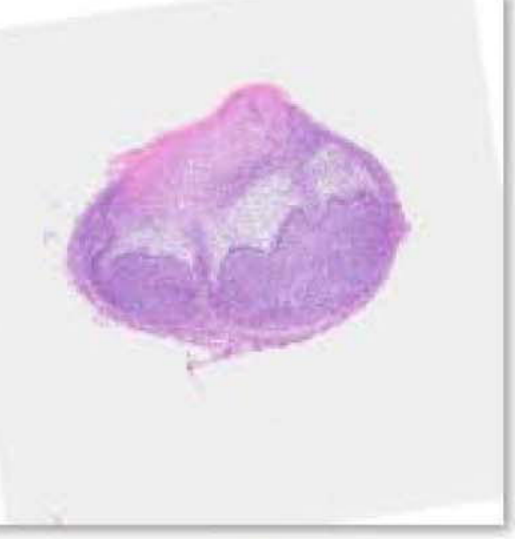


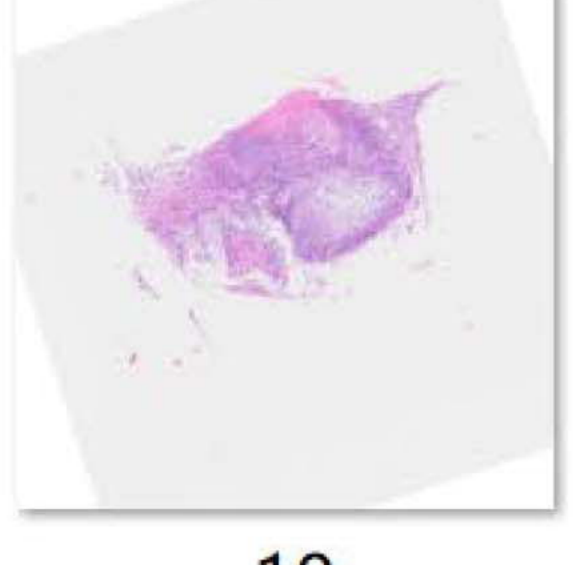










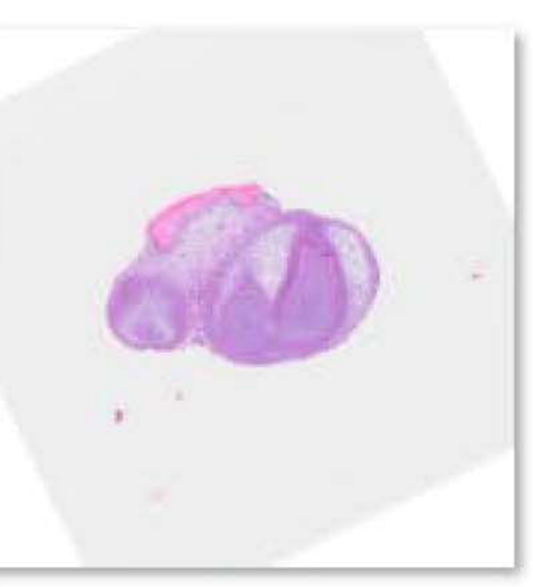






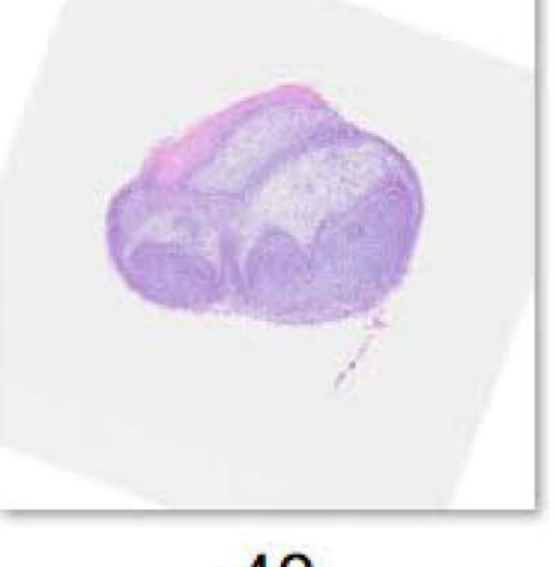




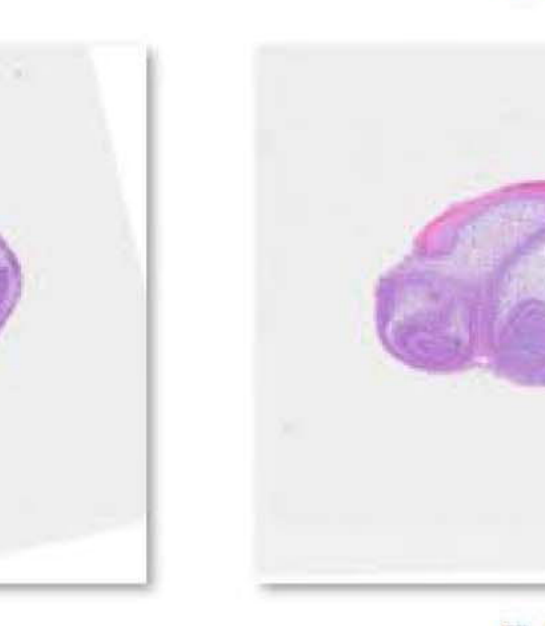










































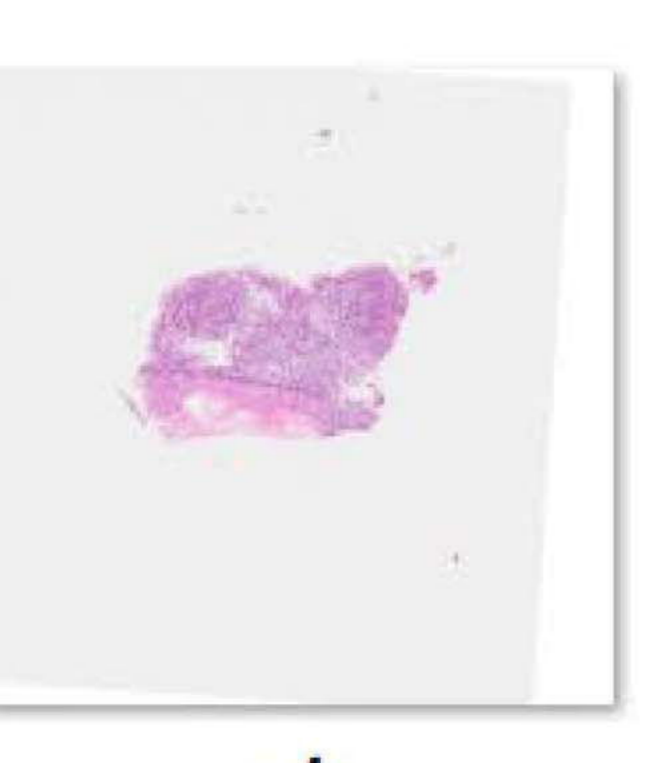




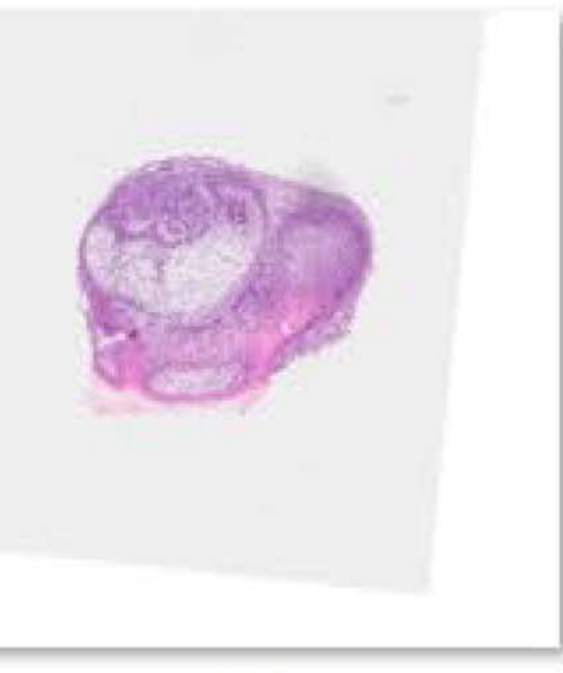










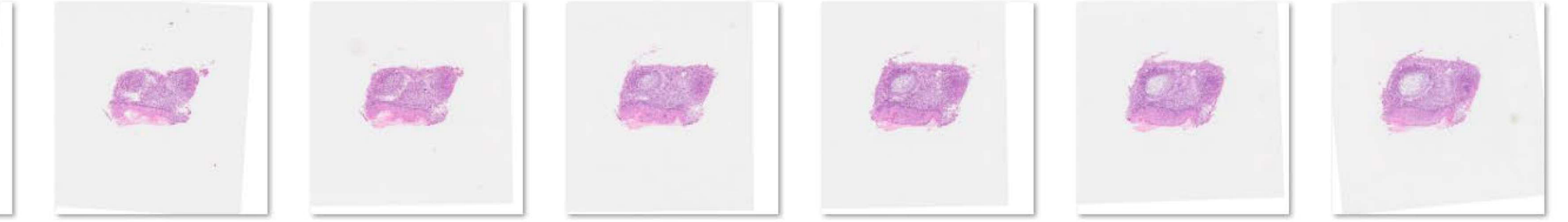




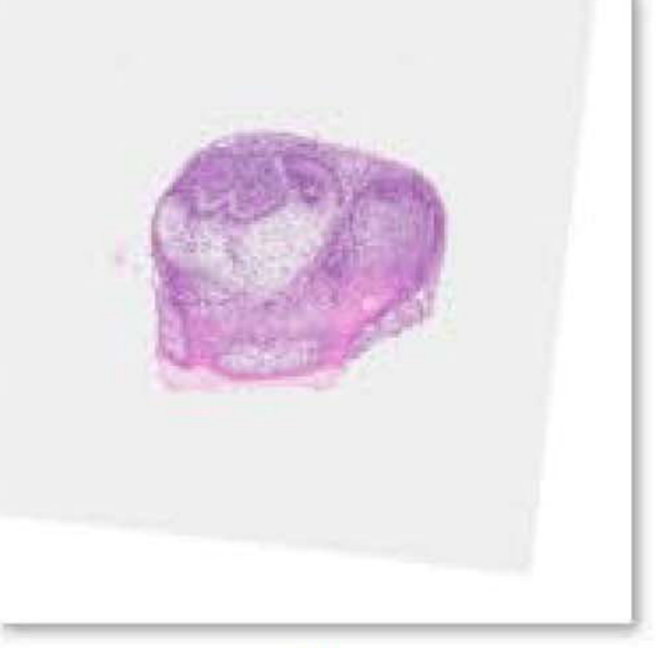




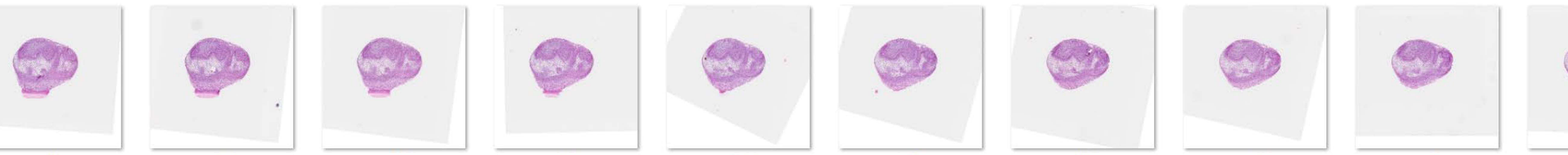


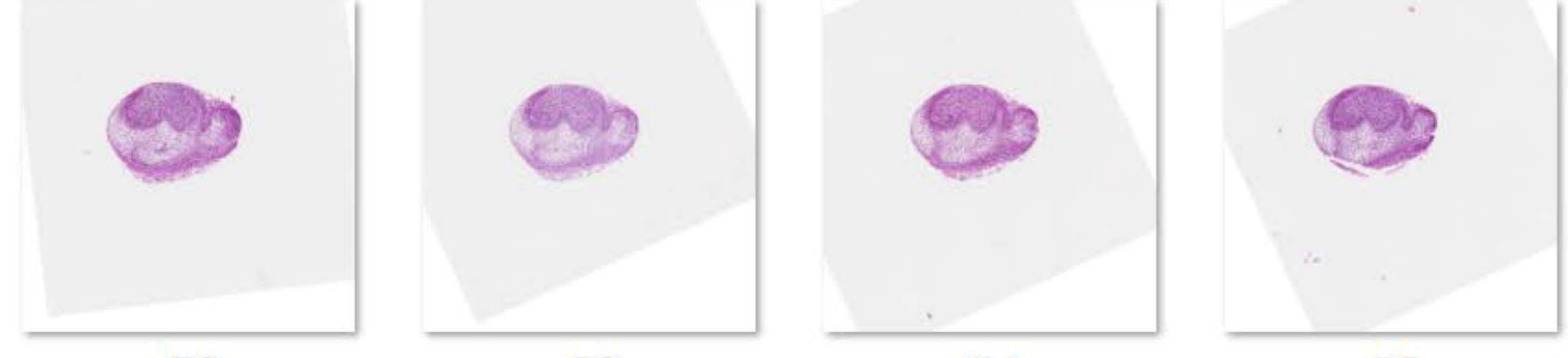






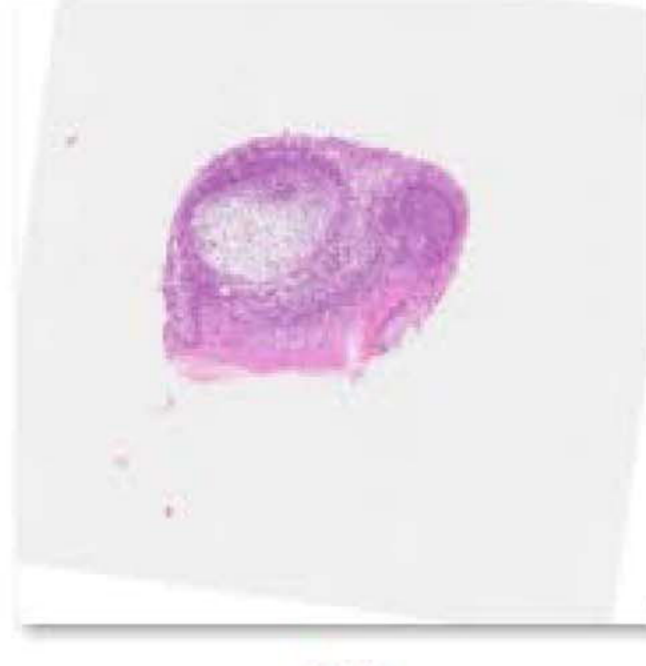


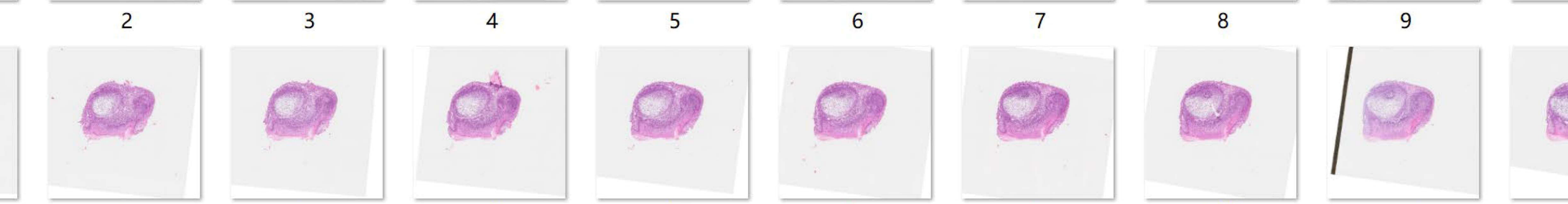






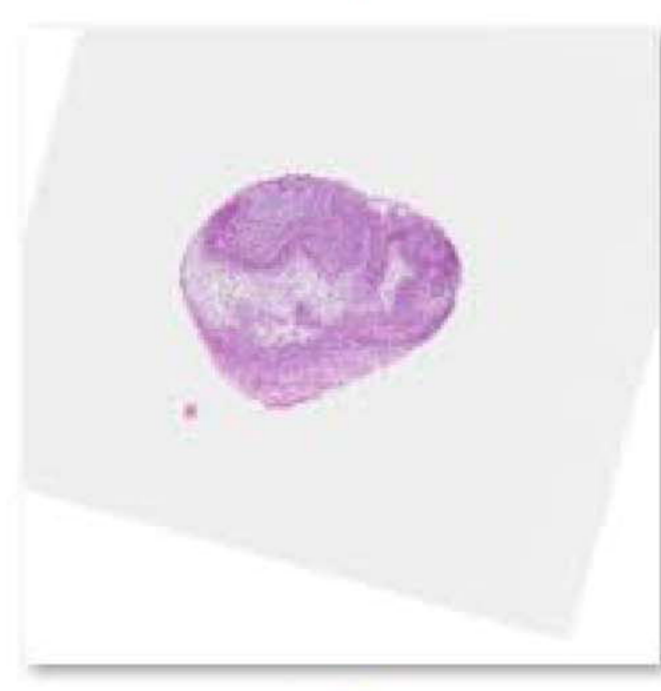








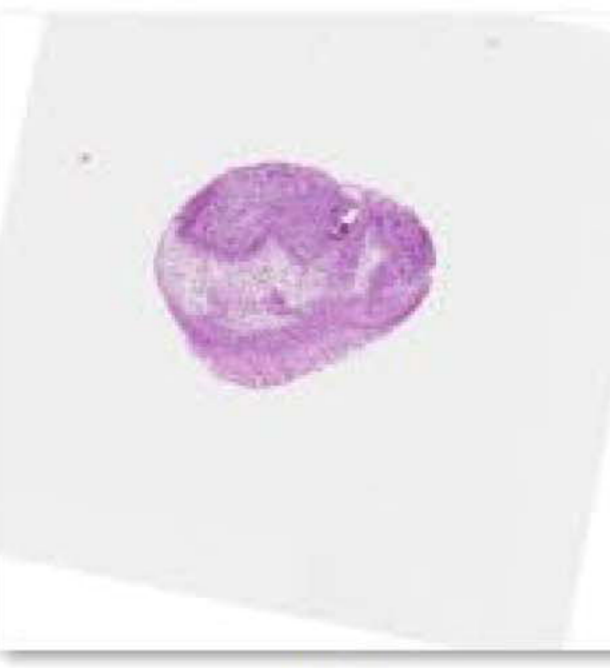




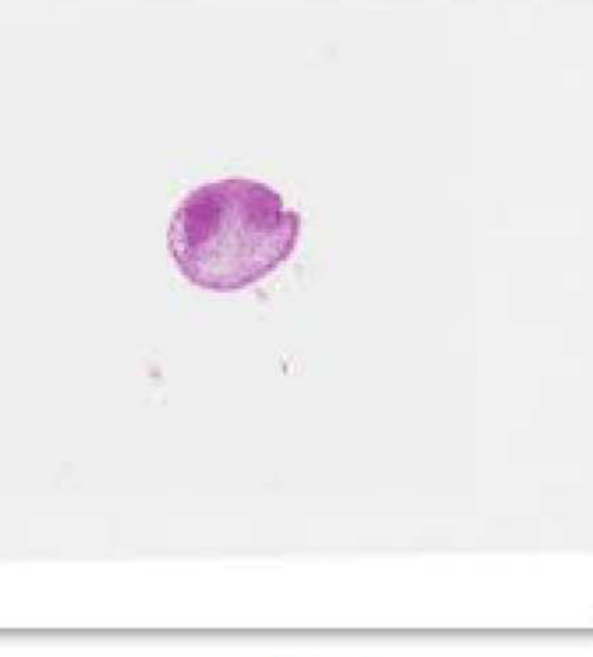


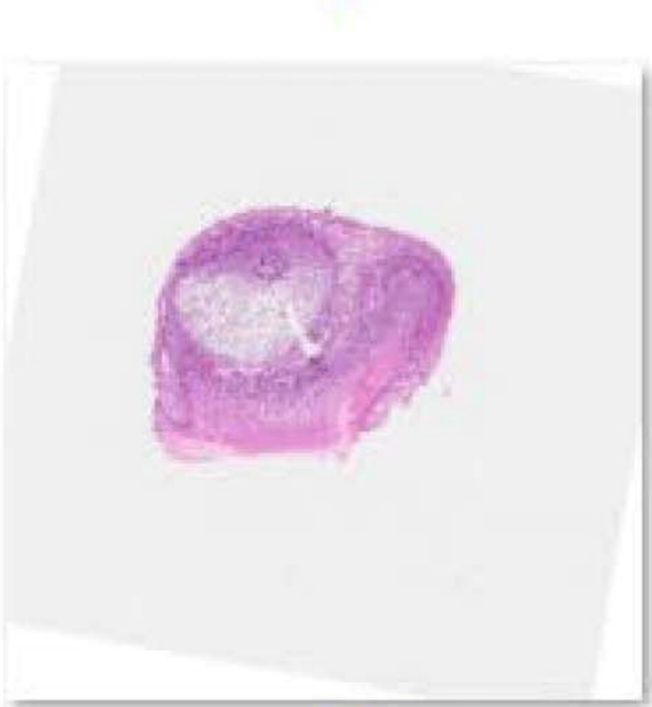


















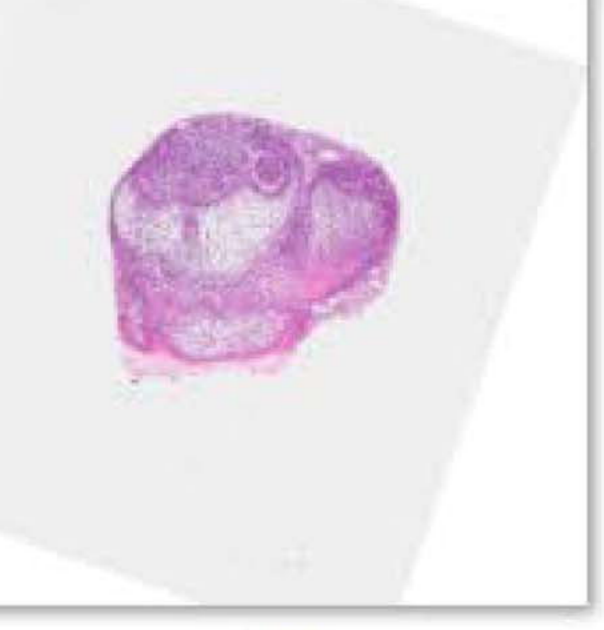






















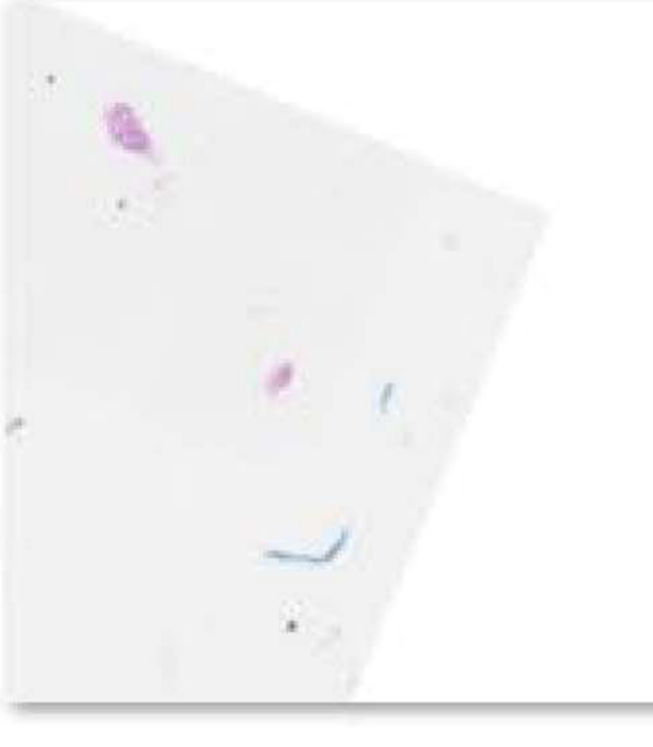






















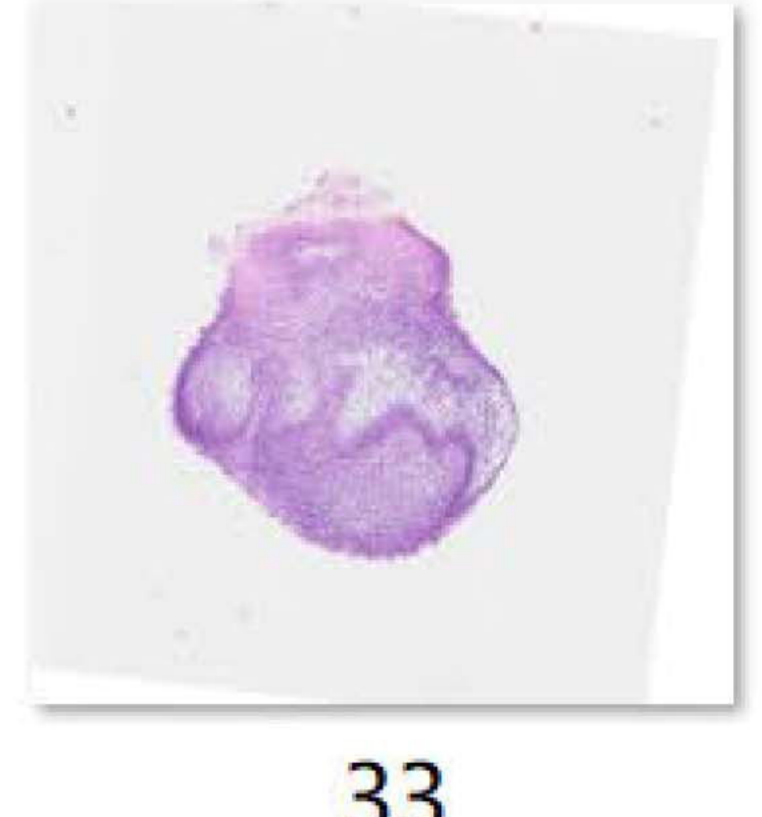


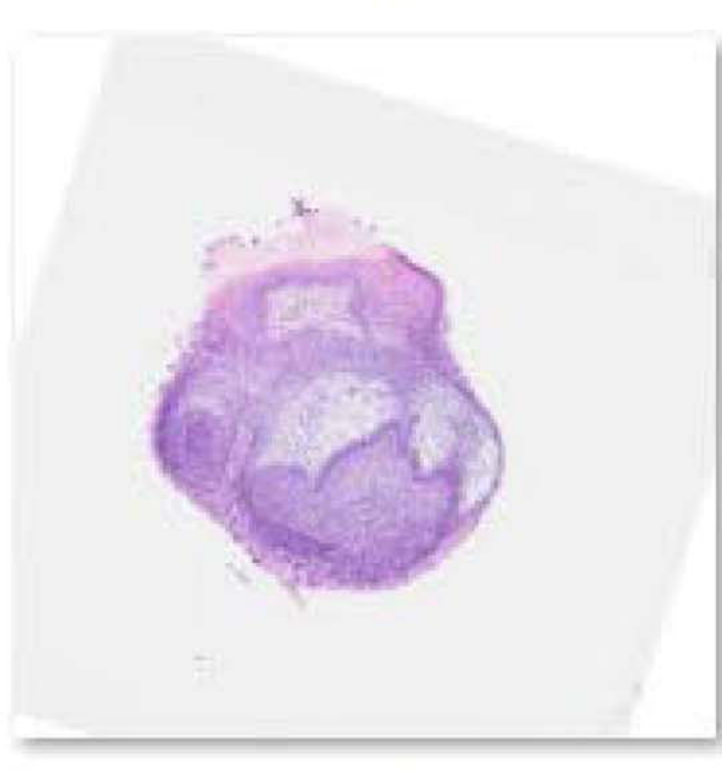






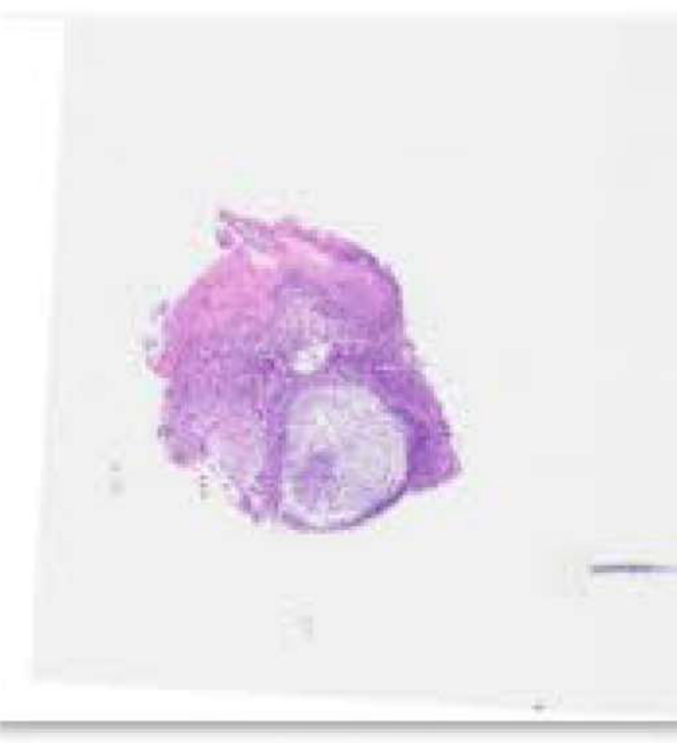


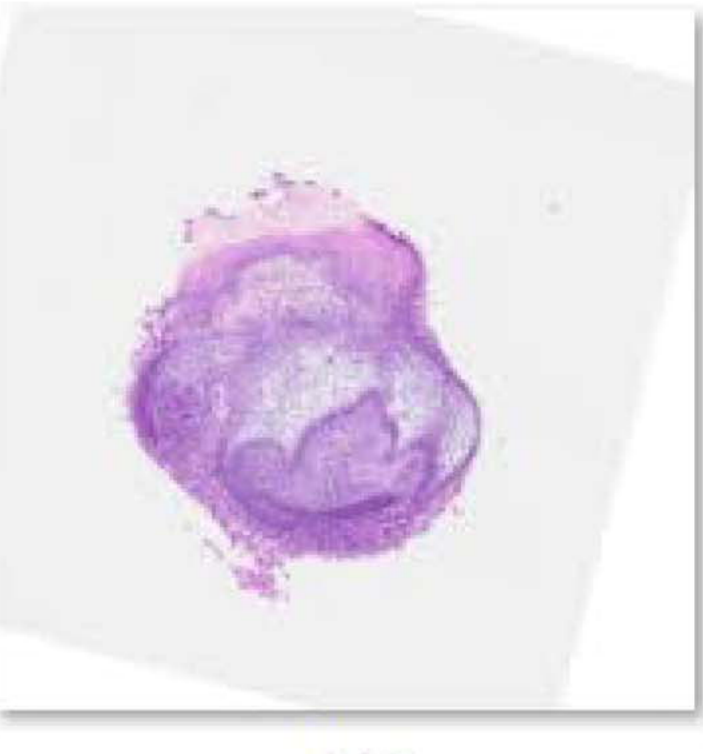


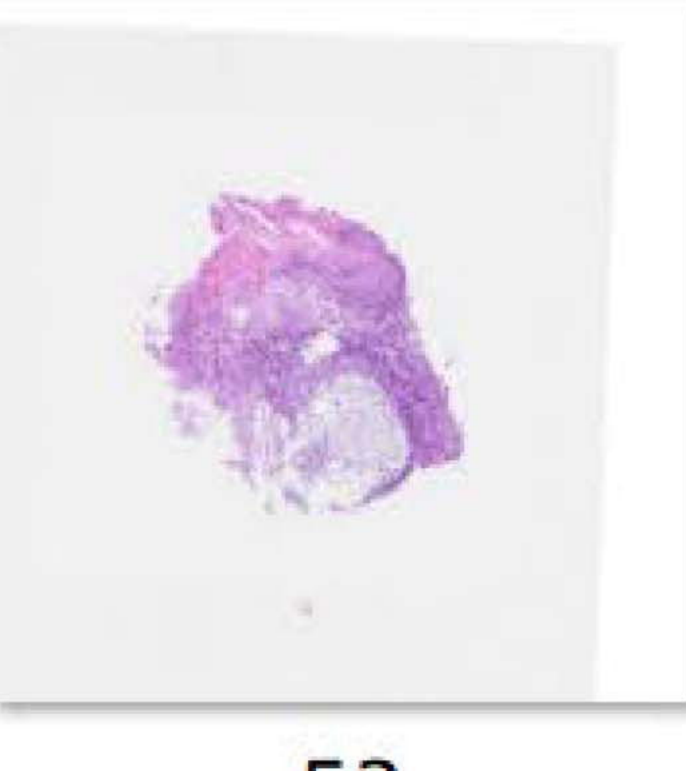




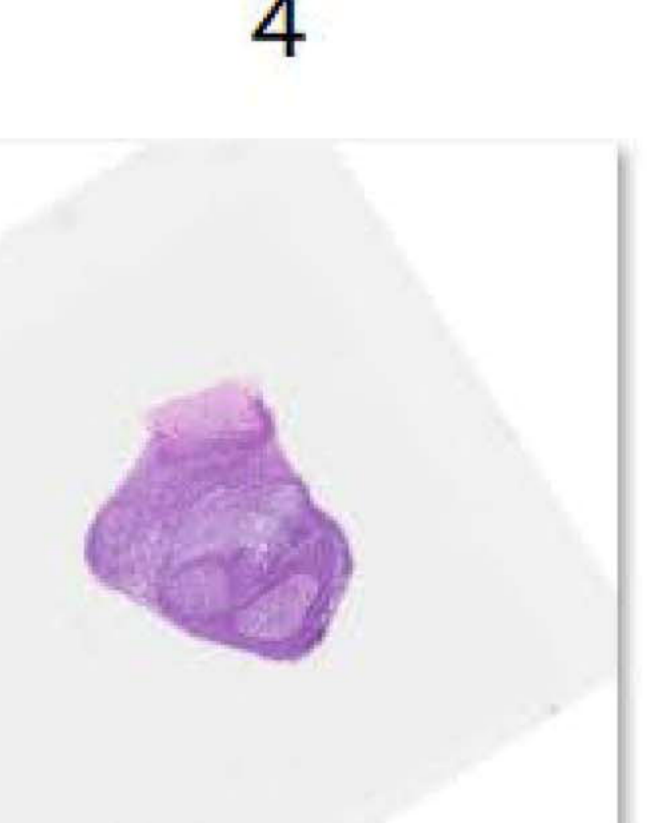


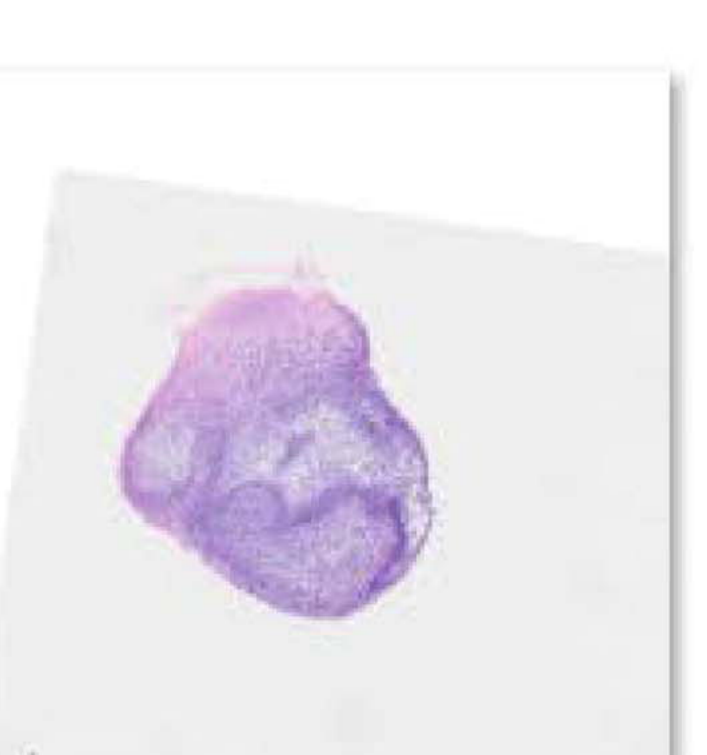










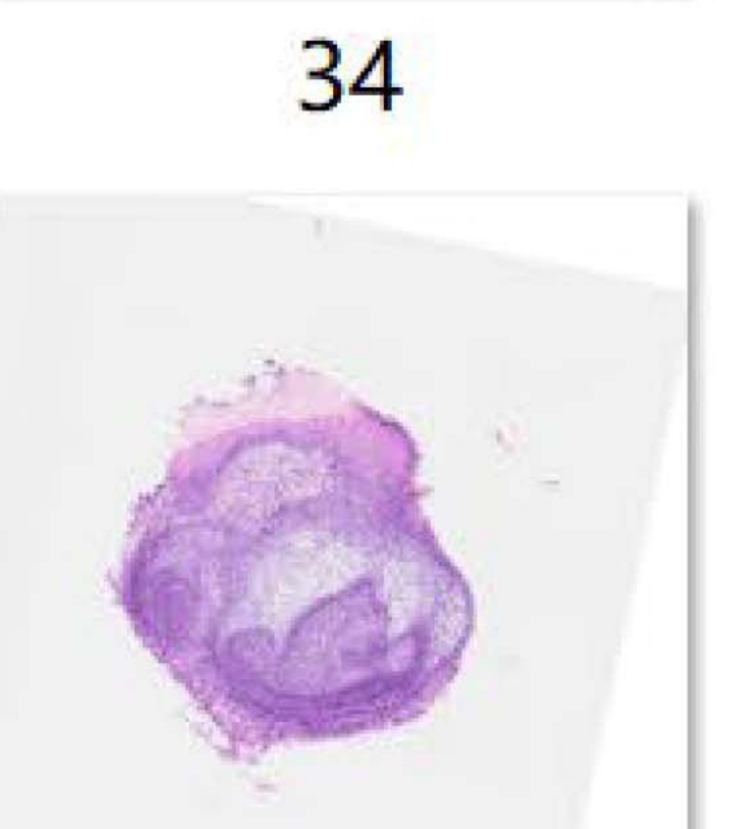














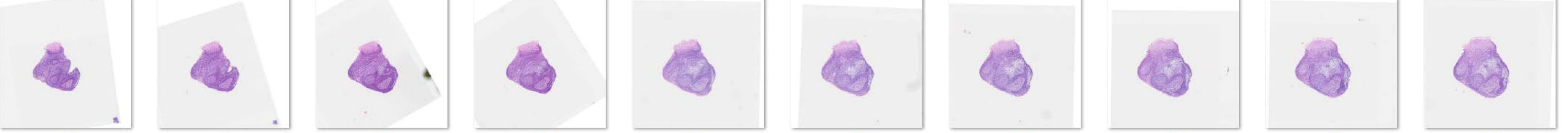








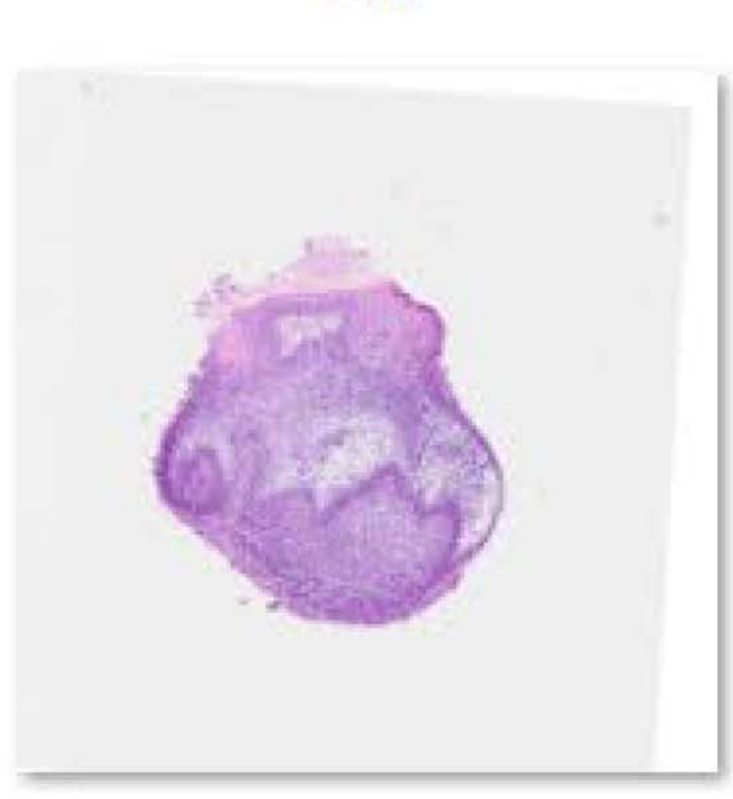




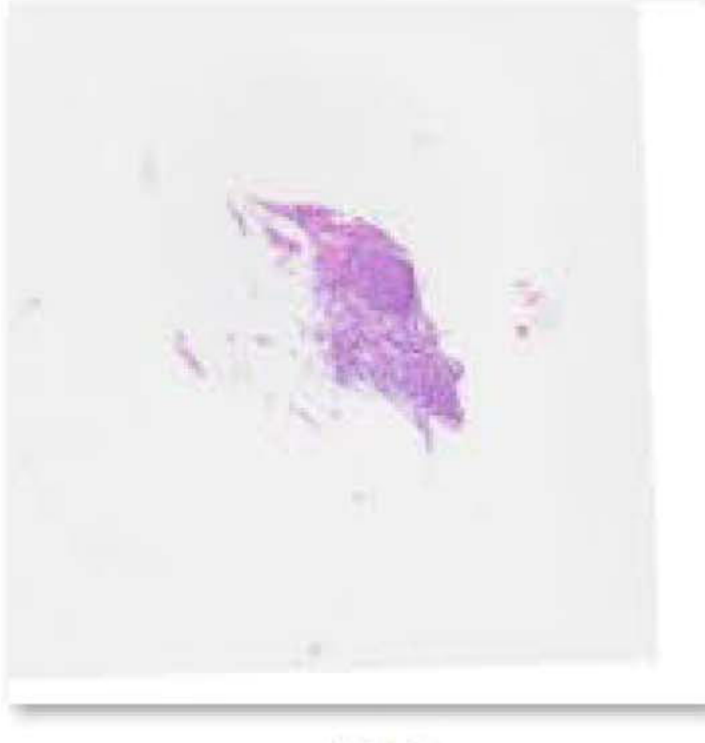






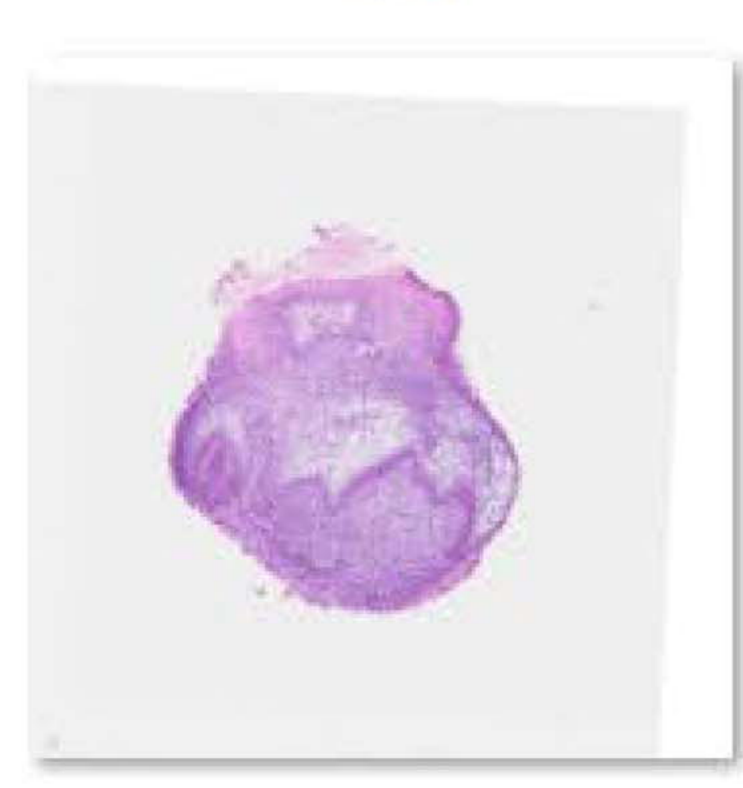


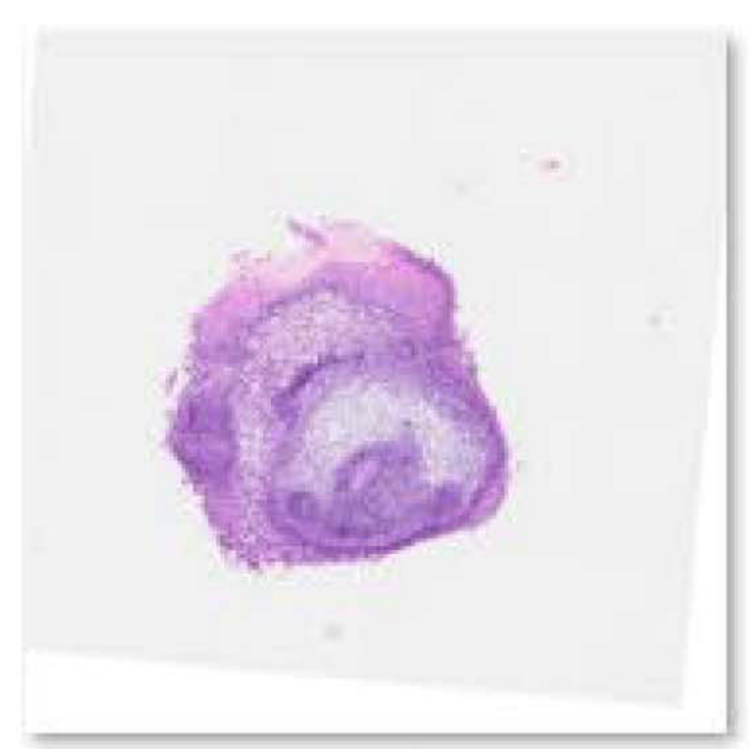








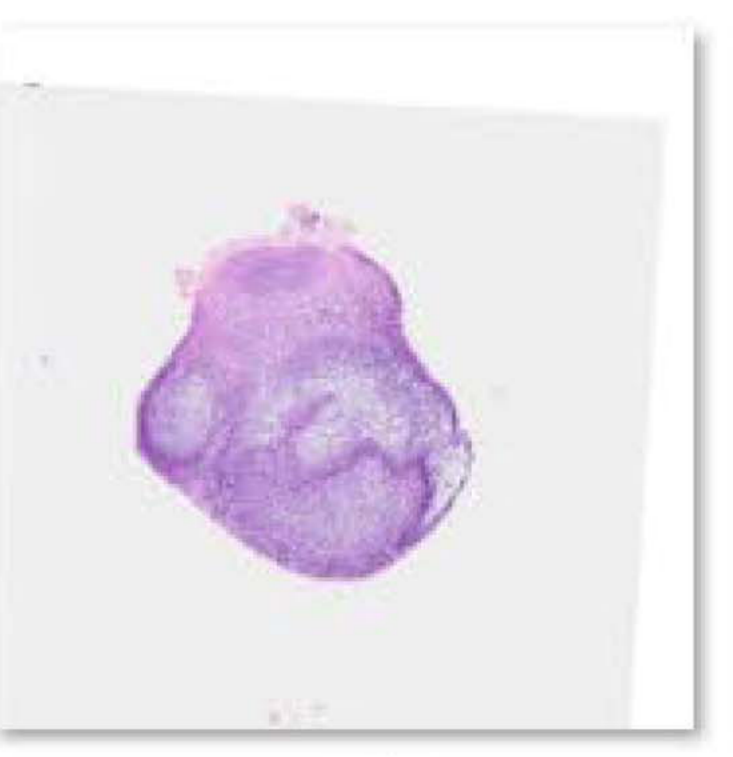






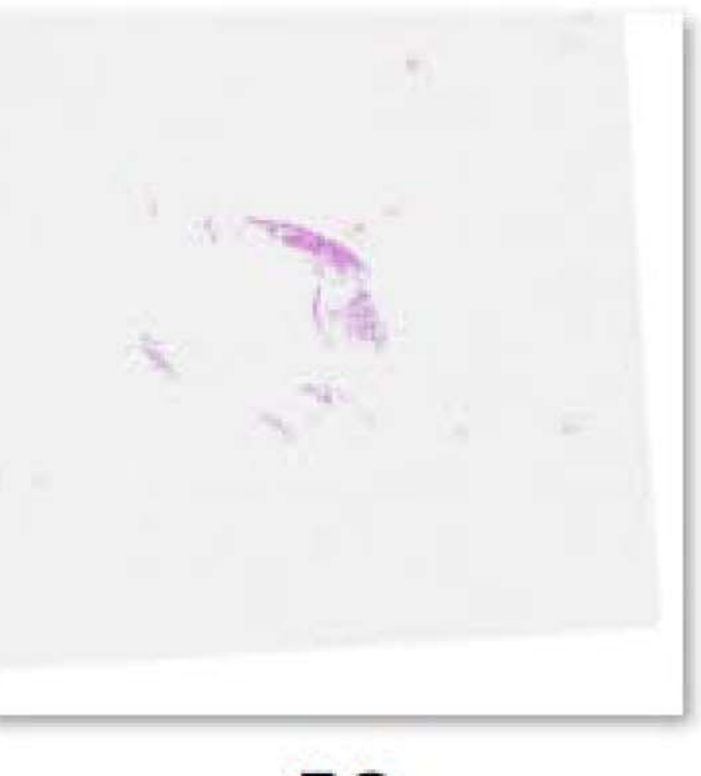


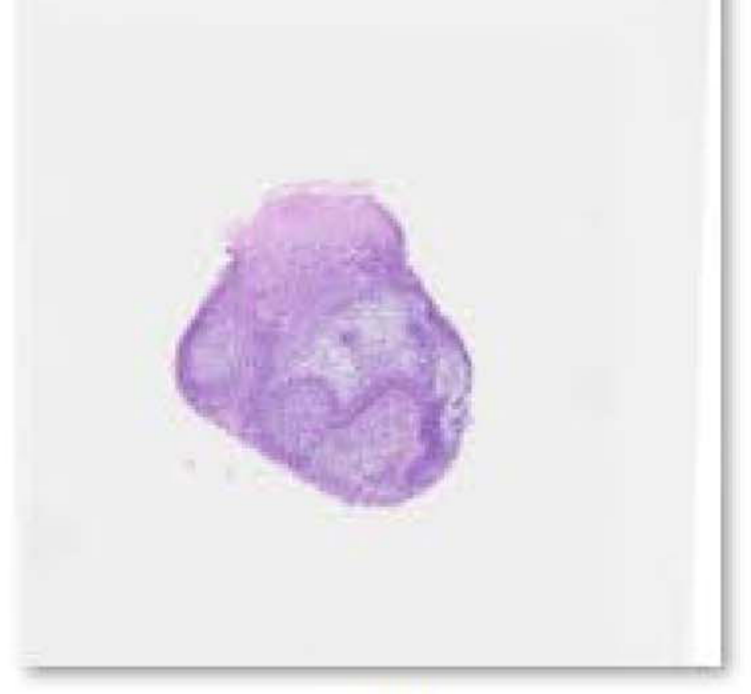




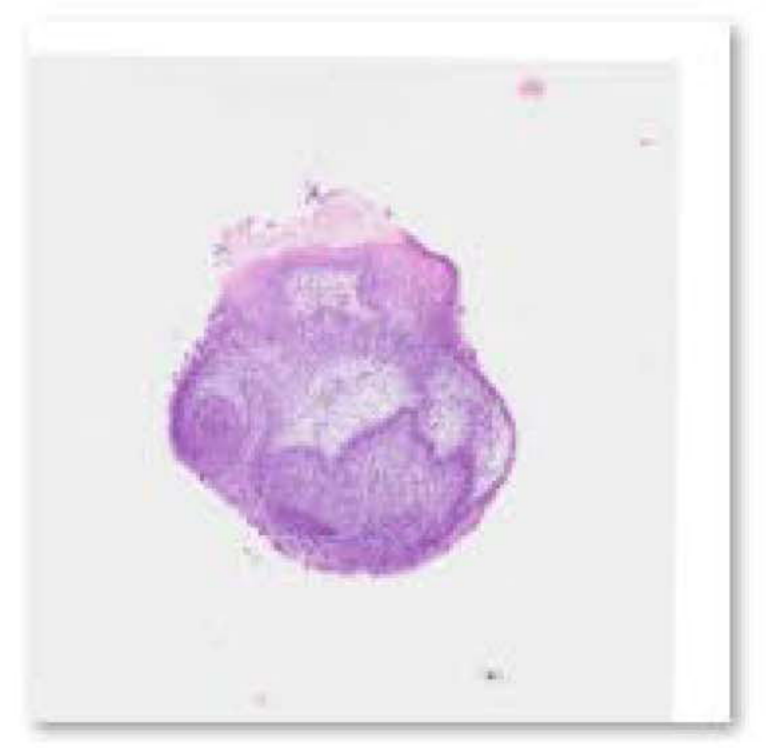










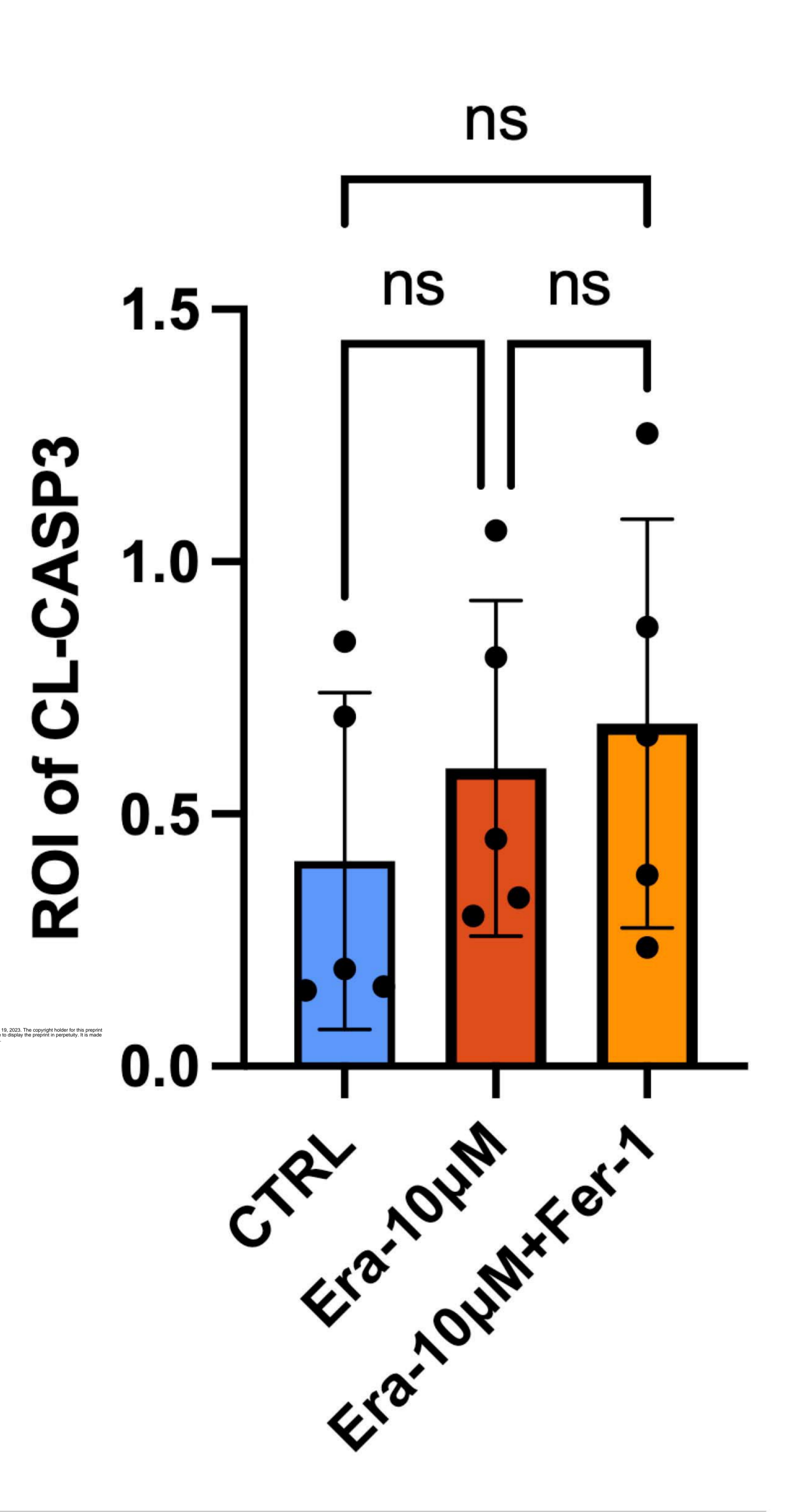




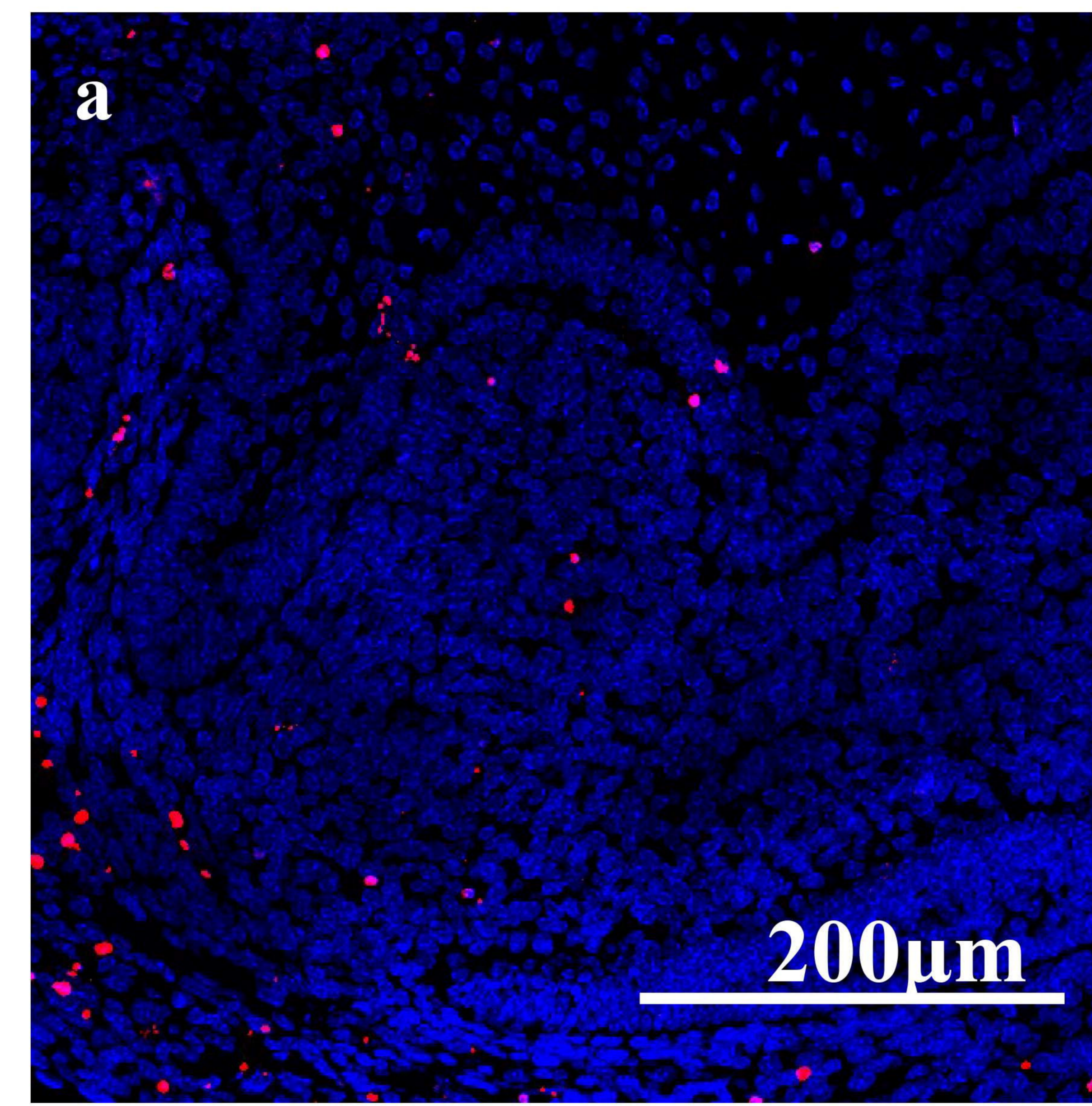




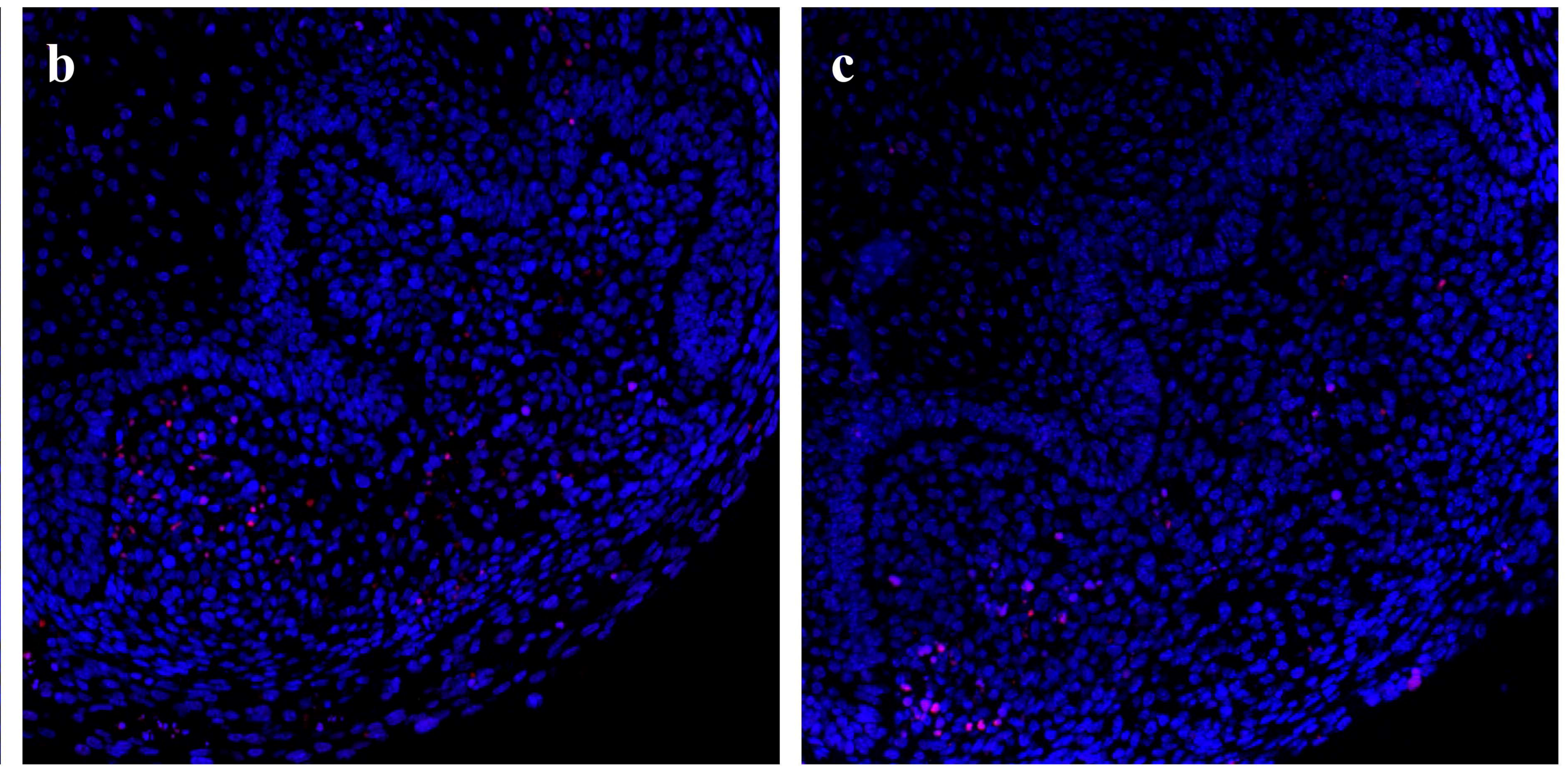
CL-CASP3



CTRL



Era-1.5µM



Era-1.5µM+Fer-1