

1 Fibroblast Stromal Support Model for Predicting Human Papillomavirus-Associated
2 Cancer Drug Responses.

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11 **Running Title:** Stroma Translational Model for HPV+ Cancer Responses

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14

15 **Abstract**

16 Currently, there are no specific antiviral therapeutic approaches targeting Human
17 papillomaviruses (HPVs), which cause around 5% of all human cancers. Specific
18 antiviral reagents are particularly needed for HPV-related oropharyngeal cancers
19 (HPV⁺OPCs) whose incidence is increasing and for which there are no early diagnostic
20 tools available. We and others have demonstrated that the estrogen receptor alpha (ER α)
21 is overexpressed in HPV⁺OPCs, compared to HPV-negative cancers in this region, and
22 that these elevated levels are associated with an improved disease outcome. Utilizing this
23 HPV⁺ specific overexpression profile, we previously demonstrated that estrogen

24 attenuates the growth and cell viability of HPV⁺ keratinocytes and HPV⁺ cancer cells *in*
25 *vitro*. Expansion of this work *in vivo* failed to replicate this sensitization. The role of
26 stromal support from the tumor microenvironment (TME) has previously been tied to
27 both the HPV lifecycle and *in vivo* therapeutic responses. Our investigations revealed that
28 *in vitro* co-culture with fibroblasts attenuated HPV⁺ specific estrogen growth responses.
29 Continuing to monopolize on the HPV⁺ specific overexpression of ER α , our co-culture
30 models then assessed the suitability of the selective estrogen receptor modulators
31 (SERMs), raloxifene and tamoxifen, and showed growth attenuation in a variety of our
32 models to one or both of these drugs *in vitro*. Utilization of these SERMs *in vivo* closely
33 resembled the sensitization predicted by our co-culture models. Therefore, the *in vitro*
34 fibroblast co-culture model better predicts *in vivo* responses. We propose that utilization
35 of our co-culture *in vitro* model can accelerate cancer therapeutic drug discovery.

36 **Importance**

37 Human papillomavirus-related cancers (HPV⁺ cancers) remain a significant public health
38 concern, and specific clinical approaches are desperately needed. In translating drug
39 response data from *in vitro* to *in vivo*, the fibroblasts of the adjacent stromal support
40 network play a key role. Our study presents the utilization of a fibroblast 2D co-culture
41 system to better predict translational drug assessments for HPV⁺ cancers. We also
42 suggest that this co-culture system should be considered for other translational
43 approaches. Predicting even a portion of treatment paradigms that may fail *in vivo* with a
44 co-culture model will yield significant time, effort, resource, and cost efficiencies.

45 **Keywords**

46 Estrogen, ER α , stroma, HPV, human papillomavirus, oropharyngeal cancer, Raloxifene,
47 Tamoxifen, therapeutics

48 **Introduction**

49 Human papillomaviruses (HPVs) are small, double-stranded DNA viruses, and
50 high-risk HPVs are known carcinogens^{1–7}. HPV is the most common sexually transmitted
51 infection in the United States (U.S.), and estimated to infect more than 80% of the
52 population at least once in their lifetime^{1–3,8–12}. HPV16 is the most prevalent genotype,
53 accounting for at least 50% of cervical cancers and approximately 90% of HPV $^+$
54 oropharyngeal cancers (HPV $^+$ OPCs)^{9,13,14}. While prophylactic HPV vaccines have
55 already begun to show remarkable efficacy in preventing infection and related diseases,
56 HPV continues to account for ~5% of worldwide cancer, and disproportionately affects
57 marginalized populations both in the U.S. and around the world^{1–3,8–12,15–17}. As such, the
58 lack of specific antiviral therapeutics available for combatting HPV-related cancers is of
59 significant concern.

60 While many cancers are on the decline, the last two decades have shown a sharp
61 increase in HPV $^+$ OPCs, for which there are no early diagnostic tools available^{4–6,18,19}.
62 HPV $^+$ OPCs are found at 4-fold higher levels in men than in women, suggesting there are
63 sex-related differences in the development of these cancers^{4,18}. Using data from The
64 Cancer Genome Atlas (TCGA), we and others have shown that the estrogen receptor
65 alpha (ER α) is overexpressed in HPV $^+$ HNC (head and neck cancers including OPCs) and
66 that these elevated levels are associated with an improved disease outcome^{18,20–26}.

67 We have also previously demonstrated that 17- β estradiol (estrogen) attenuates
68 the growth and cell viability of HPV $^+$ keratinocytes and HPV $^+$ cancer cells *in vitro*, but

69 not HPV negative (HPV⁻) keratinocytes or HPV⁺ cancer cells²⁶. Sensitization occurs via
70 numerous mechanisms: 1) at the level of viral transcription, 2) via interactions with E6
71 and E7, 3) through manipulation of cell survival and cell death pathways²⁶. Here, we
72 report that the expansion of estrogen treatment into *in vivo* *NOD-scid IL2Rg^{null}* (NSG)
73 mice revealed a lack of response to estrogen alone or in combination with radiotherapy
74 (IR).

75 Previously, estrogen has been shown to promote HPV-induced cervical disease in
76 immunocompetent mice, yet this enhancement was lost in NSGs²⁷⁻³². HPV oncogenes, in
77 conjunction with estrogen, were shown to fundamentally reprogram the tumor
78 microenvironment (TME)^{27,28,33}. We therefore sought to determine if the TME,
79 specifically the stromal support provided by fibroblasts, alters the estrogenic effects in
80 our model systems³³⁻⁴². *In vitro* co-culture studies revealed that stromal interactions
81 markedly change cell growth and viability in response to estrogen in our HPV⁺ models.

82 While our estrogen studies did not prove effective *in vivo*, ER α remains
83 overexpressed in HPV⁺OPC and HPV⁺ cervical cancers^{24-27,29,31-33,43}. Selective estrogen
84 receptor modulators (SERMs) have proven to provide a multitude of therapeutic
85 applications⁴⁴⁻⁵⁰. Analysis of K14E6/E7 transgenic mice models have previously shown
86 the possible utility of raloxifene on reduction of recurrence of cervical neoplastic disease,
87 and earlier literature suggested the utility of tamoxifen to lengthen the latent period for
88 cervical dysplasia and carcinoma in carcinogen-induced models^{45,51}. We utilized our co-
89 culture system to assess the efficacy of raloxifene and tamoxifen and showed significant
90 growth repression to one or both SERMs in a number of cancer cell lines. Our
91 preliminary assessment of these SERMs in an *in vivo* HPV⁺HNC cell line correlated with

92 our *in vitro* observations and suggests that these drugs may be useful adjuvant approaches
93 for further investigation. To our knowledge, our analysis is the first to provide evidence
94 on the utility of SERMs for HPV⁺OPCs.

95 Overall, this report further expands upon the analysis of utilizing estrogen related
96 signaling in the quest for HPV-specific antiviral approaches²⁵. While estrogen presented
97 compelling evidence *in vitro*, this report demonstrates that estrogen treatment did not
98 translate to *in vivo* models²⁶. The development of co-culture models utilizing fibroblast
99 “feeder” cells demonstrated that these supporting fibroblasts altered the response to
100 estrogen *in vitro*, modeling what was observed *in vivo*. Analysis of SERMs with this co-
101 culture model demonstrated the utility of altering estrogen-related signaling both *in vitro*
102 and *in vivo*. Therefore, co-culture with fibroblasts offers a simple and more
103 physiologically relevant environment by stimulating more of the cellular interactions
104 present in solid tumors. This co-culture model may serve to better predict drug responses
105 in other translational paradigms, not limited to HPV⁺ cancers. Co-culture allows for an
106 examination of the complex cellular responses to drug effects *in vivo*, thereby enhancing
107 the accuracy of *in vitro* therapeutic evaluations for more successful translational
108 approaches.

109 **Results**

110 **Estrogen sensitization is not observed *in vivo***

111 We have previously reported that estrogen attenuates the growth of epithelial cells
112 in an HPV⁺ dependent manner *in vitro*²⁶. We found that this occurred via both a
113 repression of transcription from the HPV16 long control region and through interactions
114 with the viral oncogenes, E6 and E7²⁶. Furthermore, estrogen treatment enhanced

115 irradiation-induced cell death in an HPV⁺ dependent manner²⁶. A logical progression was
116 to assess the response of HPV⁺ cancers to estrogen *in vivo*. Previously, our laboratory as
117 well as others have demonstrated that HeLa cells are highly responsive to estrogen
118 treatment^{26,52,53}. Consequently, experiments were designed to assess the combination of
119 estrogen and radiation treatment on HeLa xenografts in female NSG mice. Contrary to
120 our *in vitro* data, Figure 1A shows that estrogen alone, or in combination with radiation,
121 had no impact on tumor response in mice. Of note, animal weights remained consistent
122 throughout the study with all treatments (Figure 1B).

123 Previously, we had observed HPV⁺ dependent cell death following estrogen
124 treatment regardless of sex, tissue of origin, or viral genome integration status²⁶. To
125 determine if any of these factors played a role *in vivo*, we decided to expand our analysis
126 to include the male, episomal, HPV16⁺ oropharyngeal cancer cell line UMSCC104⁵⁴. In
127 addition, we added “early” estrogen supplementation that began the same day as
128 xenografts were injected, to determine if a temporal relationship was essential to the
129 estrogen treatment response. As our previous radiation treatment using 10 Gy had some
130 off-target effects in our NSGs, radiation was reduced to 5 Gy in these studies. As
131 observed in Figure 1C (animal weight Figure 1D), estrogen again had no impact on tumor
132 response in any of these conditions. These results complement the previous observations
133 by the Lambert laboratory utilizing MmuPV1²⁷. The Lambert observations demonstrated
134 that estrogenic effects *in vivo* are reliant, at least in part, on estrogen’s suppression of the
135 host immune system in immunocompetent mice; whereas estrogenic enhancement of
136 disease progression were not observed in immunodeficient NSG mice²⁷. Our results
137 indicate that the lack of estrogenic alterations in disease progression may not be

138 papillomavirus species specific, but further studies are needed to confirm this
139 observation.

140 **Stroma alters HPV-specific estrogen growth response in keratinocyte models**

141 There are numerous differences when moving from *in vitro* to *in vivo* models. An
142 increasingly recognized component of *in vivo* responses is the adjacent stromal support
143 network, or “tumor microenvironment” (TME)^{33,55,56}. In regard to HPV, evidence also
144 supports a significant role for stroma during the viral lifecycle and HPV-induced disease
145 ^{33,39,41,42}. Fibroblasts are a key component of this stromal support and can significantly
146 alter cancer resistance and therapeutic responses³³⁻⁴². We already utilize feeder layers of
147 mitomycin C (MMC) growth-arrested murine 3T3-J2 fibroblasts (referred to as J2s
148 moving forward) in the immortalization and maintenance of our primary keratinocyte
149 derived HPV⁺ epithelial cultures. Thus, we applied the same approach to investigate
150 whether this co-culture system affects the response to estrogen. MMC inactivation is a
151 supported approach to arrest the proliferation of fibroblasts, allowing for the
152 establishment of a supportive feeder layer that maintains its ability to synthesize RNA
153 and protein and provide stromal regulation of neighboring cells of interest⁵⁷. This
154 approach is widely accepted as necessary for maintenance of the HPV genome, as well as
155 a necessary component of 3D models for HPV lifecycle analysis⁵⁸⁻⁶⁴. N/Tert-1 cells
156 (telomerase immortalized foreskin keratinocytes, HPV negative), as well as HFK+E6E7
157 (foreskin keratinocytes immortalized by the viral oncogenes only), and HFK⁺HPV16
158 (foreskin keratinocytes immortalized by the entire HPV16 genome, replicating as an
159 episome), were cultured in the presence or absence of J2 cells, and treated with 15µM
160 estrogen, or vehicle control²⁶. Compared to untreated, non-cocultured N/Tert-1 cells, no

161 significant alterations in growth rate were observed following estrogen treatment or co-
162 culture with J2s (Figure 2A). Conversely, estrogen significantly repressed the growth of
163 both the E6E7 and HPV16 immortalized cell lines in the absence of stromal support
164 (Figure 2B and 2C, respectively). The presence of stromal support fibroblasts
165 significantly rescued this growth suppression (Figure 2B and 2C); coculture with
166 fibroblasts mitigates the growth suppressive effects of estrogen, *in vitro*. This implies that
167 the presence of stromal support is responsible for the differences in response to estrogen
168 treatment that we observed between our *in vitro* and *in vivo* models (Figure 1)²⁶.

169 **Stroma alters HPV-specific estrogen growth response in cancer models**

170 We sought to expand our co-culture investigations to include cancer lines. A
171 hallmark of transformation in many cancer cell lines is anchorage independent growth
172 and loss of adherence in cell culture⁶⁵. To improve our cancer cell count analysis, we
173 developed a novel, quantifiable co-culture system. Using the NuLight lentivirus system
174 (Sartorius), we transduced our cells of interest with nuclear mKate2-red and developed
175 stable cell lines; this system enabled automated cell counting and could distinguish
176 between our cells of interest and the non-labeled J2s. An additional advantage of this
177 system was the ability to monitor cellular morphology and observe any co-culture
178 influences upon colony formation and cellular distribution. We sought to determine
179 whether J2s alter cancer cell line responses to estrogen, as observed in our keratinocyte
180 models (Figure 2). We utilized four cancer cell lines in our co-culture experiments: HN30
181 – an HPV⁻ head and neck cancer (p53wt), HeLa – an HPV18⁺ integrated cervical cancer
182 (p53wt), UMSCC47 – an HPV16⁺ integrated head and neck cancer (p53wt), and
183 UMSCC104 – an HPV16⁺ episomal head and neck cancer (p53wt). As HeLa cells are

184 highly sensitive to estrogen, 1.5 μ M estrogen was utilized in experiments with these cells;
185 all other cell lines were treated with 15 μ M as previously described^{26,52}. As previously
186 shown, estrogen treatment significantly repressed cell growth in an HPV⁺ dependent
187 manner (Figures 3A,C,E,F)²⁶. Conversely, estrogenic sensitivity was no longer observed
188 in HPV⁺ cancer cells grown with feeder cells (Figures 3B,D,F,H). This was similar to the
189 loss of estrogenic sensitivity in our HPV⁺ keratinocytes (Figure 2), and in our mouse
190 studies (Figure 1A, and 1C). Again, demonstrating that stromal support alters estrogenic
191 sensitivity of HPV⁺ cancer models.

192 To determine if this repression of estrogenic sensitization was specific to mouse
193 fibroblast support in co-culture, HeLa cells were grown in the presence or absence of
194 conditioned media collected from replicating 3T3-J2s cultures, or grown in co-culture
195 with MMC-inactivated human dermal mesenchymal fibroblasts (HDFMs). Again Figure
196 4A demonstrates that HeLa cells in monoculture undergo significant growth repression in
197 the presence of estrogen. Similarly, estrogenic growth repression was observed in the
198 presence of J2 conditioned media (Figure 4B). Co-culture with HDFMs rescued
199 estrogenic growth repression in HeLa cells (Figure 4C), similar to what was observed
200 with J2s (Figure 3B). HDFMs did not repress the growth of HeLa alone (Figure 4C),
201 contrary to the growth repression observed in the presence with J2 (Figure 3B). While we
202 do not currently understand the mechanism behind this altered growth potential with
203 alternative species' fibroblasts, we find it noteworthy that estrogenic sensitivity is lost
204 with both co-culture methods, but not by incubation with media, indicating that cell-cell
205 contact is required. Moreover, this loss of sensitivity is not dependent on fibroblast
206 growth alteration of cancer cells.

207 **Stroma does not alter the response to cisplatin in cancer models.**

208 To assess the predictiveness of our co-culture model, we found it important to
209 investigate whether stroma would alter the response to other chemotherapeutic
210 approaches. HeLa, UMSCC47, UMSCC104, and HN30 cells were therefore evaluated for
211 their responsiveness to cisplatin (Figure 5A,C,E,G). Fibroblasts did not rescue the growth
212 arrest observed in all cell lines (Figures 5B,D,F,H). Cisplatin is a well-established clinical
213 treatment paradigm for HPV⁺ and HPV⁻OPCs, cervical cancer, and many other cancers^{66–}
214 ⁷⁴. We predicted that fibroblasts would not be able to rescue cancer cells from this
215 conventional chemotherapeutic agent. We postulate that fibroblasts are unlikely to
216 change the response to most currently accepted cancer treatment modalities. Instead, we
217 suggest that this model may better predict which novel therapeutics may fail translational
218 approaches. This is something we are currently investigating, and we encourage others to
219 consider this approach as well when translating treatments from *in vitro* to *in vivo*.

220 **Stroma alters HPV long control region (LCR) transcriptional regulation**

221 We previously published that estrogen represses transcription of the HPV16 long
222 control region (LCR) both in our N/Tert-1 and C33a transcription models²⁶. This
223 transcriptional repression of the LCR, in turn, downregulated the expression of early viral
224 genes in our numerous HPV⁺ keratinocyte and HPV⁺ cancer models²⁶. Expanding upon
225 these previous observations, we further wanted to determine the impact of stromal
226 support on estrogenic HPV16-LCR transcriptional regulation. N/Tert-1 cells grown in the
227 presence of stromal support had significantly enhanced levels of HPV16-LCR
228 transcription and this transcriptional regulation was no longer significantly repressed by
229 estrogen (Figure 6A); this may be one of the many contributing factors involved in the

230 loss of estrogen sensitivity. ER α was also assessed, and stroma did not appear to alter
231 protein expression in N/Tert-1 cells (Figure 6B). These results suggest that stroma is
232 highly supportive of HPV16-LCR transcriptional regulation.

233 Fibroblasts are routinely utilized to support HPV genome maintenance and the
234 viral life cycle in keratinocyte models^{33,41,57,75-82}. While this model is accepted, the full
235 mechanism of how fibroblasts aid maintenance of the viral genome as an episome has yet
236 to be elucidated. Figure 6A demonstrates that stromal support enhances HPV16-LCR
237 transcription. It was therefore important to confirm that this transcriptional regulation had
238 downstream effects. Previously, elements of the HPV upstream regulatory region,
239 another term for the viral LCR, have been implicated as a requirement for long term viral
240 persistence in keratinocytes⁸³. The McBride laboratory demonstrated that the chromatin
241 architecture of this region is important for genome partitioning and may influence
242 integration⁸³. Figure 6C confirms that HFK $^+$ HPV16 begin to integrate the viral genome
243 when grown in the absence of J2 for one week^{84,85}. The accepted practice of fibroblast co-
244 culture, may in part maintain viral episomes via influencing the transcription of LCR.
245 Thus, stroma supports viral protein expression and alters host signaling pathways in
246 keratinocytes

247 We next investigated viral protein expression and host protein signaling observed
248 in the presence or absence of fibroblast support. HFK+E6/E7, HFK $^+$ HPV16, and N/Tert-
249 1 cells (Figure 6D) were grown in the presence or absence of J2s. J2s are washed off
250 before harvesting samples, and are murine-derived so there should be limited detection
251 via human-specific antibodies. To confirm that altered protein levels observed were not
252 due to any residual J2s, lane 1 in Figure 6D demonstrates that significant specific bands

253 were not observable with the majority of our chosen antibodies with 100 μ g of J2 protein
254 input. SIRT1 does have a notable lower band which is not unexpected due to the gene
255 homology between mouse and human SIRT1 and that the immunogen was developed
256 from the amino acids 1-131 of mouse Sir2 α . γ H2AX has a low-level detectible band due
257 to cross reactivity of the antibody; it should be noted that the significant observations
258 discussed have taken this into account. Figure 6D demonstrates that HFK+E6E7, which
259 do not rely on the LCR for early gene expression, do not have altered E7 protein levels in
260 the presence or absence of fibroblasts (lanes 2,3). As fibroblasts were shown to enhance
261 LCR transcription (Figure 6A), HFK $^+$ HPV16 cells, which rely on the LCR for early gene
262 expression, have enhanced E7 levels when grown in the presence of J2s (lanes 4,5).
263 HFK+E6E7 had limited levels of p53 expression, and HFK $^+$ HPV16 had low expression
264 of p53, due to E6-targeted degradation; however significant enhancement of p53 levels in
265 the presence of J2 in both cell lines were striking^{63,64,79,86-91}. To our knowledge, this is the
266 first report to suggest why p53 is not always fully degraded in HPV $^+$ cell lines; E6
267 degradation of p53 may depend on whether or not keratinocytes are maintained on feeder
268 cells^{79,88,90,92-97}. Upregulation of p53 in the presence of J2 was also observed in N/Tert-1
269 cells (lanes 6,7) so this observation is independent from E6 or full genome expression. It
270 is worth noting that HFK $^+$ HPV16 cells are consistently maintained in co-culture with J2s,
271 and that J2s were not supplemented for the “control” conditions. This also demonstrates
272 that the stromal induced alterations of p53 protein levels are reversible (Figure 6D). Total
273 levels of pRb were not altered N/Tert-1 cells nor in HFK+E6E7; this corresponds to
274 unaltered E7 levels in HFK+E6E7 cells. Alternatively, pRb was reduced in HFK $^+$ HPV16
275 grown in the presence of J2s. Again, suggesting fibroblasts are important for viral

276 regulation of keratinocyte signaling, possibly through the LCR. A consistent observation
277 was the upregulation of γ H2AX in the presence of J2s in all of the cell lines (Figure 6D).
278 Numerous reports have demonstrated HPV viral integrity and genome stability is highly
279 reliant on DNA repair machinery, including γ H2AX; J2 enhancement of γ H2AX may be
280 another key mechanism of fibroblast regulation of viral genome stability in keratinocyte
281 models^{7,76,98–106}. Upstream of p53 and γ H2AX, SIRT1 is known to target histones and
282 non-histone substrates such as p53, and has been shown to decrease in response to DNA
283 damage¹⁰⁷. N/Tert-1 and HFK⁺HPV16 grown in the presence of J2s demonstrated a
284 significant decrease in SIRT1. The lower band observed can be associated with SIRT1
285 post-translational modifications; while this was altered by fibroblasts in both HFK+E6E7
286 and HFK⁺HPV16, conclusions were not made due to the band observed with the J2 input
287 sample^{40,107–110}. While we are still investigating additional signaling mechanisms
288 involved, these observations highlight the importance of properly controlling for whether
289 or not cell lines are grown in the presence or absence of feeder cells when considering
290 viral and host signaling events.

291 **Selective estrogen receptor modulators reduce growth rates *in vitro* and *in vivo***

292 While estrogen had no impact on the tumor response in mice, alone or in
293 combination with radiation (Figure 1), we expanded our analysis to assess if the selective
294 estrogen receptor modulators (SERMs) raloxifene or tamoxifen would be a useful
295 approach to continue to take advantage of the HPV⁺ specific overexpression of ER α . It
296 should be noted that SERMs can be agonists or antagonists of the ER α , and these
297 responses are dependent on region, cell type, and the localization and availability of
298 estrogen response elements (EREs)^{44,46,47,111}. Preclinical data supports the utility of

299 SERMs for cervical cancer, particularly the utility of raloxifene on the reduction of
300 recurrent neoplastic disease; however, SERMs have yet to be evaluated in
301 HPV⁺OPC^{47,111}.

302 HeLa, UMSCC47, and UMSCC104 HPV⁺ cancer cells grown without fibroblast
303 support exhibited significant growth repression to both SERMs (Figures 7A,C,E). While
304 fibroblast support did not alter the response to SERMs in all the cell lines, rescued growth
305 was observed in HeLa cells treated with tamoxifen while remaining responsive to
306 raloxifene (Figures 7B,D,F). It is worth noting that previous *in vivo* and clinical analysis
307 have predicted the utility of raloxifene in HPV⁺ cervical cancer response, whereas
308 tamoxifen is not recommended^{45–47,51,111}. We therefore sought to analyze SERM
309 responsiveness in an HPV⁺HNC *in vivo* model. Expansion of SERM treatment *in vivo*
310 was therefore conducted in UMSCC104 cells alone or in combination with IR to assess
311 their utility as well as determine if our co-culture model could be useful in future
312 translational approaches. For this study, 4 Gy IR was chosen to further reduce the off-
313 target issues relating to radiation use in NSGs¹¹². As observed in Figure 8A, radiation
314 alone significantly reduced tumor volume starting on Day 14, while raloxifene (Figure
315 8A) and tamoxifen alone (Figure 8B) were able to significantly reduce tumor volume
316 starting on Day 32. Furthermore, in comparison to radiation (IR) alone, tamoxifen+IR
317 significantly reduced tumor volume starting on Day 42 (Figure 8B). Kaplan-Meier
318 survival analysis demonstrated 50% survival for tamoxifen+IR on day 70 in comparison
319 to 10% survival at the same time point for IR alone or raloxifene+IR at our chosen
320 endpoint (Figure 8C). With all treatments, animal weights remained consistent
321 throughout the study (Figure 8D). Our *in vitro* co-culture system modeled our *in vivo*

322 responses, as well as those previously observed, to both estrogen and SERMs in HPV⁺
323 cervical *in vivo* models (Figure 1,3,7) ^{27,32,43,45–47,51,111}. Co-culture also predicted the
324 responsiveness of SERMs in an HPV⁺HNC *in vivo* model (Figure 7,8) and suggests that
325 the utility of SERMs for the treatment of HPV⁺OPC are worth further investigation.
326 Altogether, we suggest that the predictiveness of this co-culture system should be
327 considered in more translational approaches. Current studies are investigating the
328 molecular mechanisms behind the alterations observed when cells are grown in fibroblast
329 co-culture. These mechanistic approaches may further expand the predictive utility of our
330 co-culture model.

331 **Discussion**

332 We previously established that estrogen attenuates the growth of HPV⁺
333 keratinocyte and cancer cell lines in both an LCR and E6E7 dependent manner^{25,26}. Of
334 note, when these studies were conducted, we did not supplement fibroblast support
335 during drug treatment²⁶. Conversely, when estrogen was utilized to treat HPV⁺ xenografts
336 in NSG mice, responsiveness was no longer observed (Figure 1). This data is supportive
337 of previous observations by the Lambert laboratory²⁷. We acknowledge the importance of
338 the immune system in the context of HPV, the responsiveness to estrogen, and its
339 potential impact on translational approaches as well^{27,28,45,113}. As NSG mice possess a
340 significantly compromised immune system, we then sought to investigate the role that
341 stromal support may play in the altered responsiveness to estrogen when moving from *in*
342 *vitro* to *in vivo* models. In doing so, we found that there is an HPV-specific change in
343 response to estrogen when comparing co-culture to non-fibroblast conditions (Figure
344 2,3,4). We find that this response is, at least in part, HPV16 LCR dependent; however,

345 there are likely other mechanisms at play (Figure 4). Additional mechanisms behind the
346 observed stromal growth alterations, as well as the response to therapeutics, are currently
347 under investigation in our laboratory and will be the subject of future reports.

348 Of note, the analysis of alternative species fibroblasts and mesenchymal cell types
349 on basement membrane arrangement and growth characteristics of organotypic raft
350 cultures were conducted many years ago¹¹⁴. Tissue-specific, species-specific, and spatial-
351 specific alterations were found to impact numerous epithelial phenotypes¹¹⁴. Extracellular
352 matrix components were found to have the greatest impact in this publication. We
353 propose that the differences in extracellular matrix components between mouse and
354 human might have moderate plating efficiency or growth efficiency alterations, and
355 contribute to the altered growth potential observed in HeLa cells (Figure 3,4). As seen in
356 Figure 3, UMSSCC47 cell growth was not impacted by J2s; alternatively, UMSSCC104 and
357 HN30 cells grew better in the presence of J2. Regardless of the altered growth potential
358 of cell lines in the presence or absence of fibroblasts, the observation that estrogenic
359 sensitivity was lost in co-culture remains the same.

360 Anecdotally, is it known in the HPV field that J2s are crucial to the culture of
361 primary keratinocytes for the maintenance of viral genome stability^{75,76,104}. Expanding the
362 accepted fibroblast co-culture system for HPV⁺ primary keratinocytes, our data
363 demonstrates that this model promotes translational utility. When keratinocytes or cancer
364 cells were grown in this co-culture system, *in vivo* results were more predictable. These
365 models allowed for us to translate our estrogen results into estrogen pathway targeting
366 drugs. Previous studies have highlighted the potential utility of SERMs in HPV⁺ cervical
367 cancers, most specifically the use of raloxifene to reduce neoplastic recurrence^{45,46,48,111}.

368 We have expanded SERM analysis from cervical cancer to oropharyngeal cancer. We
369 now demonstrate that SERMs may present clinical applicability as adjuvant approaches
370 in HPV⁺OPC and future investigations are warranted. While our observations apply both
371 to primary keratinocytes and cancer models of HPV, we are currently in collaboration to
372 expand analysis to other cancer treatment models. Nevertheless, stroma is recognized to
373 contribute to the response to therapeutics and the development of resistance, and we
374 conclude that it should be more often considered in future translational
375 approaches^{33,35,39,40,92}.

376 **Materials and Methods**

377 **Cell Culture**

378 HN30 (generous gift from Hisashi Harada, VCU Philips Institute), UMSCC47 (Millipore;
379 Burlington, MA, USA), and HeLa (generous gift from Alison McBride, NIAID) cells
380 were grown in Dulbecco's modified Eagle's medium (DMEM) (Invitrogen; Carlsbad,
381 CA, USA) supplemented with 10% charcoal/dextran stripped fetal bovine serum (Gemini
382 Bio-products; West Sacramento, CA, USA). UMSCC104 (Millipore) cells were grown in
383 Eagle's minimum essential medium (EMEM) (Invitrogen) supplemented with
384 nonessential amino acids (NEAA) (Gibco) and 20% charcoal/dextran stripped fetal
385 bovine serum. N/Tert-1 cells and all derived cell lines have been described previously
386 and were maintained in keratinocyte-serum free medium (K-SFM; Invitrogen),
387 supplemented with a 1% (vol/vol) PenStrep (Gibco) and previously described
388 antibiotics^{20–22,115–120}. HFK⁺HPV16 have been previously described and were grown in
389 Dermalife-K complete media (Lifeline Technology), and maintained on inactivated
390 fibroblast feeder cells (described below)¹²¹. HFK+E6/E7 were grown in Dermalife-K

391 complete media (Lifeline Technology), and maintained on inactivated fibroblast feeder
392 cells (described below); the immortalization process is described below. Of note, we have
393 no issues with fibroblast plating efficiency, or keratinocyte viral episome maintenance
394 utilizing these keratinocyte complete media kits over the traditional use of E-media^{121,122}.
395 For all cells not directly purchased from companies, the cell type was confirmed by Johns
396 Hopkins or MD Anderson cell line authentication services, and the cells were maintained
397 at 37°C in a 5% CO₂–95% air atmosphere, passaged every 3 or 4 days, and routinely
398 monitored for mycoplasma (Sigma, MP0035).

399 **Flank xenografts for *in vivo* drug trials**

400 HeLa and UMSCC104 cells were stably transduced with a lentiviral vector for pLX304
401 Luciferase-V5 blast (generous gift from Renfeng Li, originally obtained from Kevin
402 Janes, Addgene plasmid # 98580)¹²³. Cell lines were selected with 10µg/ml blasticidin.
403 Expression was verified with bioluminescent imaging, further outlined and defined in
404 transcriptional activity analysis method detailed below.

405 Xenografting was performed using previously described methodology, in collaboration
406 with the Virginia Commonwealth University (VCU) Cancer Mouse Models Core
407 Laboratory (CMMC)^{124,125}. All experiments were conducted in accordance with animal
408 protocol AD10002330 approved by VCU Institutional Animal Care and Use Committee.
409 NOD-SCID-IL2γ receptor null (NSG) mice (6-8 weeks old) were injected with 1x10⁶
410 cells suspended in PBS and Cultrex™ basement membrane extracts (BME) (Bio-
411 techne/R&D Systems) into the right flank. HeLa studies were conducted in female mice;
412 UMSCC104 studies were conducted in male mice (chosen to mimic sex from human
413 donors). Days 1-3 post-xenograft and at varying times throughout the studies,

414 bioluminescence imaging was performed using a Xenogen IVIS-100 system (Calipers
415 Life Sciences, Hopkinton, MA) to verify xenograft establishment, growth, and possible
416 metastasis based on previously established protocols¹²⁶. Tumor volume was also
417 measured on noted dates and calculated as $V = AB^2 (\pi/6)$, where A is the longest
418 dimension of the tumor, and B is the dimension of the tumor perpendicular to A. Data
419 points presented on the graph are representative for each condition while more than 70%
420 of the animals remained in each group. HeLa cells were palpable on day 10, UMSCC104
421 cells were palpable on day 7. It should be noted that in our hands, UMSCC104 xenografts
422 are prone to ulceration. Animals were humanely euthanized when ulcerations were
423 observed.

424 **NSG estrogen delivery**

425 A combinational approach for estrogen delivery was based on modified protocols
426 designed in collaboration with the source of our obtained control and estrogen beeswax
427 pellets (0.4mg estrogen, Huntsman Cancer Institute, University of Utah)^{127,128}. Estrogen
428 was also delivered in drinking water using a protocol kindly shared online by the Wicha
429 lab¹²⁹. Briefly, a 2.7 mg/mL stock of 17-estradiol (Sigma # E2758) in ethanol was diluted
430 to a final concentration of 8 µg/mL in sterile drinking water. Pellets (control or estrogen)
431 were implanted after tumors were palpable. Drinking water supplementation also began
432 at this time point. “Early estrogen” in UMSCC104 denotes that water estrogen
433 supplementation began the same day as xenograft injections (pellets were still implanted
434 when tumors became palpable on day 7).

435 **NSG SERM delivery**

436 Treatment of mice with Raloxifene was performed as previously described by the
437 Lambert Laboratory^{45,111}. The human formulation of raloxifene hydrochloride (60mg
438 tablets; EVISTA; Eli Lilly) were purchased from Virginia Commonwealth University
439 Health System Pharmacy. Tablets were resuspended in PBS for a final concentration of
440 10 mg/ml. Mice were administered a 150 μ l drug suspension (equivalent to 1.5 mg) via
441 i.p. injection. Treatment of Tamoxifen was performed as previously described¹³⁰.
442 Tamoxifen (Sigma, T-5648) was resuspended in corn oil (Sigma C-8267) at 37°C for a
443 final concentration of 10 mg/ml. Mice were administered 100 μ l drug suspension
444 (equivalent to 1.0 mg) via i.p. injection. SERM treatments began after tumors were
445 palpable. Mice received treatment 5 days a week for 4 weeks, for a total of 20 injections.

446 **Small Animal Radiation Research Platform (SARRP) ionizing radiation delivery**

447 1 day following pellet implantation to allow for mouse recovery, targeted ionizing
448 radiation (IR) was delivered utilizing the Xstrahl SARRP. HeLa studies utilized 10 Gy.
449 Due to the radiation sensitivity of NSG mice, UMSCC104 studies reduced IR dose to 5
450 Gy, and finally to 4 Gy.

451 **Culture, conditioned media collection, and mitomycin C (MMC) inactivation of**
452 **3T3-J2 mouse feeder cells**

453 3T3-J2 immortalized mouse fibroblasts (J2) were grown in DMEM and supplemented
454 with 10% FBS. Fresh media was exchanged twice a week; conditioned media was spun
455 down as 500 rcf to remove any residual cells^{131,132}. 80-90% confluent plates were
456 supplemented with 4 μ g/ml of MMC in DMSO (Cell Signaling Technology) for 4-6 hours
457 at 37°C. MMC-supplemented medium was removed and cells were washed with 1xPBS.
458 Cells were trypsinized, spun down at 500 rcf for 5 mins, washed once with 1xPBS, spun

459 again, and resuspended. Quality control of inactivation (lack of proliferation) was
460 monitored for each new batch of mitomycin-C. 100-mm plate conditions were
461 supplemented with 1×10^6 J2 and 6-well plate conditions were supplemented with 1×10^5
462 every 2-3 days; for longer term cultures, any remaining J2s were washed off with 1x PBS
463 and new J2 were continually supplemented every 2-3 days.

464 **Culture and mitomycin C (MMC) inactivation of human dermal mesenchymal
465 fibroblast feeder cells**

466 Human dermal mesenchymal fibroblasts (HDFM) were grown, treated, and quality
467 controlled as described for the above J2 protocol. 6-well plate conditions were
468 supplemented with 1×10^5 every 2-3 days.

469 **Generation of E6E7-immortalized keratinocytes**

470 Primary keratinocytes from single donors were obtained from LifeLine Cell
471 Technologies¹²¹. Cells were cultured on collagen-coated plates for lentiviral delivery of
472 HPV16 E6E7, using the pLXSN16E6E7 plasmid (Addgene plasmid # 52394, a gift from
473 Denise Galloway)¹³³. Following selection with G418 (72mM), cells were cultured on
474 mitomycin-C inactivated 3T3-J2 fibroblasts. HFKs were cultured in DermaLife-K
475 Complete media (LifeLine Cell Technologies) and E6E7 expression was confirmed by
476 qRT-PCR. Of note, these cells were generated at the same time and utilizing the same
477 donors as the previously described HFK-HPV16¹²¹.

478 **Co-culture of keratinocytes in the presence or absence of inactivated fibroblasts**

479 N/Tert-1, HFK E6E7, or HFK HPV16 cells were seeded at 5×10^5 in 100-mm plates in the
480 presence or absence of previously seeded J2s (at least 6 hours prior). Twenty-four hours
481 later, noted cells were supplemented with 15 μ M 17 β -estradiol (estrogen). Forty-eight

482 hours after estrogen supplementation, plates were washed to remove residual J2 and cells
483 were trypsinized and counted. For analysis of 1 week time point, HFK E6E7 or HFK
484 HPV16 cells were seeded at 1×10^5 in 100-mm plates in the presence or absence of
485 previously seeded J2s. J2s were re-supplemented every 2-3 days as previously described.
486 Pellets from these experiments were utilized for subsequent immunoblotting or DNA
487 analysis, detailed below.

488 **Generation of stable nuclear labeled cells with Incucyte® Nuclight Lentivirus**

489 mKate2 Incucyte® Nuclight Lentivirus (puro) cells were generated according to the
490 Sartorius product guide protocol (Sartorius cat# 4476), using a MOI of 3 or 6, depending
491 on cell type. Cells generated were maintained in 1 μ g/ml puromycin supplemented media.
492 Fluorescence was routinely monitored by BZ-X TexasRed filter via the Keyence BZ-
493 X800 inverted fluorescence microscope.

494 **Co-culture of nuclear-labeled cancer cell lines in the presence or absence of
495 inactivated fibroblasts**

496 Stable mKate2-puro HeLa, UMSCC47, UMSCC104, or HN30 cells were seeded in
497 triplicate at 1×10^4 per well in 6-well plates in the presence or absence of previously
498 seeded J2 cells, or HDFM cells (at least 6 hours prior – fibroblast type is noted for each
499 experiment). Twenty-four hours after seeding, day 0 images were captured in brightfield
500 and TexasRed with the Keyence BZ-X800 Image Cytometer. Noted drugs were
501 supplemented immediately after this initial imaging: 1.5 μ M (HeLa) or 15 μ M 17 β -
502 estradiol (Sigma), 10 μ M Tamoxifen (MP Biomedicals), 10 μ M Raloxifene (Cayman
503 Chemical Company), or 10-20 μ M Cisplatin (APExBIO). Cytometry images were again
504 captured on day 1 and day 3. UMSCC104 were washed on day 3 after imaging, new J2

505 and drug were supplemented, and additional images were captured on day 5 and 7. Ten
506 fields of view were randomized per well for all conditions. Cell count image cytometry
507 batch analysis was performed using the Keyence BZ-X800 Image Analyzer software. All
508 conditions utilized set analysis conditions from a single randomized control image and
509 applied to all data points automatically to reduce variability and bias. Data is presented as
510 fold of control from day 0. Representative images are presented in Figure 9.

511 **Immunoblotting**

512 Specified cells were washed with 1X PBS and trypsinized. Pellets were washed with 1X
513 PBS and resuspended in 5X packed cell volume of NP40 buffer (50mM Tris-HCl Ph 7.5,
514 150mM NaCl, 1%NP-40, 5mM EDTA) supplemented with Roche cOmplete protease
515 inhibitor and Roche PhosSTOP phosphatase inhibitor. Cell-lysis buffer suspension was
516 incubated on ice for 30 min with occasional agitation, then centrifuged for 15 min at
517 14,000 rcf at 4 °C. Supernatant protein concentration was measured via the Bio-Rad
518 protein estimation assay according to manufacturer's instructions. 100 µg protein samples
519 were heated at 95 °C in 4x Laemmli sample buffer (Bio-Rad) for 5 min. Noted samples
520 were run down a Novex 4–12% Tris-glycine gel (Invitrogen) and transferred onto
521 a nitrocellulose membrane (Bio-Rad) at 30V overnight, or 100V for 1 hour using the wet-
522 blot transfer method. Membranes were blocked with Odyssey (PBS) blocking buffer
523 (diluted 1:1 with 1X PBS) at room temperature for 1 hour and probed with indicated
524 primary antibody diluted in Odyssey blocking buffer. Membranes were washed with PBS
525 supplemented with 0.1% Tween (PBS-Tween) and probed with the indicated Odyssey
526 secondary antibody 1:10,000 (goat anti-mouse IRdye 800CW or goat anti-rabbit IRdye
527 680CW) diluted in Odyssey blocking buffer. Membranes were washed three times with

528 PBS-Tween and an additional wash with 1X PBS. Infrared imaging of the blot was
529 performed using the Odyssey CLx Li-Cor imaging system. Immunoblots were quantified
530 using ImageJ utilizing GAPDH as the internal loading control. The following primary
531 antibodies were used for immunoblotting in this study at 1:1000, unless otherwise noted:
532 ER α (Abcam, ab32063), GAPDH 1:10,000 (Santa Cruz, sc-47724), pRb (Santa Cruz, sc-
533 102), p53 (Cell Signaling Technology, CST-2527 and CST-1C12), γ H2AX 1:500 (Cell
534 Signaling Technology, CST-80312 and CST-20E3), SIRT1 (EMD Millipore 07-131), β -
535 actin (Santa Cruz, sc-47778), E7 1:500 (Santa Cruz, sc-6981).

536 **Transfection and transcriptional activity analysis.**

537 N/Tert-1 cells were plated at a density of 5×10^5 in 100-mm dishes. The following day,
538 the previously described plasmids for pGL3 basic, pGL3 control, or pHPV16-LCR-Luc
539 were transfected Lipofectamine 2000 (according to the manufacturer's instructions,
540 ThermoFisher Scientific). Twenty-four hours after transfection, cells were washed, and
541 noted cells were supplemented with 15 μ M 17 β -estradiol; J2s were also supplemented at
542 this time point for noted conditions to reduce likelihood of altered transfection efficiency.
543 Forty-eight hours after transfection, cells were harvested utilizing Promega reporter lysis
544 buffer and analyzed for luciferase using the Promega luciferase assay system.
545 Concentrations were normalized to protein levels, as measured by the Bio-Rad protein
546 assay dye. Relative fluorescence units (RFU) were measured using the BioTek Synergy
547 H1 hybrid reader.

548 **Exonuclease V assay**

549 PCR based analysis of viral genome status was performed using methods described by
550 Myers *et al.*⁸⁴. Briefly, 20 ng genomic DNA was either treated with exonuclease V

551 (RecBCD, NEB), in a total volume of 30 μ l, or left untreated for 1 hour at 37°C followed
552 by heat inactivation at 95°C for 10 minutes. 2 ng of digested/undigested DNA was then
553 quantified by real time PCR using a 7500 FAST Applied Biosystems thermocycler with
554 SYBR Green PCR Master Mix (Applied Biosystems) and 100 nM of primer in a 20 μ l
555 reaction. Nuclease free water was used in place of the template for a negative control.
556 The following cycling conditions were used: 50°C for 2 minutes, 95°C for 10 minutes, 40
557 cycles at 95°C for 15 seconds, and a dissociation stage of 95°C for 15 seconds, 60°C for
558 1 minute, 95°C for 15 seconds, and 60°C for 15 seconds. Separate PCR reactions were
559 performed to amplify HPV16 E6 F: 5'- TTGCTTTCGGGATTTATGC-3' R: 5'-
560 CAGGACACAGTGGCTTTGA-3', HPV16 E2 F: 5'-
561 TGGAAGTGCAGTTGATGGA-3' R: 5'- CCGCATGAACTTCCCATACT-3', human
562 mitochondrial DNA F: 5'-CAGGAGTAGGAGAGAGGGAGGTAAG-3' R: 5'-
563 TACCCATCATAATCGGAGGCTTG -3', and human GAPDH DNA F: 5'-
564 GGAGCGAGATCCCTCCAAAAT-3' R: 5'- GGCTGTTGTCATACTTCTCATGG-3'

565 **Reproducibility, research integrity, and statistical analysis**

566 All *in vitro* experiments were carried out at least in triplicate in all of the cell lines
567 indicated. All cell lines were bought directly from sources indicated, or typed via cell line
568 authentication services. All images shown are representatives from triplicate experiments.
569 *In vivo* experiments were designed in collaboration with the VCU Massey Cancer Center
570 animal core and biostats core for sample size justification and statistical power analysis.
571 Quantification is presented as mean +/- standard error (SE). Student's t-test or analysis of
572 variance (ANOVA) were used to determine significance as appropriate: * p<0.05,
573 **p<0.01, ***p<0.001

574 **Data availability**

575 Following the 2023 NIH data management and sharing policy, all data resulting from the
576 development of projects will be available in scientific communications presented at
577 conferences and in manuscripts that will be published in peer-reviewed scientific
578 journals. Data will be deposited in the Open Science Framework (OSF) platform. OSF
579 can be accessed at <https://osf.io>. VCU is an OSF institutional member, and OSF is an
580 approved generalist repository for the 2023 NIH data management and sharing policy.

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939
940
941 **Figure Legends Figure**

942 **Figure 1: Estrogen fails to sensitize *in vivo*.**

943 1A. HeLa cells are an integrated HPV18+ female cervical cancer cell line, we therefore
944 chose to utilize female NOD-scid IL2Rgnull (NSG) mice for this treatment paradigm; we
945 did not choose to ovariectomize these mice. Day 0 marks the date at which cells were
946 injected for xenografts. Tumors were palpable on day 10, mice were randomized.
947 Estrogen alone (E=0.4mg pellet + 8 µg/ml water supplementation *ad libitum*), radiation
948 alone (IR=10 Gy), as well as the combinational approach (E+IR) were monitored for
949 effects on tumor volume by calipers. 1B. Mice were monitored for weight throughout the
950 study. 1C. UMSSC104 cells are an episomal HPV16+ male oropharyngeal line, we

951 therefore chose to utilize male NSG mice for this treatment paradigm. Day 0 marks the
952 date at which cells were injected for xenografts; early estrogen water supplementation
953 began on this day, pellets were later added when tumors were palpable (8 μ g/ml water
954 supplementation *ad libitum*). Tumors were palpable on day 7. Estrogen alone (0.4mg
955 pellet + 8 μ g/ml water supplementation *ad libitum*), radiation alone (5 Gy), as well as the
956 combinational approach were monitored for effects on tumor volume by calipers. 1D.
957 Mice were monitored for weight throughout the study.

958

959 **Figure 2: Fibroblasts significantly reduce HPV-specific estrogenic sensitization in**
960 **keratinocytes.** (A) N/Tert-1, (B) HFK+E6E7, and (C) HFK+HPV16 cells were seeded
961 on day 0 and grown in the presence or absence of J2s that had been seeded at least 6
962 hours prior. Cells were washed to remove J2s in noted conditions, trypsinized, and
963 counted on day 3. *, p < 0.05; ***, p < 0.001.

964

965 **Figure 3: Fibroblasts significantly reduce HPV-specific cancer estrogenic**
966 **sensitization.** J2s were seeded in the morning and noted nuclear-labeled cancer cells
967 were seeded at least 6 hours after: HeLa (A,B), UMSCC47 (C,D), UMSCC104 (E,F),
968 HN30 (G,H). Co-culture images for quantitation were taken the following morning and
969 are set at day 0, estrogen (E) was added immediately after initial imaging on day 0. Cells
970 were again imaged at day 1 and day 3. UMSCC104 cells were grown for an additional
971 time point; these were replenished with new J2s and estrogen on day 3 (post-imaging)
972 and day 5 and imaged again on day 7. Within same graphs *p<0.05, **p<0.001. J2s
973 altered growth rates for some of the cell lines and graphs are presented as separate for

974 visual simplicity, but experiments were run concurrently; comparing top and bottom
975 graphs $p<0.05$ J2 increased growth, $&p<0.05$ J2 decreased growth.

976

977 **Figure 4: Human fibroblasts (HDFM) significantly reduce HPV-specific cancer**
978 **estrogenic sensitization.** HeLa were seeded into standard media (A), J2 conditioned
979 media (B), or HDFMs were seeded in the morning and nuclear-labeled HeLa cells were
980 seeded at least 6 hours after (C). Co-culture images for quantitation were taken the
981 following morning and are set at day 0, estrogen (E) was added immediately after initial
982 imaging on day 0. Cells were again imaged at day 1 and day 3. $*p<0.05$

983

984 **Figure 5: Fibroblasts do not alter cisplatin sensitivity.** J2s were seeded in the morning
985 and noted nuclear-labeled cancer cells were seeded at least 6 hours after: HeLa (A,B),
986 UMSCC47 (C,D), UMSCC104 (E,F), HN30 (G,H). Co-culture images for quantitation
987 were taken the following morning and are set at day 0, noted concentrations of cisplatin
988 were added immediately after initial imaging on day 0. Cells were again imaged at day 1
989 and day 2. New J2s or media were replenished on day 3, and imaged again on day
990 5. Within same graphs $****p<0.001$. J2s altered growth rates for some of the cell lines
991 and graphs are presented as separate for visual simplicity, but experiments were run
992 concurrently; comparing top and bottom graphs $p<0.05$ J2 increased growth, $&p<0.05$
993 J2 decreased growth.

994

995 **Figure 6: Stroma supports transcriptional regulation, viral protein expression, and**
996 **episomal maintenance; estrogenic transcriptional regulation is lost with stroma.** (A)

997 N/Tert-1 cells were transfected with 1 μ g of pgl3 basic backbone, 1 μ g of pgl3 control
998 (positive control), or 1 μ g LCR and grown in the presence or absence of 15 μ M estrogen
999 and/or J2s that had previously been seeded. Forty-eight hours after transfection, a
1000 luciferase-based assay was utilized to monitor levels of LCR transcription. Data are
1001 presented as relative fluorescence units (RFU), normalized to total protein concentration
1002 as monitored by a standard bovine serum albumin (BSA) assay. ANOVA **, $P < 0.01$;
1003 ***, $P < 0.001$ (B) N/Tert-1 cells were grown in the presence or absence of J2s. Cells
1004 were washed to remove J2, then lysed and analyzed via western blotting for ER α .
1005 GAPDH was used as a loading control. (C) HFK $^+$ HPV16 cells were seeded on day 0 and
1006 grown in the presence or absence of J2s for 1 week. Cells were washed to removed J2,
1007 then lysed and analyzed for DNA expression of E2 and E6 via the exonuclease V assay,
1008 in comparison to GAPDH and mitochondrial DNA controls. Results are presented as
1009 percent integration as calculated from the cut ratio of matched GAPDH. * $P < 0.05$. (D)
1010 HFK+E6E7 (lanes 2,3) and HFK $^+$ HPV16 (lanes 4,5) and N/Tert-1 (lanes 6,7) cells were
1011 seeded on day 0 and grown in the presence or absence of J2s for 1 week. Lane 1 is an
1012 input lysate from J2 alone to control for any background level of expression in fibroblasts
1013 that were not removed via washing. Cells were washed to remove J2s in noted conditions,
1014 trypsinized, lysed, and analyzed via western blotting for SIRT1, pRb, p53, γ H2aX, and
1015 E7. GAPDH and β -actin were utilized as loading controls.

1016
1017 **Figure 7: Fibroblast co-culture demonstrates SERMs are worth assessing in an**
1018 **HPV+ *in vivo* model.** J2s were seeded in the morning and noted nuclear-labeled cancer
1019 cells were seeded at least 6 hours after: HeLa (A,B), UMSCC47 (C,D), UMSCC104

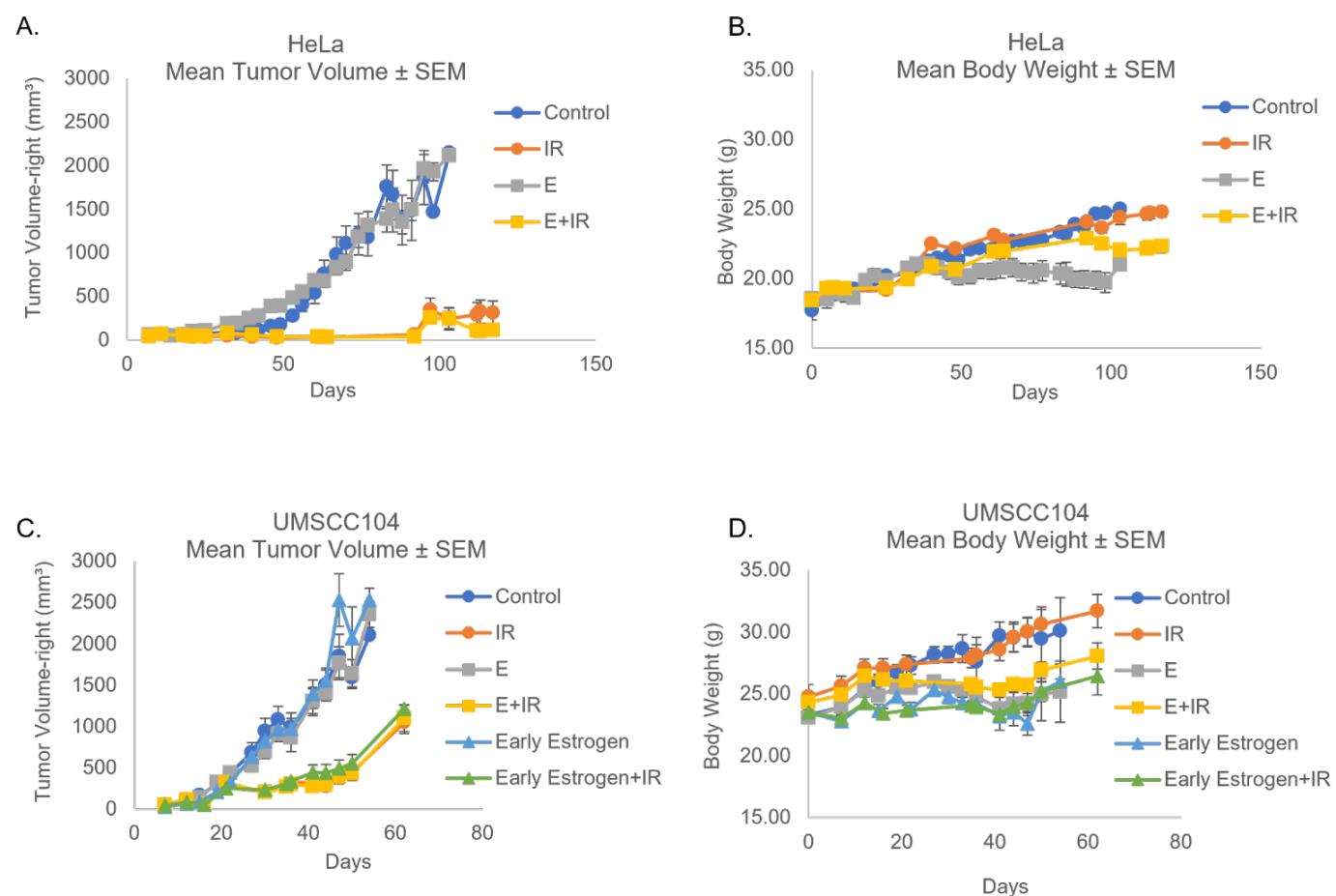
1020 (E,F). Co-culture images for quantitation were taken the following morning and are set at
1021 day 0, 10 μ g raloxifene (R) or 10 μ g tamoxifen (T) were added immediately after initial
1022 imaging on day 0. Cells were again imaged at day 1 and day 3. UMSCC104 cells were
1023 grown for an additional time point; these were replenished with new J2s and drugs on day
1024 3 (post-imaging) and day 5 and imaged again on day 7. Within same graphs *p<0.05,
1025 ***p<0.001. J2s altered growth rates for some of the cell lines and graphs are presented
1026 as separate for visual simplicity, but experiments were run concurrently.

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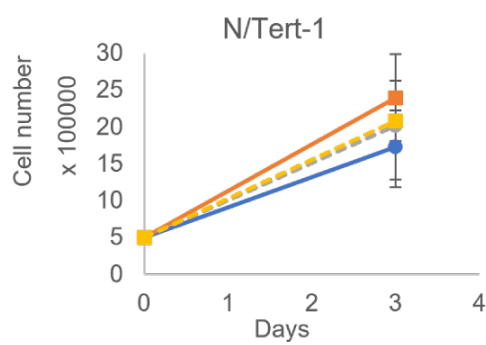
1028 **Figure 8: Fibroblast *in vitro* co-culture SERM assessment in UMSCC104 cells**
1029 **predicted the utility *in vivo*.** As described in Figure 1, UMSCC104 cells were injected
1030 for xenografts in male NOD-scid IL2Rgnull (NSG) mice. Day 0 marks the date at which
1031 cells were injected for xenografts. (A,B) Tumors were palpable on day 7. 1-Control, 2-
1032 Raloxifene alone (1.5 mg), 3-Tamoxifen alone (1.0 mg), 4-radiation alone (4 Gy) (IR), as
1033 well as the combinational approaches (5-Raloxifene+IR, 6-Tamoxifen+IR) were
1034 monitored for effects on tumor volume by calipers. These experiments were run
1035 concurrently but presented on separate graphs for visual clarity (groups have been
1036 numbered for this clarity). Data points shown for Mean Tumor Volume are representative
1037 of conditions with at least 70% of animals remaining. (C) Kaplan-Meier Survival plots
1038 the survival curve of the animals treated. This data includes any mice that needed to be
1039 sacrificed due to tumor ulceration. (D) Mice were monitored for weight throughout the
1040 study. *p<0.05, **p<0.01 ***p<0.001 from control, \$p<0.05 from IR alone, colors of *
1041 or \$ are matched to colors of corresponding conditions that are significant from control or
1042 radiation alone, respectively.

1043

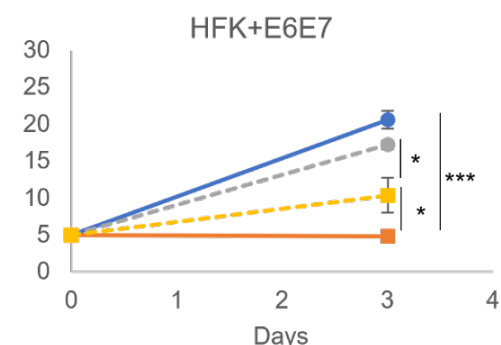
1044 **Figure 9. Representative cell culture images quantified via Keyence.** Images
1045 presented are day 5 of UMSCC104 cell cultures in the presence or absence of J2s. Images
1046 were captured at 10x in brightfield and TexasRed. All cell lines at all time points were
1047 counted utilizing the same control-established automated parameters to ensure
1048 reproducibility.



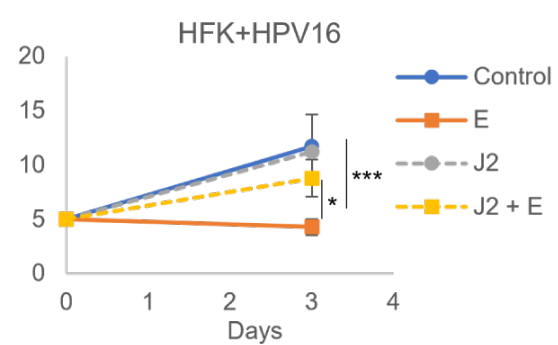
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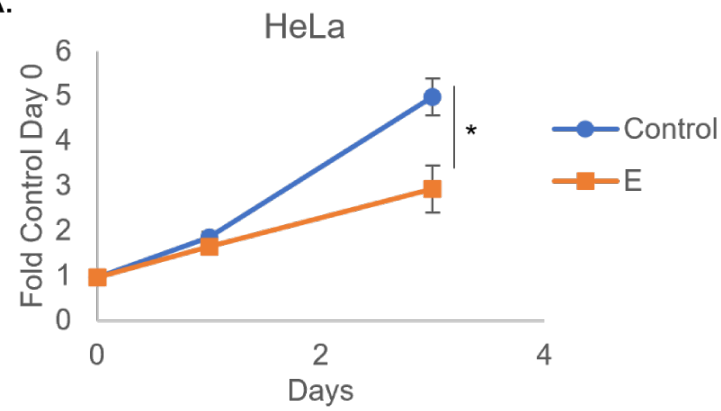
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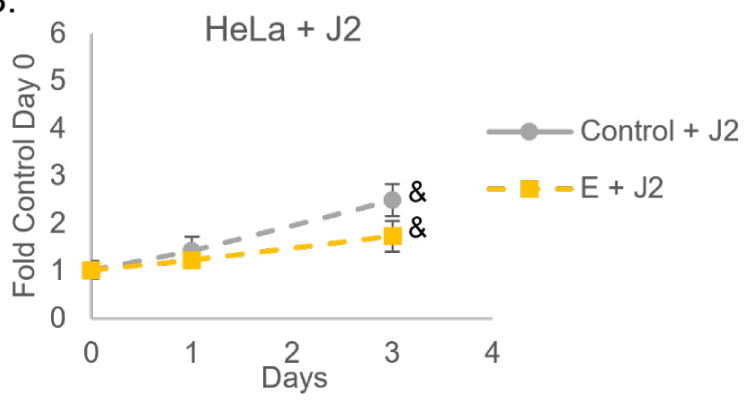
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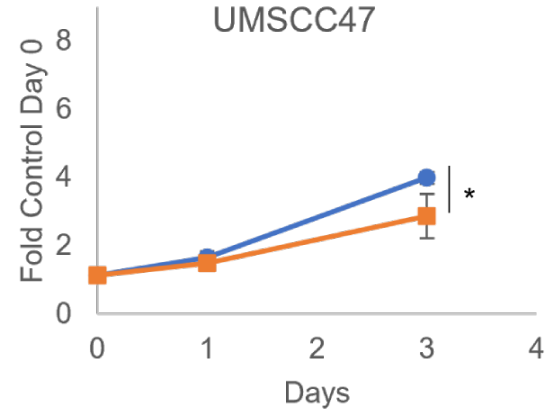
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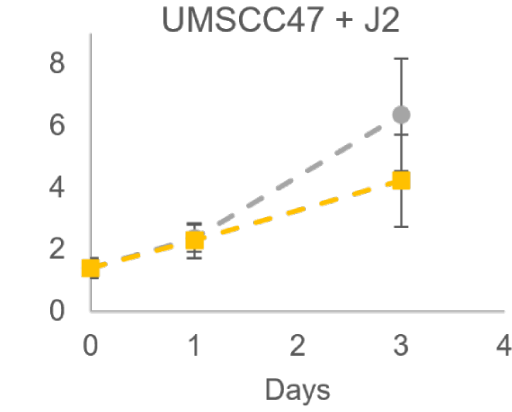
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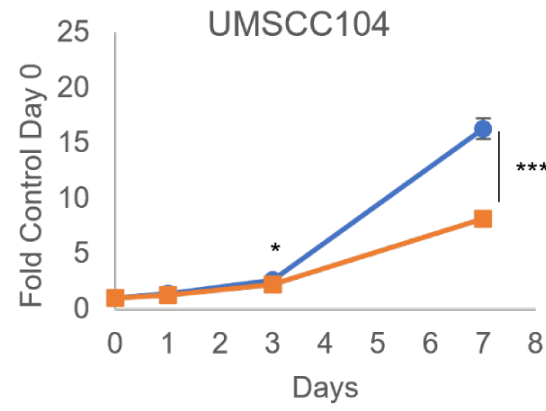
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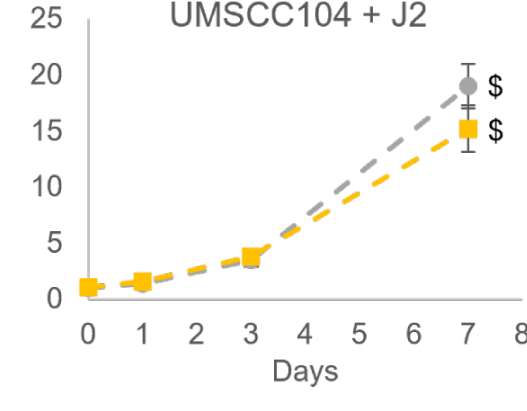
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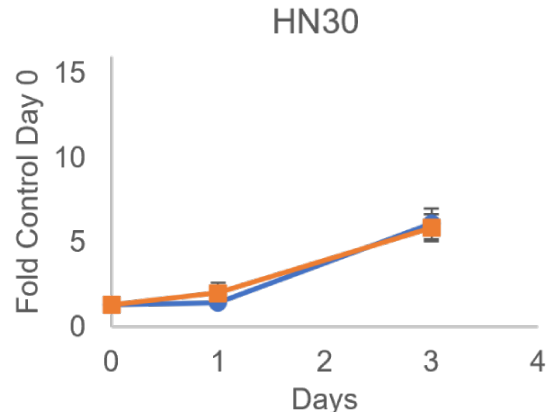
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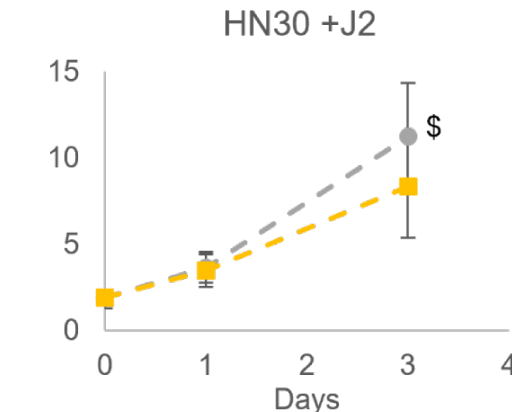
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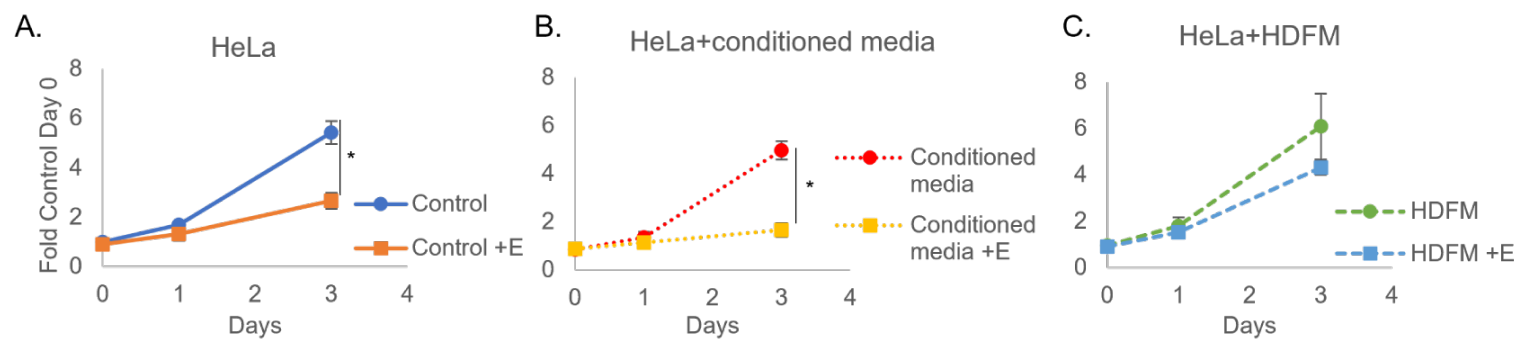


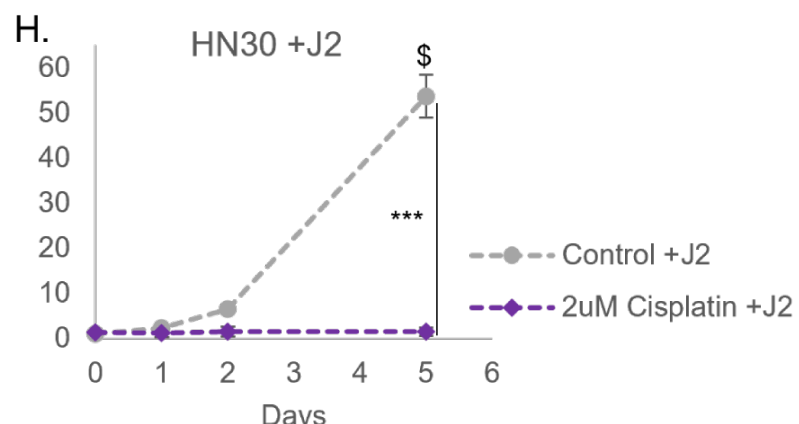
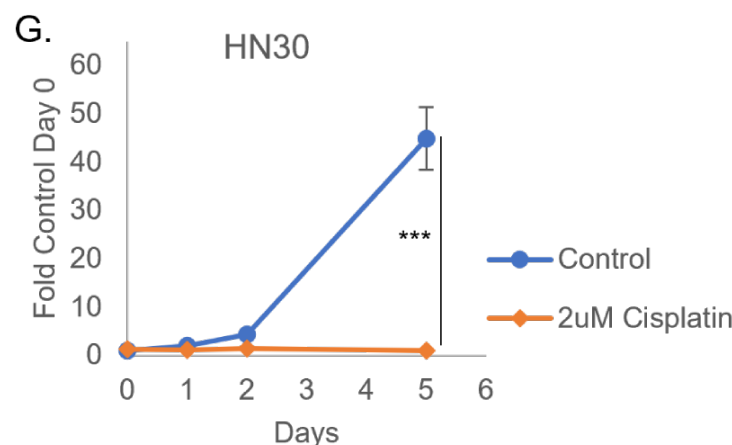
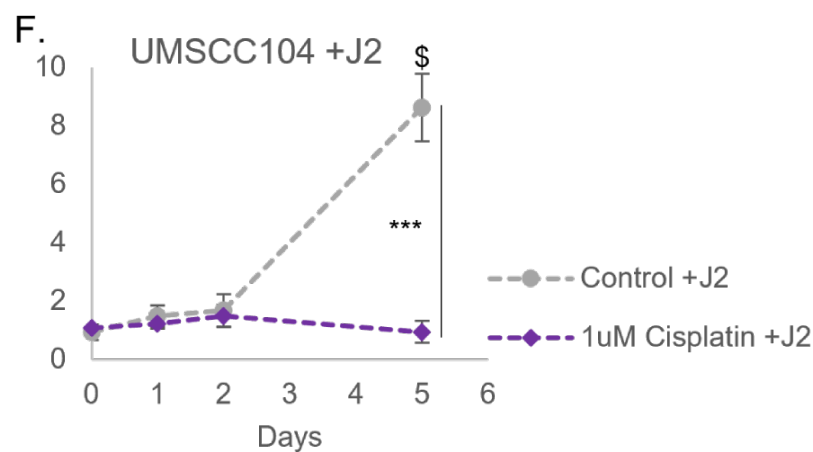
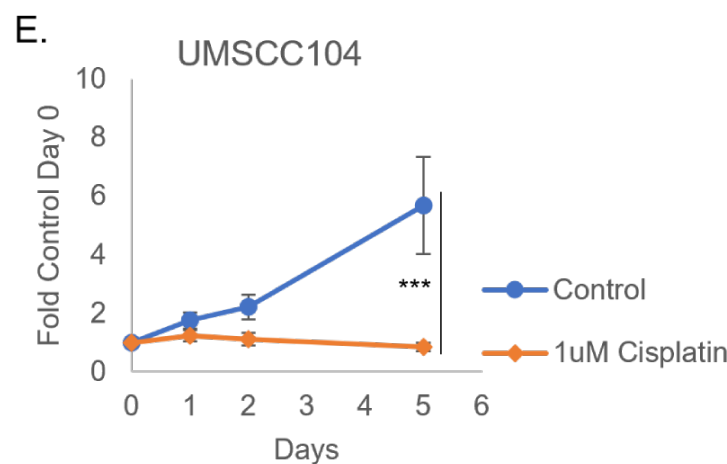
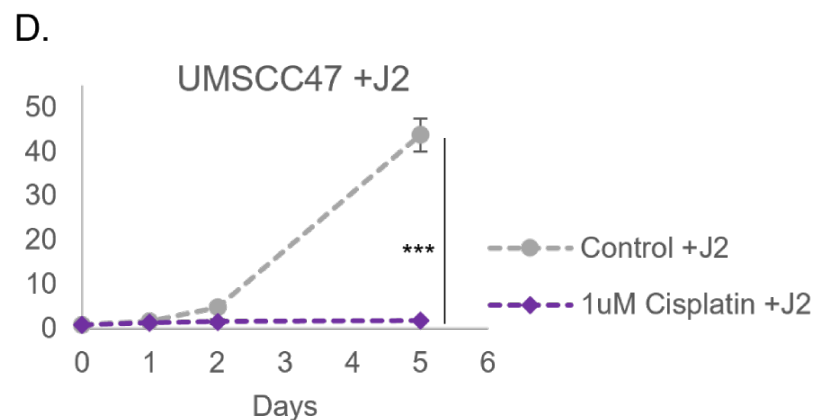
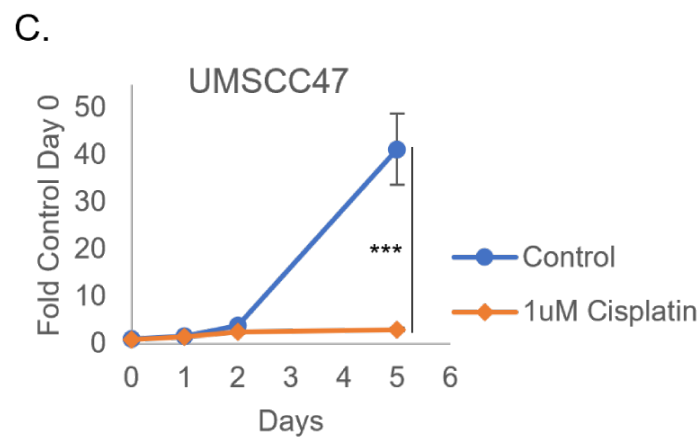
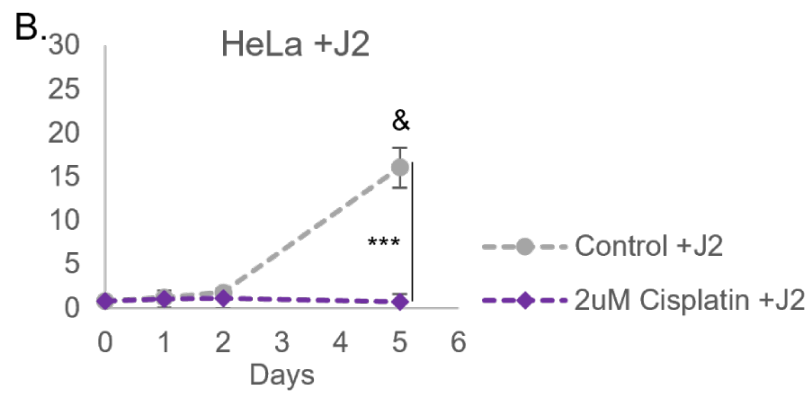
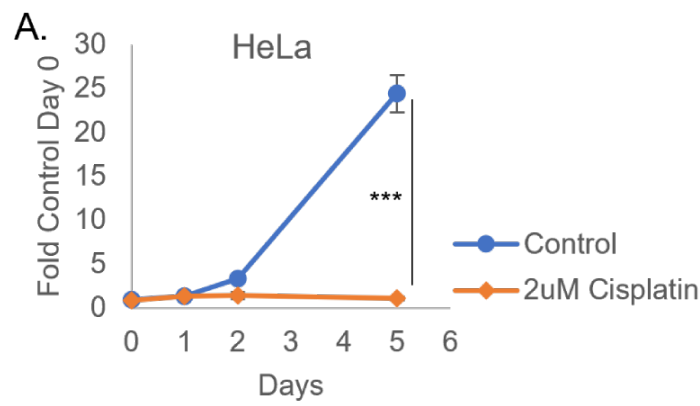
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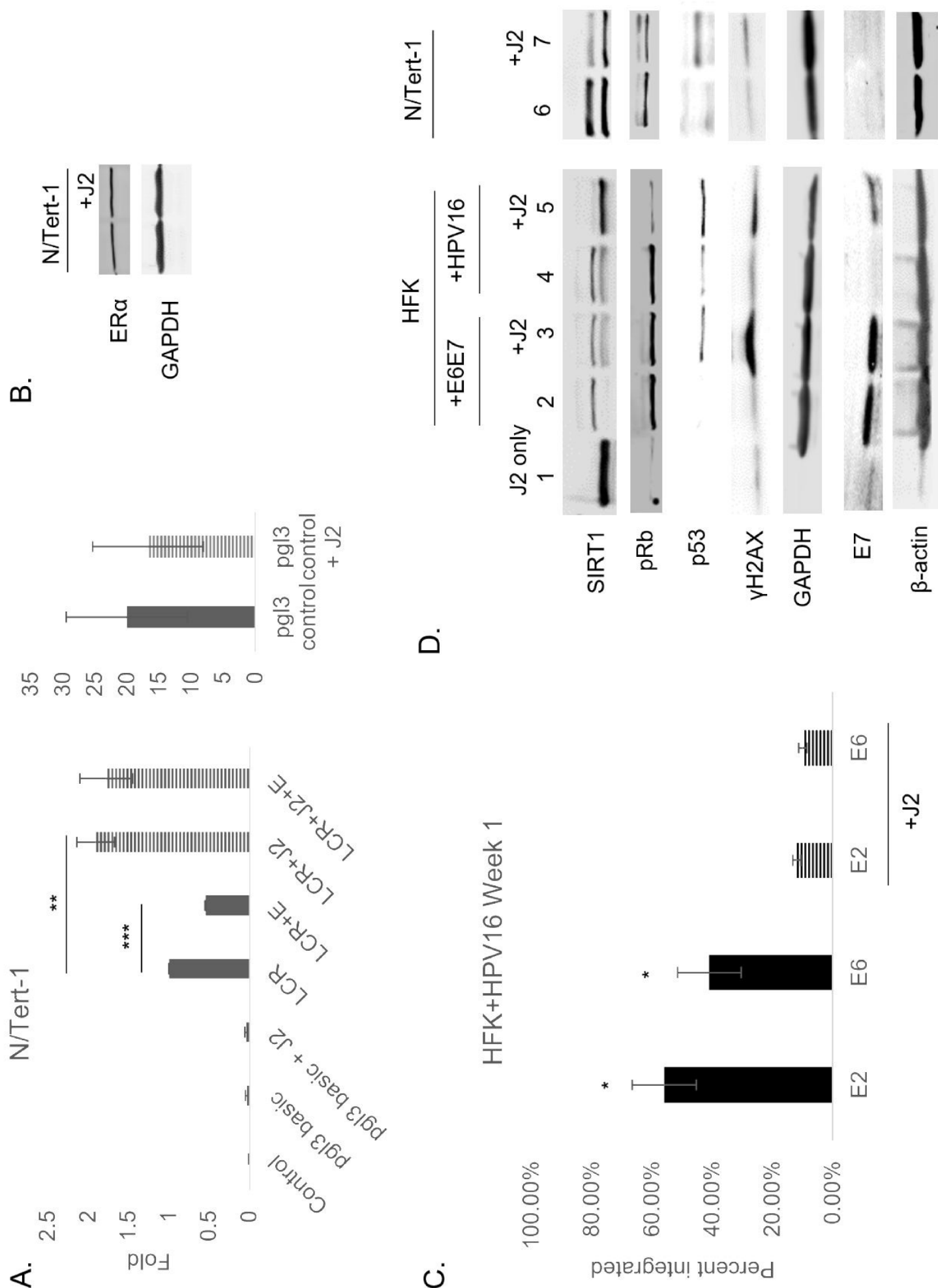


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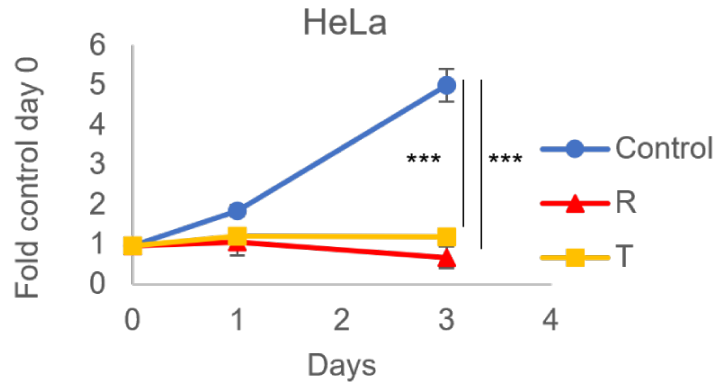




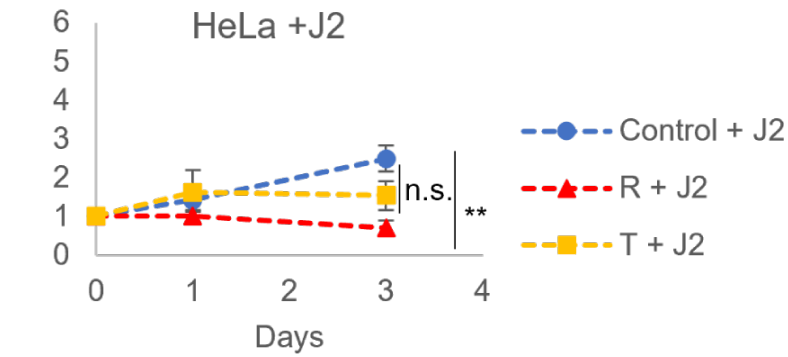




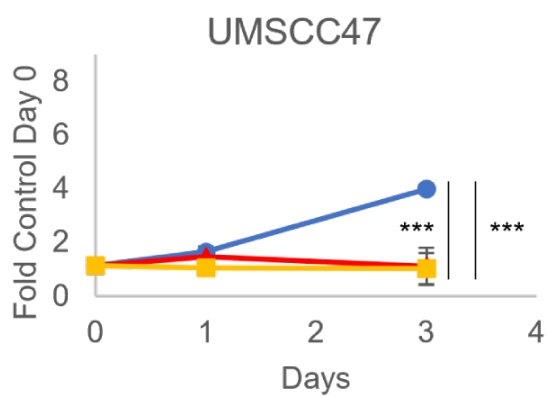
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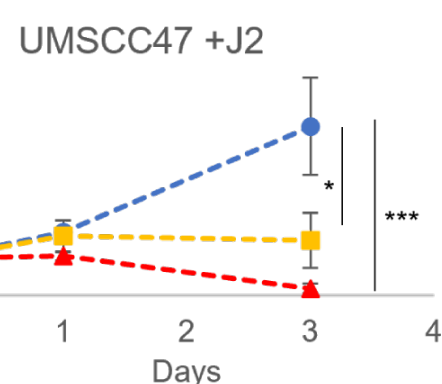
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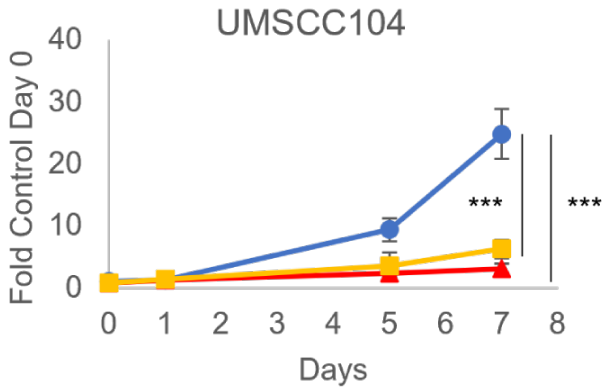
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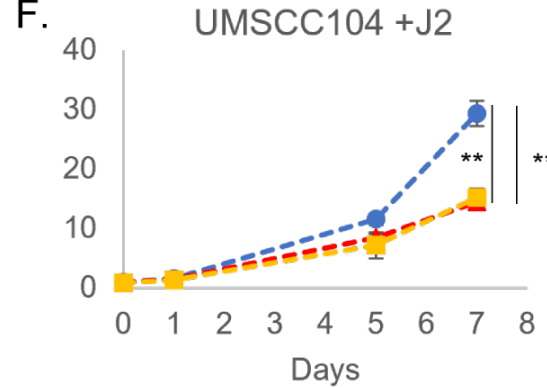
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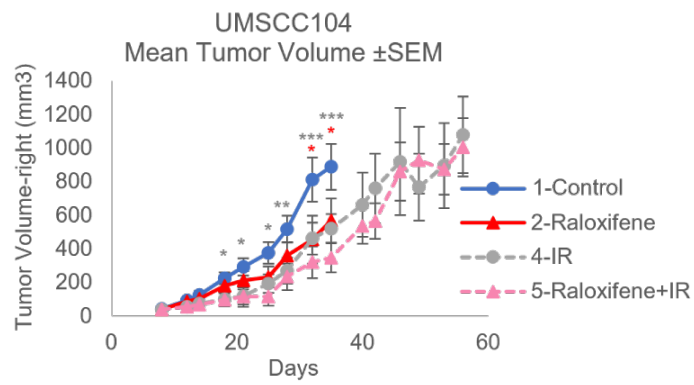
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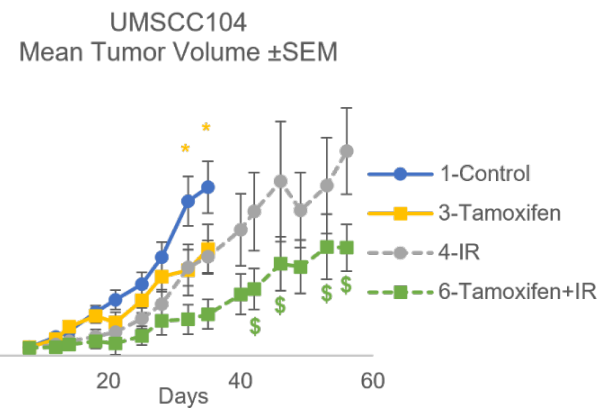
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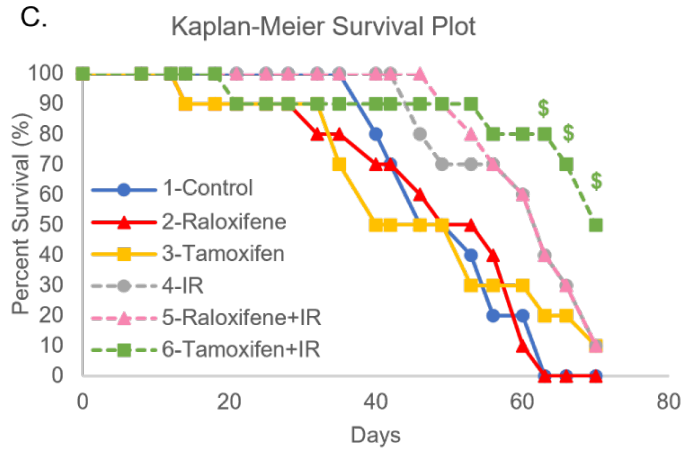
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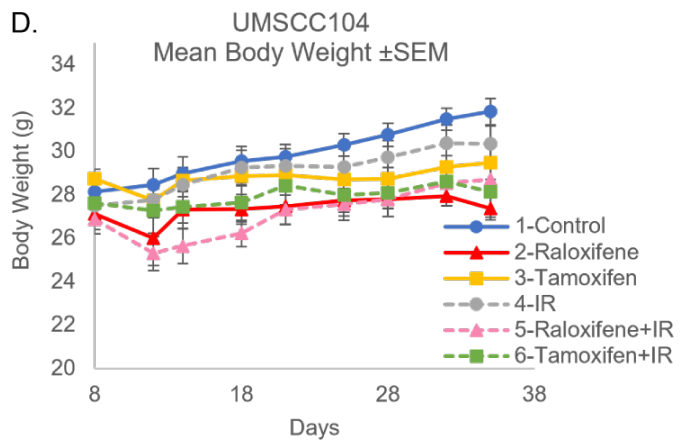
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C.



D.



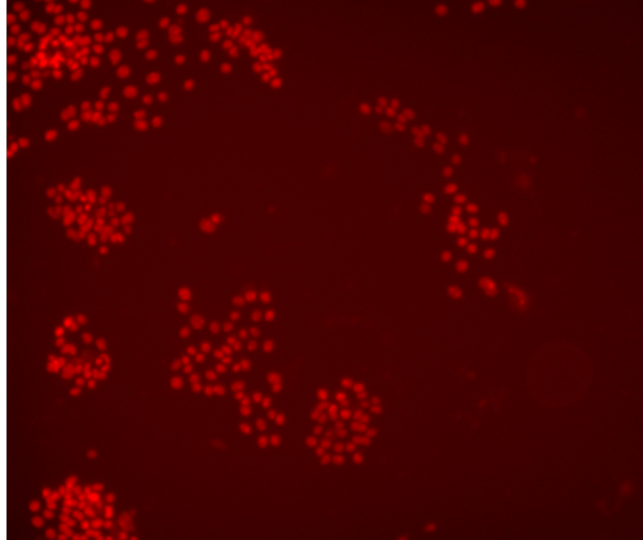
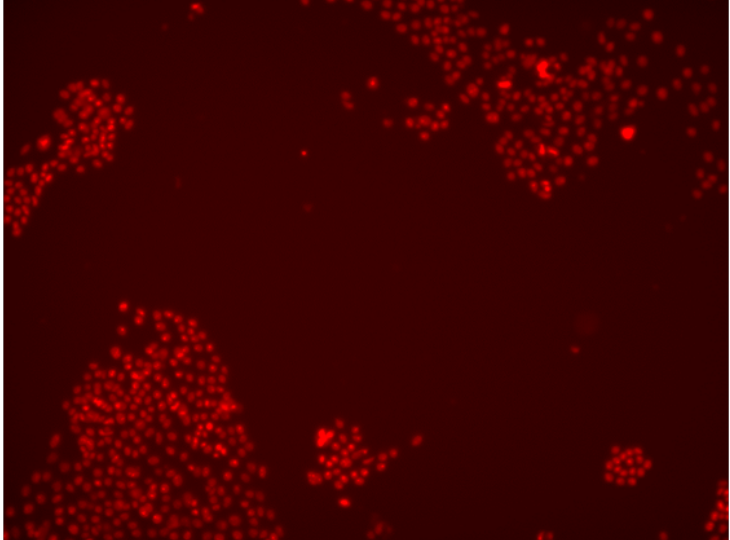
Traditional Cell Culture

Fibroblast Co-Culture

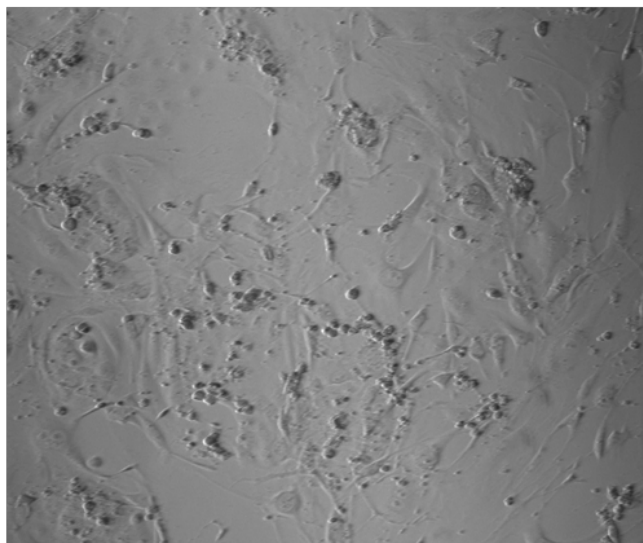
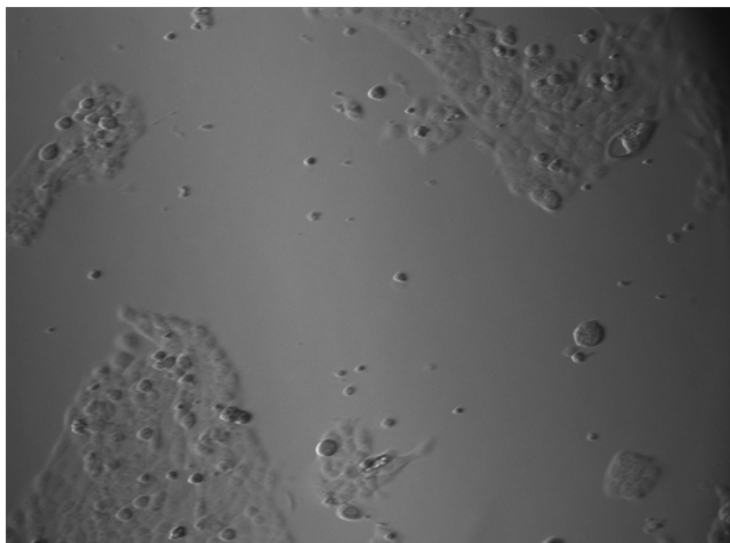
UMSCC104

UMSCC104 + J2

mKate₂-red



brightfield



merged

