1 Hotspot *ESR1* mutations are multimodal and contextual

2 modulators of breast cancer metastasis

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61	

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- 64

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106 Abstract

- 107 Constitutively active estrogen receptor-α (ER/ESR1) mutations have been identified in
- 108 approximately one third of ER+ metastatic breast cancers. Although these mutations are known
- 109 mediators of endocrine resistance, their potential role in promoting metastatic disease has not
- 110 yet been mechanistically addressed. In this study, we show the presence of *ESR1* mutations
- 111 exclusively in distant but not local recurrences in five independent breast cancer cohorts. In
- 112 concordance with transcriptomic profiling of ESR1 mutant tumors, genome-edited ESR1 Y537S
- and D538G mutant cell models exhibited a reprogrammed cell adhesive gene network via
- alterations in desmosome/gap junction genes and the TIMP3/MMP axis, which functionally
- 115 conferred enhanced cell-cell contacts while decreasing cell-extracellular matrix (ECM) adhesion.
- 116 In vivo studies showed *ESR1* mutant cells were associated with larger multi-cellular circulating
- 117 tumor cell (CTC) clusters with increased compactness compared to *ESR1* WT CTCs. These
- 118 preclinical findings translated to clinical observations, where CTC clusters were enriched in
- 119 patients with *ESR1*-mutated metastatic breast cancer. Conversely, context-dependent migratory
- 120 phenotypes revealed co-targeting of Wnt and ER as a vulnerability in a D538G cell model.
- 121 Mechanistically, mutant ESR1 exhibited non-canonical regulation of several metastatic
- 122 pathways, including secondary transcriptional regulation and de novo FOXA1-driven chromatin
- 123 remodeling. Collectively, these data provide evidence for *ESR1* mutation-modulated metastasis
- 124 and suggest future therapeutic strategies for targeting *ESR1* mutant breast cancer.

125

126 Statement of Significance

- 127 Context and allele-dependent transcriptome and cistrome reprogramming in mutant ESR1 cell
- 128 models elicit diverse metastatic phenotypes related to cell adhesion and migration, which can
- 129 be pharmacologically targeted in metastatic breast cancer.

130

131 Introduction

- 132 More than 70% of breast cancers express estrogen receptor-α (ER/ESR1). Antiestrogen
- 133 therapies, including depletion of estradiol (E2) by aromatase inhibitors (Als) or
- 134 antagonizing ER activity by Selective Estrogen Receptor Modulators/Degraders
- 135 (SERMs/SERDs), are conventional treatments for ER+ breast cancer. Development of
- 136 resistance to these endocrine therapies, however, remains a clinical and socioeconomic
- 137 challenge (1,2).
- 138 30-40% of endocrine-resistant metastatic breast cancer (MBC) is enriched in ESR1
- 139 somatic base pair missense mutations (3-5), that can be detected in the blood of
- patients with advanced disease (6,7). Clinically, ligand binding domain (LBD) *ESR1*
- 141 mutations correlate with poor outcomes in patients with advanced disease (6,8,9).
- 142 Recent work from our group and others has uncovered a crucial role for these *ESR1*
- 143 hotspot mutations in driving constitutive ER activity and decreased sensitivity towards
- 144 ER antagonists (10-12). Moreover, structural investigation of the two most frequent
- 145 mutations, variants Y537S and D538G, has demonstrated that *ESR1* mutations stabilize
- 146 helix 12 (H12) in an agonist conformation, thereby providing a mechanistic explanation
- 147 for constitutive ER activity (13).
- 148 The identification of *ESR1* mutations in endocrine resistant MBC suggests that mutant
- 149 ER may not only mediate endocrine resistance but also have an unappreciated role in
- 150 enabling metastasis. Indeed, recent *in vivo* studies showed that mutant ER can promote
- 151 metastasis (14,15), and *in vitro* studies showed a gain of cell motility (15,16) and growth
- 152 in 3D culture (17). Although epithelial-mesenchymal transition (EMT) has been
- 153 described as one potential explanation for the Y537S mutant (18), overall mechanisms
- remain largely unclear. In order to identify personalized therapeutic vulnerabilities in
- 155 patients harboring *ESR1* hotspot mutations, there is an urgent need to decipher the
- 156 mechanistic underpinnings and precise roles of mutant ER in the metastatic progression
- 157 using comprehensive approaches and model systems.
- 158 Previous transcriptomic profiling performed by us and others has revealed a context-
- 159 dependence of *ESR1* mutation effects, as well as significant differences between the
- 160 two most frequent hotspot mutations, Y537S and D538G (11,12,14,15,19). Differentially

- 161 expressed genes vary widely following expression of the mutations in their respective
- 162 cell line model, however, both Y537S and D538G maintain distinction from the E2-
- 163 dependent wild-type (WT) ER transcriptome. Similarly, comparison of the WT and
- 164 mutant ER cistromes has also revealed context-dependent and allele-specific effects on
- 165 ER recruitment (11,14). Furthermore, we recently showed that *ESR1*-mutant
- 166 transcriptomic reprogramming is associated with epigenetic remodeling (19). While
- 167 these findings imply that in the setting of high molecular diversity in tumors and patients,
- 168 somatic *ESR1* mutations have the potential to trigger different metastatic phenotypes,
- 169 this phenomenon has yet to be investigated.
- 170 In this study, we explore metastatic gain-of-function phenotypes in genome-edited
- 171 ESR1 mutant models under the guidance of transcriptomic changes detected in clinical
- samples. We identify mechanisms underlying context and allele-specific metastatic
- 173 phenotypes, and subsequently confirm alterations in a number of potential therapeutic
- 174 targets in metastatic tumors. We believe that our systematic bedside-to-bench approach
- 175 will ultimately lead to improved metastasis-free outcomes and prognosis for patients
- 176 with ER+ tumors.
- 177

178 Materials and methods

- 179 Additional details and references are provided in the Supplementary Materials and
- 180 Methods section.
- 181 Human tissue studies from the Womens Cancer Research Center (WCRC) and

182 Charite cohorts

- 183 All patients enrolled were approved within IRB protocols (PRO15050502) from the
- 184 University of Pittsburgh and Charite Universitaetsmedizin Berlin. Informed consent was
- obtained from all participating patients. Biopsies were obtained and divided into distant
- 186 metastatic or local recurrent tumors. Genomic DNA was isolated from formalin fixed
- 187 paraffin embedded (FFPE) samples and *ESR1* mutation status was detected with
- droplet digital PCR (ddPCR) targeting Y537S/C/N and D538G mutations in pre-
- amplified *ESR1* LBD products as previously reported (7).
- 190

191 CTCs analysis from the NU16B06 Cohort

- 192 A retrospective cohort comprising 151 Metastatic Breast Cancer (BC) patients
- 193 characterized for CTCs, and ctDNA at the Robert H. Lurie Comprehensive Cancer
- 194 Center of Northwestern University (Chicago, IL) between 2015 and 2019 was analyzed.
- 195 Patients' enrollment was performed under the Investigator Initiated Trial (IIT) NU16B06
- 196 independently from treatment line. The overall baseline staging was performed
- 197 according to the investigators' choice, CTCs and ctDNA collection was performed prior
- 198 to treatment start. CTC enumeration was performed though the CellSearch™
- 199 immunomagnetic System (Menarini Silicon Biosystems). Mutations in ESR1 (hotspots
- 200 D538 and Y537) and PIK3CA (hotspots E453 and H1047) were detected by either
- 201 ddPCR assay using the QX200 ddPCR System (Bio-Rad) or through the
- 202 Guardant360[™] high sensitivity next-generation sequencing platform (Guardant Health,
- 203 CA). More details for CTC enumeration, mutation detection and statistical analysis can
- 204 be found in Supplementary Materials and Methods.
- 205

206 Cell culture

- 207 Genome-edited MCF7 (RRID: CVCL_0031) and T47D (RRID: CVCL_0553) ESR1
- 208 mutant cell models from different sources were maintained as previously described

209 (12,19,20). Hormone deprivation was performed for all experiments, unless otherwise210 stated.

211

212 Reagents

213 17β-estradiol (E2, #E8875) was obtained from Sigma, and Fulvestrant (#1047),

214 carbenoxolone disodium (#3096) and EDTA (#2811) were purchased from Tocris.

- LGK974 (#14072) and T-5224 (#22904) were purchased from Cayman. Marimastat
- 216 (S7156) was obtained from SelleckChem. Recombinant human Wnt3A (5036-WN-010)
- 217 was purchased from R&D Systems. For knockdown experiments, siRNA against
- 218 FOXA1 (#M-010319), DSC1 (#L-011995), DSC2 (#L-011996), GJA1 (#L-011042) and
- 219 GJB2 (#L-019285) were obtained from Horizon Discovery. Desmosome and scramble
- 220 peptides were designed based on previous studies (21,22) and synthesized from
- 221 GeneScript. Peptide sequences are presented in Supplementary Table S10.
- 222

223 Animal Studies

224 Long term metastatic evaluation: 4-week old female *nu/nu* athymic mice were 225 ordered from The Jackson Laboratory (002019 NU/J, RRID: IMSR JAX:002019) 226 according to University of Pittsburgh IACUC approved protocol #19095822. MCF7 and 227 T47D ESR1 mutant cells were hormone deprived and resuspended in PBS with a final 228 concentration of 10⁷ cells/ml. 100µl of cell suspension was then injected via tail vein into 229 nude mice with 7 mice per group. Mice were under observation weekly. According to the 230 IACUC protocol, if greater than 50% of mice in any group show predefined signs of 231 euthanasia, the entire cohort needs to be euthanized. Cohorts were euthanized at 13 232 weeks for MCF7 cell-injected mice and 23 weeks for T47D cell-injected mice. Macro-233 metastatic tumors and potential organs (lung, liver, UG tract) for metastatic spread were 234 harvested. Solid macro-metastatic tumors (non-lymph node) were counted for 235 comparison. All tissues were processed for FFPE preparation and hematoxylin and 236 eosin (H&E) staining by the Histology Core at Magee Women's Research Institute. 237 Macro-metastatic tumor FFPE sections were further evaluated by a trained pathologist. 238 Micro-metastatic lesions in the lung were further examined and quantified by 239 immunofluorescence staining as described in supplementary materials and methods.

240 Short term CTC cluster assessment: 4-week old female *nu/nu* athymic mice were 241 ordered from The Jackson Laboratory (002019 NU/J) according to University of 242 Pittsburgh IACUC approved protocol #19095822. MCF7 WT and mutant cells were 243 stably labelled with RFP-luciferase by infection with the pLEX-TRC210/L2N-TurboRFP-244 c lentivirus plasmid. Labelled cells were hormone deprived and resuspended in PBS at 245 a final concentration of 10⁷ cells/ml. 100µl of cell suspension was then injected into 246 nude mice with 6 mice per group via an intracardiac left ventricle injection. Post-injected 247 mice were immediately imaged using the IVIS200 in vivo imaging system (124262, 248 PerkinElmer) after D-luciferin intraperitoneal injection to confirm successful cell delivery 249 into the circulation system. All mice were euthanized after one hour of injection and their 250 whole blood were extracted via cardiac puncture and collected into CellSave 251 Preservative Tubes (#790005, CellSearch). Blood samples were mixed with 7ml of 252 RPMI media and shipped to University of Minnesota for CTC enrichment. CTCs were 253 extracted using an electric size-based microfilter system (FaCTChekr) and stained with 254 antibody against pan-cytokeratins (CK) and DAPI. Slides with stained CTCs were 255 manually scanned in a blind manner and all visible single CTCs or clusters were imaged 256 under 5X or 40X magnification respectively. To set up criteria for identifying CTC 257 clusters via images, we analyzed seven single CTCs with intact CK signal distribution 258 and calculated the average nuclei-edge to membrane distance (x). Inter-nuclei-edge 259 distance greater than 2x for any two CTCs were excluded in CTC cluster calling. All 260 measurements were performed in a blind manner. Details of filter and staining are 261 included in the supplementary materials and methods.

262

263 **qRT-PCR**

MCF7 and T47D cells were seeded in triplicates into 6-well plates with 120,000 and 90,000 cells per well respectively. After desired treatments, RNA was and cDNA was synthesized using iScript kit (#1708890, BioRad, Hercules, CA). qRT-PCR reactions were performed with SybrGreen Supermix (#1726275, BioRad), and the $\Delta\Delta$ Ct method was used to analyze relative mRNA fold changes with *RPLP0* measurements serving as the internal control. All primer sequences can be found in Supplementary Table S10.

271 Immunoblotting

272 After desired treatments, cells were lysed with RIPA buffer spiked with a fresh protease

- and phosphatase cocktail (Thermo Scientific, #78442) and sonicated. Protein
- 274 concentrations were quantified using the Pierce BCA assay kit (Thermo Fisher,
- 275 #23225). 80-120µg of protein for each sample was loaded onto SDS-PAGE gels, and
- then transferred onto PVDF membranes. The blots were incubated with the following
- antibodies: desmocollin 1 (sc-398590, RRID: AB_2894905), desmoglein 2 (sc-80663,
- 278 RRID: AB_2093438), plakophilin (sc-33636, RRID: AB_2164139), connexin 26 (sc-
- 279 7261, RRID: AB_2110895) and cFOS (sc-52, RRID: AB_2106783) from Santa Cruz;
- 280 ER-α (#8644, RRID: AB_2617128), HA (#3724, RRID: AB_1549585), Non-phospho-β-
- 281 catenin (#19807, RRID: AB 2650576), Histone H3 (#4499, RRID: AB 10544537), AIF
- 282 (#5318, RRID: AB 10634755), GSK3β (Ser9, #5558, RRID: AB 10013750), phospho-
- 283 GSK3α (Ser21, #9316, RRID: AB 659836), GSK3β (#12456, RRID: AB 2636978) and
- 284 GSK3α (#4337, RRID: AB_10859910) from Cell Signaling Technology; β-catenin
- 285 (#610154, RRID: AB_397555) from BD; Tubulin (T6557, RRID: AB_477584) and
- connexin 43 (C6219, RRID: AB_476857) from Sigma Aldrich; and *TIMP3* (ab39184,
- 287 RRID: AB_2204971) from Abcam.
- 288

289 IncuCyte Live Cell Imaging System

- 290 Wound scratch assay. MCF7 or T47D cells were seeded at 150,000 cells/well into
- 291 Imagelock 96-well plates (Essen Bioscience, #4379) pre-coated with Matrigel (Corning,
- 4356237). Wounds were scratched in the middle of each well using a Wound Maker
- 293 (Essen Bioscience, #4493). Desired treatments mixed with 5µg/ml of proliferation
- blocker Mitomycin C (Sigma-Aldrich, #10107409001) were loaded after two washes with
- 295 PBS. The IncuCyte Zoom system was used to record wound images every 4 hours and
- wound closure density was calculated using the manufacturer's wound scratch assay
- 297 module. For the dominant negative *TCF4* overexpression experiment, Myc-tagged
- 298 DN*TCF4* plasmids (Addgene, #32729) were transiently transfected into targeted cells
- for a total of 24 hours before being subjected to the wound scratch assay.
- 300 Aggregation rate assay. 3,000 MCF7 or 4,000 T47D cells were seeded into 96-well
- 301 round bottom ultra-low attachment plates (Corning, #7007) with 100µl of respective

- 302 media in each well. Cell aggregation was monitored by the IncuCyte living imaging
- 303 system every hour. Spheroid areas were normalized to time 0.
- 304

305 Calcein-labelled cell-cell interaction assay

306 MCF7 and T47D cells were seeded into black-walled 96 well plate at 150,000 cells per 307 well to achieve a fully confluent monolayer after 24 hours. Separate cultures of cells 308 were digested and labelled with 1µM calcein AM (BD Pharmingen, #564061) for 30 309 minutes in room temperature. 40,000 labelled cells were loaded on top of the previously 310 plated monolayers and incubated for 1 hour at 37°^C. Cells were washed three times 311 after incubation by manually pouring out the PBS washing agent. The plates were read 312 using Victor X4 plate reader (PerkinElmer) under the excitation and emission 313 wavelength of 485/535nm. Cell-cell adhesion ratios were calculated by dividing the 314 post-wash readouts to the pre-wash readouts after each wash. For the vacuum 315 aspiration method, we used a standard laboratory vacuum pump with a modified speed 316 of approximately 100 ml/minutes. Adhesion ratios after three washes were plotted 317 separately for each independent experiment.

318

319 Ibidi microfluidic system

320 MCF7 and T47D ESR1 mutant cells were hormone deprived for 3 days and diluted to 321 10⁶ cells in 14ml of respective media before being loaded into the ibidi pump system 322 (ibidi, #10902). Cells were constantly flowing with 15dynes/cm of shear stress for two 323 hours before immediate imaging after being seeded back into a flat bottom ULA plate. 324 For each group, six wells were imaged twice. Time zero (T0) cells were also imaged as 325 the initial time point control. Cell numbers in clusters or non-clusters were manually 326 counted. Cell cluster ratios were calculated by dividing the cell numbers in clusters to 327 the total number of cells. Cell clustering grade was calculated by the cell numbers 328 present in each cluster. For CBX treatment, cells were pre-treated with 100µM CBX for 329 two days before being added to the flow chamber. For the desmosome blocking 330 peptides treatment, 75µM of each DSC1, DSC2, DSG1 and DSG2 peptide or 150µM of 331 each scramble peptide were pre-mixed into cell suspension for flow experiments.

332

333 Cell-ECM adhesion assay

- 334 30,000 cells/well were seeded into collagen I coated (Thermo Fisher Scientific,
- A1142803) or uncoated 96-well plates. For the ECM array assay, cells were
- resuspended and loaded into the ECM array plate (EMD Millipore, ECM540). After a 2-
- hour incubation at 37°^C, the plates were washed with PBS three times, and attached
- 338 cells were quantified using the FluoReporter kit (Thermo Fisher Scientific, F2962).
- 339 Adhesion ratios were calculated by dividing the remaining cell counts in the washed
- 340 wells to the initial cell counts in pre-washed plates. For *TIMP3* overexpression, the
- 341 PRK5M-TIMP3 plasmid (Addgene, #31715) was transfected into targeted cells, which
- 342 was subjected to the adhesion assay after a 24-hour transfection period.
- 343

344 Chromatin-immunoprecipitation (ChIP)

- 345 ChIP experimentation was performed as previously described (23). The
- immunoprecipitation was performed using ERα (sc543, RRID: AB_631471) and rabbit
- 347 IgG (sc2027, RRID: AB_737197) antibodies (Santa Cruz Biotechnologies). Histone 3
- 348 acetylation at K27 site (ab4729, RRID: AB_2218291), and Histone 3 di-methylation at
- 349 K4 site (ab7766, RRID: AB_2560996) and FOXA1 (ab23738, RRID: AB_2104842)
- antibodies were obtained from Abcam. Detailed ChIP-seq analysis are provided in the
- 351 Supplementary Material and Methods.
- 352

353 Statistical Analysis

- 354 GraphPad Prism software version 7 and R version 3.6.1 were used for statistical
- analysis. All experimental results included biological replicates and were shown as
- 356 mean ± standard deviation, unless otherwise stated. Specific statistical tests were
- 357 indicated in corresponding figure legends. All tests were conducted as two-tailed, with a
- 358 p<0.05 considered statistically significant. Drug synergy was calculated based on the
- 359 Bliss independence model using the SynergyFinder (<u>https://synergyfinder.fimm.fi/</u>) (24).
- 360 Bliss synergy scores were used to determine synergistic effects.

361

362 Data Availability Statement

- 363 The ER and FOXA1 ChIP-seq data has been deposited onto the Gene Expression
- 364 Omnibus database (GSE125117 and GSE165280). All publicly available resources
- 365 used in this study are summarized in Supplementary Table S11. All raw data and scripts
- 366 are available upon request from the corresponding author.

367 Results

368 Significant enrichment of *ESR1* mutations in distant metastases compared to 369 local recurrences

370 We compared *ESR1* mutation frequencies between distant metastatic and locally 371 recurrent tumors. A combination of four previously reported clinical cohorts (MSKCC, 372 METAMORPH, POG570 and IEO) showed that while 155/877 distant metastases (18%) 373 harbored ESR1 mutations, none were found in 44 local recurrences (Table 1 and 374 Supplementary Table S1) (25-28). To expand upon this observation, we additionally 375 screened 75 ER+ recurrent tumors from the Women's Cancer Research Center 376 (WCRC) and Charite Hospital for ESR1 hotspot (Y537S/C/N and D538G) mutations 377 using highly sensitive droplet digital PCR (ddPCR). We identified 12 ESR1 mutation-378 positive cases among the distant metastases (25%), whereas none of the local 379 recurrences were *ESR1* mutation-positive (Table 1 and Supplementary Table S2). 380 There was no significant difference in time to recurrence for patients with distant vs local 381 recurrences (Supplementary Fig. S1A & Table S3), making it less likely that the 382 observed differences could simply be due to duration of time to recurrence between 383 local and distant recurrences, as was previously suggested (6). If however, we compare 384 time to distant recurrence between tumors with WT and mutant ESR1, we observed 385 significantly longer time to recurrence in two of the five cohorts (Supplementary Fig 386 S1B). For three of the cohorts (WCRC/Charite, POG570, and MSKCC), details on lines 387 of therapies was available, and we observed that in two of the cohorts, patients with 388 distant metastases had on average significantly more lines of therapy compared to 389 those with local recurrences (Supplementary Fig. S1C and S1D), and that patients with 390 ESR1 mutant tumors had been exposed to more lines of therapies than those with WT 391 tumors (Supplementary Fig. S1E). Finally, we repeated our comparative analysis of 392 frequency of ESR1 mutations in local and distant recurrences restricting it to patients 393 exposed to endocrine therapies and confirmed significant enrichment of ESR1 394 mutations in distant metastasis (Supplementary Table S1). Thus, while collectively 395 these analyses recapitulated that *ESR1* mutations are arising primarily as an outcome 396 of therapeutic selection, their more frequent occurrences in distant compared to local

397 recurrences suggest a potential role in promoting metastasis in addition to conferring398 endocrine resistance.

ESR1 mutant tumors show a unique transcriptome associated with multiple metastatic pathways

To identify candidate functional pathways mediating the metastatic properties of *ESR1*mutant cells, we compared WT and *ESR1* mutant tumor transcriptomes from four
cohorts of ER+ metastatic tumors: our local WCRC cohort (46 *ESR1* WT and 8 mutant
tumors) (29-31) and three previously reported cohorts - MET500 (34 *ESR1* WT and 12
mutants tumors), POG570 (68 *ESR1* WT and 18 mutants tumors) and DFCI (98 *ESR1*WT and 32 mutants tumors) (14,27,32) (Fig. 1A & Supplementary Table S4).

- 407 Although principal component analyses on global transcriptomes did not segregate
- 408 *ESR1* WT and mutant tumors (Supplementary Fig. S2A), both "Estrogen Response
- 409 Early" and "Estrogen Response Late" signatures were significantly enriched in ESR1
- 410 mutant tumors in 3 out of 4 cohorts, with a trend towards enrichment in the fourth cohort
- 411 (Fig. 1B). These results recapitulate the observation of ER hyperactivation as a result of
- 412 hotspot mutations, previously described in other preclinical studies (12,14,20).
- 413 Differential gene expression analysis identified a considerable number of altered genes
- that were associated with *ESR1* mutations (Fig. 1C & Supplementary Table S5), which
- 415 further inferred functional alterations in various metastasis-related pathways.
- 416 Remarkably, "Cell-To-Cell Signaling & Interaction" and "Cell Movement" were featured
- 417 among the top five altered pathways for *ESR1* mutant tumors in all four cohorts (Fig.
- 418 **1D)**.
- 419 In addition to the broad effects associated with *ESR1* mutations, we next questioned
- 420 whether different *ESR1* mutant variants could display divergent functions. A meta-
- 421 analysis of the five above-mentioned ER+ MBC cohorts examining *ESR1* mutations
- 422 underscored D538G (37%) and Y537S (24%) as the predominant variants (Fig. 1E).
- 423 Given the challenge of merging RNA-seq data sets from multiple cohorts due to
- 424 immense technical variations, we selectively compared mutation variant specific
- 425 transcriptomes of ten Y537S- or eight D538G-harboring tumors to the WT counterpart

426 (n=32) respectively from the DFCI cohort, which provided the largest numbers and thus 427 maximized statistical power. Aligning enrichment levels of 50 hallmark gene sets for the 428 two mutant variants again confirmed "Estrogen Response Early" and "Estrogen 429 Response Late" as the top co-upregulated pathways (Fig. 1F), with Y537S tumors 430 displaying higher ER activation (Supplementary Fig. S2B), consistent with cell line 431 studies (12,33). The similar observation was also validated in MET500 and POG570 432 cohort regardless of the smaller sample size (Supplementary Fig. S2B). We also identified enriched cell cycle related pathways (E2F targets, G2M checkpoint and mitotic 433 434 spindle) and metabolic related pathways (fatty acid, bile acid and xenobiotic 435 metabolisms) in Y537S and D538G tumors, respectively, implying that different ESR1 436 mutant variants might hijack distinct cellular functions to promote malignancy. Again, 437 similar trends of these mutant-variant pathways were recapitulated in POG570 cohort 438 (Supplementary Fig. S2C). Taken together, these results provide support that despite 439 mutant variant-specific alterations, ESR1 mutations might broadly mediate metastatic 440 phenotypes through effects on cell-to-cell interactions and cell movement. We next 441 validated the *in silico* results using previously established genome-edited MCF7 and 442 T47D cell line models (12).

443 ESR1 mutant-cells exhibit stronger cell-cell adhesion

We first addressed the enrichment of cell-cell interaction signaling in the mutant tumors
through morphological inspection of cell cluster formation in suspension culture (Fig.
2A). We observed more compact cell clusters in MCF7 and T47D mutant cell lines
compared to their WT counterparts after six days of suspension culture. A time course
study confirmed enhanced cluster formation 24-48hrs past cell seeding (Supplementary
Fig. S3A). Similar observations were made in individual clones, eliminating the
possibility for clonal effects (Supplementary Fig. S3B).

- 451 Since *ESR1* mutant cells displayed significantly increased ligand-independent growth in
- 452 suspension (Fig. 2B), we sought to rule out the possibility that increased cluster
- 453 formation was simply a result of increased cell number by assessing cell-cell adhesive
- 454 capacity using multiple approaches in short term culture (within 1 day). We therefore

455 directly quantified homotypic cell-cell interactions by measuring the adhesion of calcein-456 labelled ESR1 WT or mutant cells. This assay showed that both MCF7 mutant cells 457 exhibited significantly stronger cell-cell adhesion compared to the WT cells (Fig. 2C). In 458 T47D cells, a similar effect was observed, but was limited to the T47D-Y537S mutant 459 cells (Supplementary Fig. S4A). These assays were complemented by quantification of 460 cell aggregation rates as a direct reflection of cell-cell adhesion, which confirmed faster 461 aggregation in MCF7-Y537S/D538G and T47D-Y537S cells (Fig. 2D & Supplementary 462 Fig. S4B-S4D). In addition, these stronger cell-cell adhesive properties were also 463 reproduced in additional ESR1 mutant cell models from other laboratories (19,20)

464 (Supplementary Fig. S4E and S4F).

465 Cell-cell interaction has been reported to affect several stages of metastasis, including 466 collective invasion, intravasation, dissemination and circulation (34-36). To test whether 467 ER mutations may affect tumor cell-cell adhesion in circulation, we utilized a microfluidic 468 pump system to mimic arterial shear stress. Comparing representative images before 469 and after 2 hours of microfluidic flow, we found MCF7 ESR1 mutant cells had a greater 470 tendency to aggregate together (Fig. 2E and 2F). Larger clusters comprised of five or 471 greater cells were more prevalent in the ESR1 mutant cell lines, whereas smaller two-472 cell clusters were diminished (Fig. 2G). A similar phenotype was also identified in 473 additional MCF7 ESR1 mutant cells and in our T47D-Y537S cell line (Supplementary 474 Fig. S5A-S5I), consistent with our observations in static conditions. In an additional 475 orthogonal approach, we utilized a quantitative microfluidic fluorescence microscope 476 system simulating blood flow (37). Quantification of dynamic adhesion events 477 normalized to adhesion surfaces revealed a consistent enhanced cell-cell adhesion 478 capacity of ESR1 mutant MCF7 cells (Supplementary Fig. S5J-S5K, Supplementary 479 videos 1-3). Together, these results show that hotspot ESR1 mutations confer 480 increased cell-cell attachment under static and fluidic conditions, and that the effect size 481 is dependent upon mutation type and genetic backgrounds. These findings are at odds 482 with increased EMT features (18), and indeed the majority of ESR1 mutant models and 483 tumors did not show increased EMT signature or increased expression of EMT marker 484 genes (Supplementary Fig. S6A-S6D).

485 We next sought to assess whether this unexpected phenotype translated into numbers 486 of CTC clusters and subsequent metastasis in vivo. One hour post intracardiac injection 487 into athymic mice, circulating MCF7 WT and mutant cells were enriched from blood 488 using a previously described electrical CTC filtering method (38) (Fig. 2H). 41%-81% of 489 CTC clusters were composed of both cancer and non-cancer cells (Supplementary Fig. 490 S7A). Despite no difference in the average amount of single CTCs and CTC clusters 491 per mouse between the WT and mutant *ESR1* (Supplementary Fig. S7B & S7C), we 492 found that overall MCF7-Y537S mutant cells were significantly enriched in clusters with 493 greater than 2 cells (Fig. 2I). Furthermore, guantification of inter-nuclei distances 494 between two-cell clusters revealed denser MCF7-Y537S clusters (Fig. 2J), supporting 495 stronger MCF7-Y537S cell-cell interactions in an in vivo blood circulation environment. 496 The data from the MCF7-D538G mutant cells did not recapitulate the adhesive 497 phenotype we discerned in vitro, suggesting mutation site-specific interactions with the 498 in vivo microenvironment potentially affect cluster formation.

499 We next performed tail vein injection and monitored bloodborne metastatic development 500 in longer-term *in vivo* experiments without estradiol supplement (Fig. 2K). We observed 501 multiple distant macro-metastatic tumors in 4/6 (67%) MCF7-Y537S mutant cell-injected 502 mice (Fig. 2L), likely as an outcome of the well-established ligand-independent cell 503 growth. In contrast, distant macro-metastatic tumor was observed in only one mouse of 504 MCF7-D538G group (1/7) and none in MCF7-WT group (0/7) (Fig. 2M, left panel). The 505 enhanced macro-metastasis observed in MCF7-Y537S but not D538G mutant was 506 consistent with our *in vivo* CTC clustering experiment, opening up the possibility that the 507 enhanced CTC clustering ability might confer an additional metastatic advantage. We 508 detected no difference in lung micro-metastatic foci areas between WT and mutant cell-509 injected mice, potentially due to a high baseline of MCF7 lung colonization capacity 510 (Fig. 2M, right panel). In contrast to our results with MCF7 cells, we only discerned one 511 macro-metastatic tumor from each T47D mutant group (Y537S: 1/6; D538G: 1/7) and 512 none in T47D-WT group (0/7) after 23 weeks of injection (Fig. 20, left panel), 513 underpinning its less aggressive behavior as compared to MCF7 cells (39,40). 514 However, both T47D-Y537S and T47D-D538G mutant cells resulted in enlarged lung

515 micro-metastases (Fig. 2N and 2O, right panel).

516 Encouraged by our *in vitro* and *in vivo* findings, we next examined CTC clusters in 517 patients with ESR1 mutant tumors. Taking advantage of a recent CTC sequencing 518 study (41), we sought to generate CTC cluster gene signatures. Differential gene 519 expression analysis in two patients with ER+ disease who had at least two CTC clusters 520 and single CTCs sequenced identified CTC cluster enriched genes (Supplementary Fig. 521 S8A and Table S6), which we subsequently applied to our RNA-seq dataset with 51 522 pairs of ER+ primary-matched metastatic tumors (44 ESR1 WT and 7 mutant) merged 523 from the WCRC and DFCI cohorts. ESR1 mutant metastatic tumors exhibited 524 significantly higher enrichment of CTC cluster-derived gene signatures (Supplementary

525 Fig. S8B and S8C).

526 To examine the interplay between *ESR1* mutations, numbers of CTCs, and clinical 527 outcome, we analyzed a cohort of 151 patients with MBC. Median age at the first blood 528 draw for CTCs enumeration was 55 years (IQR: 44 - 63 years), 76 patients (50.3%) 529 were diagnosed with ER+ HER2-negative MBC, 38 (25.2%) with HER2-positive MBC 530 and 37 (24.5%) with TNBC. Bone (49.7%), lymph nodes (41.1%), lung (34.4%) and liver 531 (34%) were the most common sites of metastasis (Supplementary Table S7). Median 532 number of CTCs was 1 (IQR: 0-10), clusters were detectable in 14 patients (9.3%) (Fig. 533 2P) and in this subgroup the median number of clustered CTCs (i.e., number of CTCs 534 involved in clusters) was 15.5 (IQR: 4 - 20). Clusters with CTCs >4 and ≤4 were 535 detected in 10 (6.6 %), and 4 (2.7%) cases, respectively. Among patients without 536 clusters (90.7%), 101 (66.89%) and 36 (23.84%) were respectively classified as stage 537 IV Indolent (< 5 CTCs) and Aggressive (\geq 5 CTCs) according to our previous study (42) 538 (Supplementary Table S7). Mutations in hotspots D538 and Y537 of ESR1 were 539 detected in 30 patients (19.9%), while mutations in hotspots E453 and H1047 of 540 PIK3CA were detected in 40 patients (26.5%) (Supplementary Table S7). Median 541 follow-up was 30.8 months. A significant association was observed between ESR1 542 genotype status and clustered CTCs > 4 (P = 0.029) (Fig. 2Q), a significant association 543 was retained after adjusting for MBC subtype (OR: 5.51, 95%C.I.: 1.29 - 23.52 P = 544 0.021). A similar trend was highlighted in the ER+ HER2-negative subgroup specifically 545 (Supplementary Fig. S8D). No association was observed with respect to PIK3CA 546 (P=0.725). Notably, patients with > 4 clustered CTCs experienced the worse prognosis

- 547 with respect to Stage IV indolent in terms of OS both in the general population (Fig. 2R)
- 548 (P < 0.0001) and in the ER+ HER2-negative subgroup (Supplementary Fig. S8E) (P <
- 549 0.0001). After adjusting for MBC subtype, >4 clustered CTCs and Stage IV aggressive
- 550 without clusters retained their prognostic impact (respectively HR: 15.50, 95%CI: 6.90 -
- 551 34.82. P < 0.001; HR: 2.37, 95%CI: 1.38 4.06. P = 0.002).

552 Mutant *ESR1* cells show increased desmosome gene and gap junction gene

553 families

554 To elucidate the mechanism of enhanced cell-cell adhesion, we investigated the 555 enrichment of four major cell-cell junction subtypes – desmosomes, gap junctions 556 (connexons), tight junctions and adherens junctions within the cell model RNA-seq data 557 (12) (Supplementary Table S6). Enrichment of the desmosome gene and gap junction 558 gene families was observed in both MCF7-Y537S/D538G and T47D-Y537S cells (Fig. 3A). Tight junctions were enriched in WT cells, and there were no differences in the 559 560 adherens junction gene family expression (Supplementary Fig. S9A). Individual gene 561 expression analysis (FC>1.2, p<0.05) identified 18 commonly upregulated desmosome 562 genes and 4 gap junction genes in both MCF7 ESR1 mutant cell lines (Fig. 3B). In 563 addition to keratins, induction of classical desmosome genes DSC1/2, DSG1/2 and 564 *PKP1*, and gap junction genes *GJA1*, *GJB2* and *GJB5* were observed and validated by 565 qRT-PCR in MCF7 cells (Fig. 3D). Higher protein levels were also observed for DSC1, 566 DSG2, PKP1, GJA1 (Cx43), and GJB2 (Cx26) (Fig. 3C). Immunofluorescence staining 567 revealed significantly higher DSG2 expression in MCF7-Y537S at cell-cell contact 568 surfaces, with a trend observed in MCF7-D538G (Fig. 3E). Consistent with the weaker 569 in vitro cell-cell adhesion phenotypes in T47D mutant cells, we observed less 570 pronounced desmosome and gap junction gene expression changes in T47D-Y537S 571 cells (Supplementary Fig. S9B). We validated the overexpression of the key 572 desmosome and gap junction genes in RNA-seq datasets from seven additional ESR1 573 mutant cell models and performed further validation studies in two of them 574 (Supplementary Fig. S9C-S9E) (11,15,19). Moreover, mining RNA-seg data from recently reported ESR1 WT and mutant ex vivo CTC models (43), we observed 575 576 overexpression of three gap junction and desmosome genes in the ESR1 mutant CTC

577 lines (Supplementary Fig. S9F). Finally, the top upregulated desmosome and gap
578 junction genes (Supplementary Table S6) were also found significantly enriched in intra579 patient matched primary and metastatic lesions with *ESR1* mutations (Fig. 3F).

580 We next investigated the functional roles of the reprogrammed adhesome in the ESR1 581 mutant MCF7 cells. Transient individual knockdown of DSC1, DSC2, GJA1 or GJB2 did 582 not cause significant changes in adhesion in either ESR1 mutant line (Supplementary 583 Fig. S10A). However, we found compensatory effects observed in the desmosome and 584 gap junction knockdowns as exemplified by increased GJA1 levels after DSC1 or DSC2 585 knockdown (Supplementary Fig. S10B). The adhesive phenotype was disrupted, 586 however, with an irreversible pan-gap junction inhibitor, Carbenoxolone (CBX), or with 587 blocking peptide cocktails against desmocollin1/2 and desmoglein1/2 proteins. Both 588 treatments caused significant inhibition of cell-cell aggregation in static conditions 589 (Supplementary Fig. S10C & S10D) as well as diminished cluster propensities and size 590 in microfluidic conditions (Fig. 3G-3L), suggesting redundancy in the mutant-driven 591 reprogrammed desmosome and connexon pathways. In summary, MCF7-592 Y537S/D538G and T47D-Y537S mutants showed increased expression of desmosome

and gap junction gene family components, which contributes to our observed enhancedcell-cell adhesion phenotype.

We next investigated the mechanisms underlying the elevated desmosome and gap 595 596 junction components in ESR1 mutant cells. Because hotspot ESR1 LBD mutations are well-described as conferring constitutive ER activation, we first examined if these cell-597 598 cell adhesion target genes are direct outcomes of ligand-independent transcriptional 599 programming. Interrogating publicly available RNA-seg and microarray datasets of six 600 estrogen treated ER+ breast cancer cell lines (12,23,44,45), we found limited and 601 inconsistent E2 induction of all examined cell-cell adhesion genes when compared to 602 classical E2 downstream targets such as *GREB1* and *TFF1* (Supplementary Fig. S11A). 603 Surprisingly, mining our MCF7 ESR1 mutant cell model ER ChIP-seq data (46) showed 604 an absence of proximate Y537S or D538G mutant ER binding sites (± 50kb of TSS) at 605 desmosome and connexon target gene loci. These results suggest that the

606 reprogrammed cell-cell adhesome is not a direct consequence of mutant ER genomic607 binding.

608 We therefore hypothesized that these altered adhesion target genes might be regulated 609 via a secondary downstream effect of the hyperactive mutant ER. A seven-day siRNA 610 ER knockdown assessment identified GJA1 as the only target gene that could be 611 blocked in mutant cells following ER depletion, whereas, strikingly, DSC1, DSG1, GJB2 612 and GJB5 mRNA levels were increased in all cell lines (Fig. 3M). This was congruent 613 with ESR1 knockdown in five additional ER+ parental cell lines, with the majority 614 exhibiting a decrease in GJA1 expression levels (Supplementary Fig. S11B). To unravel 615 potential intermediate transcription factors (TFs) involved in the secondary regulation, 616 we examined the levels of TFs previously reported to regulate GJA1 expression (47) 617 (Supplementary Fig. S11C). Among those, the AP1 family component FOS (cFos) was 618 identified as the top TF upregulated in ESR1 mutant cells in a ligand-independent 619 manner. In addition, the AP1-associated transcriptional signature was also significantly 620 enriched in MCF7 ESR1 mutant cells (Supplementary Fig. S11D), and hence we tested 621 if GJA1 overexpression was dependent on the cFOS/AP1 transcriptional network. 622 Higher cFOS mRNA and protein levels in *ESR1* mutant cells were confirmed, which 623 declined along with GJA1 levels after ESR1 knockdown (Fig. 3N & Supplementary Fig. 624 S11E). Importantly, pharmacological inhibition of cFOS-DNA binding partially rescued 625 GJA1 overexpression in ESR1 mutant cells (Fig. 30, Supplementary Fig. S11F-S11G). 626 In conclusion, our results denote GJA1 as an indirect target of mutant ER through 627 activation of the cFOS/AP1 transcriptional axis in MCF7 cell models.

628 Since the majority of the cell-cell adhesion targets altered in the ESR1 mutant cells 629 were not direct ER target genes (Supplementary Fig. S11A & S11B), we investigated 630 potential impacts of epigenetic remodeling on these targets. Using our recently reported 631 ATAC-seq dataset from T47D ESR1 mutant cells (19), we observed that one of the 632 connexon targets, GJB5, exhibited increased chromatin accessibility at its gene locus in T47D-Y537S cells (Supplementary Fig. S12A & S12B), suggesting that epigenetic 633 634 activation modulates gene expression in this particular context. We further evaluated 635 active histone modifications on our target gene loci in the MCF7 model. We observed

- 636 enhanced H3K27ac and H3K4me2 recruitment in both MCF7-Y537S and D538G cells
- 637 at the nearest two histone modification sites around the DSC1 and DSG1 loci, the two
- 638 most upregulated desmosome component genes in MCF7 mutant cells (Fig. 3P),
- 639 suggesting activation of desmosome genes via an indirect ER-mediated epigenetic
- 640 activation (Fig. 3Q).

641 *ESR1* mutations promote reduced adhesive and enhanced invasive properties via 642 altered *TIMP3*-MMP axis

643 In addition to altered cell-cell adhesion, metastasis is also mediated by coordinated 644 changes in cell-matrix interaction (48,49). Therefore, we assessed whether mutant ER 645 affects interaction with the extracellular matrix (ECM). Computational analysis showed 646 inverse correlation between ECM receptor pathway signatures and ESR1 mutation 647 status in the DFCI cohort with the same trend appearing in 2/3 of the remaining cohorts 648 (Fig. 4A, Supplementary Fig. S13A & Table S6). Employing an adhesion array on seven 649 major ECM components, we observed that the MCF7 ESR1 mutant cell lines 650 consistently lacked adhesive properties on almost all ECM components with the 651 exception of fibronectin, and T47D ESR1 mutant cells displayed reduced adhesion on 652 collagen I, collagen II and fibronectin (Fig. 4B). Considering that collagen I is the most 653 abundant ECM component in ER+ breast cancer (Supplementary Fig. S13B), we 654 repeated the adhesion assay on collagen I (Fig. 4C & 4D; Supplementary Fig. S13C & 655 S13D) and similarly found reduced adhesion in both ER mutant cells. In an orthogonal 656 approach, we visualized and quantified adhesion in a co-culture assay on collagen I 657 using differentially labelled ESR1 WT and mutant cells, which confirmed significantly 658 decreased adhesive properties in the mutant cells (Supplementary Figure S13E & 659 S13F). Of note, ESR1 mutant adhesion deficiency on collagen I was also observed in 660 two additional ESR1 mutant models (Supplementary Fig. S13G).

We sought to investigate the molecular mechanisms underlying the unique defect of
collagen I adhesion in *ESR1* mutant cells. There was no consistent change in
expression of members of the integrin gene family, encoding well-characterized direct
collagen I adhesion receptors, in our cell line models (Supplementary Fig. S14A and

665 Supplementary Table S6). We therefore hypothesized that another gene critical in 666 regulation of ECM genes might be altered and to test this directly, we performed gene 667 expression analysis of 84 ECM adhesion-related genes using a qRT-PCR array 668 (Supplementary Table S8). Pairwise comparisons between each mutant cell line and 669 corresponding WT cells revealed a strong context-dependent pattern of ECM network 670 reprogramming, with more pronounced effects in MCF7 cells (Fig. 4E). Intersection 671 between Y537S and D538G mutants showed 23 and 1 consistently altered genes in 672 MCF7 and T47D cells, respectively (Fig. 4F). TIMP3, the gene encoding tissue 673 metallopeptidase inhibitor 3, was the only shared gene between all four mutant cell 674 models (Fig. 4F), and we confirmed its decreased expression at the mRNA (Fig. 4G & 675 Supplementary Fig. S14B) and protein level (Fig. 4H), as well as in other genome-676 edited *ESR1* mutant models (Supplementary Fig. S14C). E2 treatment represses 677 TIMP3 expression, suggesting that its downregulation in ESR1 mutant cells is likely due 678 to ligand-independent repressive ER activity (Supplementary Fig. S14C). 679 Downregulation of TIMP3 was found in several (but not all) tamoxifen resistant MCF7 680 models, but long-term estradiol deprived (LTED) MCF7 showed upregulation 681 (Supplementary Fig. S14D). Further, changes in TIMP3 were not seen in other LTED 682 models, suggesting that alteration of TIMP3 by mutant ESR1 is complex and warrants 683 further investigation. Overexpression of TIMP3 rescued the adhesion defect in ESR1 684 mutant cells (Figure 4I, 4J & Supplementary Fig. S14E), with no impact on cell 685 proliferation (Supplementary Fig. S14F). Collectively, these data imply a selective role 686 for TIMP3 downregulation in causing the decreased cell-matrix adhesion phenotype of 687 the ESR1 mutant cells, consistent with a critical role for TIMP3 in metastasis in other 688 cancer types (50,51).

689 Given the role of *TIMP3* as an essential negative regulator of matrix metalloproteinase 690 (MMP) activity , we compared MMP activity between *ESR1* WT and mutant cells. A pan-691 MMP enzymatic activity assay revealed significantly increased MMP activation in all 692 mutant cells (Fig. 4K & 4L), indicating that the *ESR1* mutant cells have increased 693 capacity for matrix digestion. This was validated in spheroid-based invasion assays in 694 which cells were embedded in collagen I (Fig. 4M) but without notable growth 695 differences (Supplementary Fig. S15A & S15B). This was additionally visualized in co696 culture spheroid invasion assays using differentially labelled T47D ESR1 WT and 697 mutant cells, which showed an enrichment of ESR1 mutant cells at the leading edge of 698 the spheroids (Supplementary Fig. S15C). Lastly, we tested if MMP blockade could 699 repress ESR1 mutant-modulated invasive and adhesive alterations. Marimastat 700 treatment substantially reduced the invasive phenotype of ESR1 mutant cells without 701 inhibiting growth (Fig. 4N, 4O & Supplementary Fig. S15D). Furthermore, the reduced 702 adhesive property was rescued by Marimastat treatment in ESR1 mutant cells 703 (Supplementary Fig. S15E). These data demonstrate that decreased TIMP3 expression, 704 resulting in increased MMP activation causes enhanced matrix digestion associated 705 with decreased adhesion to ECM, ultimately conferring invasive properties to ESR1 706 mutant cells.

707 *De novo* FOXA1-mediated Wnt pathway activation enhances of the T47D-D538G 708 cell migration

709 T47D D538G cells showed increased *in vivo* tumorigenesis despite showing less

- pronounced adhesive phenotypes compared to T47D Y537S and MCF7 Y537S/D538G
- 711 cells. Reasoning mutation and context-dependent metastatic activities of the mutant ER
- 712 protein and having identified "Cellular Movement" as another top hit in our initial
- pathway analysis of differentially expressed genes in *ESR1* mutant tumors (Fig. 1D), we
- assessed potential differences in cellular migration between the different models.
- 715 Wound scratch assays identified significantly increased cell motility in the T47D-D538G
- model (Fig. 5A & 5B), but not in T47D-Y537S (Fig. 5B) or MCF7 mutant cells
- 717 (Supplementary Fig. S16A & S16B). This enhanced motility was shared between the
- three individual T47D-D538G clones again excluding potential clonal artifacts
- 719 (Supplementary Fig. S16C & S16D). Furthermore, we observed a different morphology
- of T47D-D538G cells at the migratory leading edges (Fig. 5C) further confirmed by
- 121 larger and stronger assembly of F-actin filaments at the edge of T47D-D538G cell
- clusters (Supplementary Fig. S16E-S16H). To mimic collective migration from a cluster
- of cells, we utilized a spheroid-based collective migration assay on type I collagen (Fig.
- 5D). The distance to the leading edges of T47D-D538G mutant cells was significantly
- ⁷²⁵ longer compared to WT spheroids (Fig. 5E). In orthogonal approaches, enhanced

migratory capacities of T47D-D538G cells were observed in co-culture assay using
labelled T47D-WT and D538G cells (Supplementary Fig. S16I & S16J) and in Boyden
chamber transwell assays (Supplementary Fig. S16K & S16L). Finally, in T47D
overexpression models, we also observed significantly enhanced migration in D538G
compared to WT overexpressing cells (Supplementary Fig. S17A-S17E).

731 To understand the mechanisms underlying the migratory phenotype of T47D-D538G 732 cells we identified pathways uniquely enriched in these cells. GSEA identified endocrine 733 resistance-promoting pathways (e.g. E2F targets) in both T47D mutants, whereas Wnt-734 β -catenin signaling was one of the uniquely enriched pathways in T47D-D538G (Fig. 735 5F). Hyperactivation of the canonical Wnt- β -catenin pathway was further confirmed by a 736 Top-Flash luciferase assay (Supplementary Fig. S18A). We also observed increased 737 phosphorylation of GSK3 β and GSK3 α as well as β -catenin (both total and nuclear) 738 protein levels in T47D-D538G cells (Fig. 5G, Supplementary Fig. S18B and S18C). 739 Stimulation of T47D-WT cells with Wnt3A was not able to increase the migration to the 740 same level of D538G cells (Supplementary Fig. S18D), suggesting that Wnt activation is 741 a required but not sufficient factor in driving this phenotype. To address the potential 742 clinical relevance of these findings, we utilized the porcupine inhibitor LGK974, which 743 prevents the secretion of Wnt ligands and is currently being tested in a clinical trial for patients with advanced solid tumors including breast cancer (NCT01351103) (52,53). 744 745 Treatment with LGK974 resulted in a 20% and 40% inhibition of T47D ESR1 WT and 746 D538G mutant cell migration respectively (Fig. 5H and Supplementary Fig. S18E) yet 747 had no effect on cell proliferation (Supplementary Fig. S18F). We next studied the 748 combination of LGK974 and the selective ER degrader (SERD), Fulvestrant, in 749 migration assays, in which we detected significant synergy (Fig. 51), suggesting that 750 combination therapy co-targeting the Wnt and ER signaling pathways might reduce the 751 metastatic phenotypes of Wnt hyperactive ESR1 mutant tumors.

- 752 We sought to decipher the mechanisms underlying T47D-D538G Wnt hyperactivation.
- 753 First, a set of Wnt component genes were identified to be uniquely enriched in tumors
- with D538G but not other mutant variants in the DFCI cohort (Supplementary Fig.
- 755 S18G). Comparing the fold changes of canonical Wnt signaling positive regulators

756 between T47D-Y537S and T47D-D538G mutant cells, we identified eight candidate 757 genes exhibiting pronounced enrichment in T47D-D538G cells (Fig. 5J), including 758 ligands (e.g. WNT6A), receptors (e.g. LRP5) and transcriptional factors (e.g. TCF4). 759 With the exception of *LRP5*, none of these candidate genes were induced by E2 760 stimulation in T47D ESR1 WT cells (Supplementary Fig. S19A). Lack of consistent E2 761 regulation was confirmed in five additional ER+ breast cancer cell lines (Supplementary 762 Fig. S19B). Hence, we alternatively hypothesized that D538G ER might gain *de novo* 763 binding sites proximal to Wnt pathway genes allowing their induction. We mapped ER 764 binding globally by analyzing ER ChIP-sequencing in T47D WT and ESR1 mutant cells. 765 Consistent with previous studies (14,20), mutant ER were recruited to binding sites 766 irrespective of hormone stimulation (Supplementary Fig. S19C & Table S9). However, 767 none of the mutant ER bound regions mapped to identified Wnt pathway genes (± 50kb 768 of TSS), again suggesting a lack of direct canonical ER regulation. Moreover, short-term 769 fulvestrant treatment only weakly dampened T47D-D538G cell migration (Fig. 5K & 5M) 770 suggesting that ER activation may not be an essential prerequisite for enhanced cell

migration in D538G cells.

Given our recent findings of enriched FOXA1 motifs in gained open chromatin of T47D-

D538G cells (19), we decided to validate this pivotal *in silico* prediction, focusing on our

observed migratory phenotype. In contrast to the limited effects of ER depletion,

strikingly, FOXA1 knockdown fully rescued the enhanced migration in T47D-D538G

cells (Fig. 5L & 5N), indicating a more dominant role of FOXA1 in controlling T47D-

777 D538G cell migration. Ligand-independent 2D growth of T47D-D538G cells was

inhibited by both fulvestrant and FOXA1 knockdown (Supplementary Fig. S19D),

suggesting a canonical ER-FOXA1 co-regulatory mechanism in growth, distinguished

from the role of FOXA1 in the regulation of migration.

To further explore how FOXA1 contributes to the migratory phenotype, we performed

782 FOXA1 ChIP-sequencing to decipher the genomic binding profiles. We identified

approximately 30,000 peaks in T47D WT cells regardless of E2 stimulation and a ~1.6

fold increase in binding sites of the Y537S (61,934) and D538G (54,766) ER mutants

785 (Supplementary Fig. S20A & Supplementary Table S9). PCA distinctly segregated all

four groups (Fig. 5O), suggesting unique FOXA1 binding site redistribution. Comparison
of binding intensities revealed 14%, 28% and 21% FOXA1 binding sites were altered in
WT+E2, Y537S and D538G groups, respectively, with a predominant gain of binding
intensities in the two T47D mutants (Fig. 5P and Supplementary Fig. S20B).

790 Since FOXA1 is a well-known essential pioneer factor of ER in breast cancer, we 791 examined interplay between FOXA1 and WT and mutant ER. Interestingly, both Y537S 792 (39%) and D538G (25%) ER binding sites showed a significantly lower overlap between 793 FOXA1 compared to the WT+E2 group (56%), albeit with the increased number of 794 gained mutant FOXA1 binding sites (Supplementary Fig. S20C). This discrepancy suggests that FOXA1 exhibits a diminished ER pioneering function and instead might 795 796 contribute to novel functions via gained *de novo* binding sites. Co-occupancy analysis 797 using isogenic ATAC-seq data (19) uncovered that the open chromatin of T47D-D538G 798 cells was more associated with FOXA1 binding sites compared to WT and T47D-Y537S 799 cells (Fig. 5Q). FOXA1 binding intensities were also stronger in D538G ATAC-sites 800 (Supplementary Fig. S20D). Collectively, these results provide evidence that FOXA1 801 likely plays a critical role in the D538G mutant cell to reshape its accessible genomic 802 landscape.

803 We further investigated the impact of the gained FOXA1-associated open chromatin on 804 transcriptomes, particularly exploring ESR1 mutant-specific genes. Intersection of the 805 gained FOXA1- and ATAC-sites for annotated T47D-D538G genes with non-canonical 806 ligand-independence identified 25 potential targets that could be attributed to de novo 807 FOXA1 bound open chromatin, exemplified by *PRKG1* and *GRFA* as top targets (Fig. 808 5R & Supplementary Fig. S21A). Notably, one of our identified D538G specific Wnt 809 regulator genes, TCF4, was uncovered in this analysis. Higher TCF4 expression in 810 T47D-D538G cells was validated by gRT-PCR and furthermore this increased 811 expression could be fully blocked following FOXA1 knockdown (Supplementary Fig. 812 S21B). Additionally, stronger FOXA1 recruitment at the TCF4 gene locus was validated 813 via ChIP-gPCR (Supplementary Fig. S21C and S21D). Importantly, overexpression of 814 dominant negative *TCF4* strongly impaired cell migration in T47D-D538G, while it only 815 slightly affected WT cells (Fig. 5S). Together, these results support that FOXA1 binding

- 816 site redistribution leads to novel chromatin remodeling and enhanced expression of
- 817 genes with roles in metastases including *TCF4*, which subsequently activate Wnt-driven
- 818 migration in T47D-D538G cells.

819

820 Discussion

821 Hotspot somatic mutations clustered in the LBD of ER represent a prevalent molecular 822 mechanism that drives antiestrogen resistance in ~30% of advanced ER+ breast 823 cancer. There is an urgent need for a deeper understanding of this resistance 824 mechanism in order to develop novel and personalized therapeutics. Utilizing clinical 825 samples, in silico analysis of large datasets, and robust and reproducible 826 experimentation in multiple genome-edited cell line models, our study uncovers complex 827 and context-dependent mechanisms of how ESR1 mutations confer gain-of-function 828 metastatic properties. We identified ESR1 mutations as multimodal metastatic 829 modulators hijacking adhesive and migratory networks, and thus likely influencing 830 metastatic pathogenesis and progression. Mechanistically, we uncovered novel ER-831 indirect regulation of metastatic candidate gene expression, distinct from previously 832 described (11,12,54) canonical ligand-independent gene induction. Nonetheless, some 833 limitations were noted in our study, such as the lack of *in vivo* validation of studied 834 therapeutic approaches and lack of proposed target validation in clinical specimens. In 835 addition, our numbers for clinical samples of paired primary-metastatic tumors harboring 836 ESR1 mutations is finite, necessitating validation in future studies with larger clinical 837 cohorts.

838 We discovered enhanced cell-cell adhesion via upregulated desmosome and gap 839 junction networks in cell lines and clinical samples with ESR1 mutations. These 840 transcriptional alterations are associated with a specific clinical phenotype characterized 841 not only by treatment resistance, but also by high CTC count and a different metastatic 842 organotropism (55,56). We propose that this key alteration may support increased 843 metastases in ER mutant tumors through facilitating the formation of homo- or 844 heterotypic CTC clusters, providing a favorable environment for CTC dissemination, as 845 previously described (34). This idea is further supported by previous data showing 846 upregulation of the desmosome gene plakoglobin (JUP) which may play a role in a CTC 847 cluster formation signature (34). We observed increased expression of plakophilin, 848 desmocollin, and desmoglein in ESR1 mutant cells, suggesting the importance of the 849 broad desmosome network reprogramming for functional cell clustering activity.

850 Moreover, enhanced gap junction genes might potentiate intercellular calcium signaling, 851 facilitating the prolonged survival of various metastatic cell types tethered to ESR1 852 mutant cells *en route* (57). Dissociation of CTC cluster using Na+/K+ ATPase inhibitors 853 decreased metastasis in vivo (41). In addition, previous studies have validated the anti-854 tumor effects of FDA-approved gap junction blockers carbenoxolone in vivo (58). Our 855 results warrant additional preclinical studies using drugs targeting desmosome and gap 856 junctions, with the ultimate goal of applying these treatments in a CTC-targeted clinical 857 trial to improve outcomes for patients harboring breast cancers with *ESR1* mutations.

858 Previous studies using similar ESR1 mutant cell models described enhanced migratory 859 properties (15,16), but no mechanistic explanations were uncovered. Here we identify a 860 critical role for Wnt- β -catenin signaling and show that co-targeting of Wnt and ER 861 resulted in synergistic inhibition of cell migration. Intriguingly, the strong effect we 862 observed on migration was unique to T47D-D538G cells, a discovery that was made 863 possible through our use of multiple genome-edited mutation models. This finding might 864 help explain the higher frequency of D538G mutations in metastatic samples, despite 865 the stronger endocrine resistance phenotype of Y537S mutation (5,12,14,33). Markedly, 866 although we highlighted the up-regulation of TCF4 as an outcome of de novo FOXA1 867 reprogramming, it is plausible that other increased Wnt regulators including receptors 868 (e.g. LRP5) and ligands (e.g. WNT6A) are also associated with the migratory 869 phenotypes. Hence LGK974, a Wnt secretion inhibitor, could efficiently abrogate this 870 phenotype. Of note, slightly higher Wnt activity and β -catenin accumulation were also 871 observed in T47D-Y537S cells, but this failed to convert into a migratory phenotype. It is 872 possible that some genes uniquely regulated by Y537S ER in T47D cells might inhibit 873 migratory phenotypes. For instance, the gap junction component, connexin 43, which is 874 exclusively upregulated in T47D-Y537S cells, has been reported to play an inhibitory 875 role in epithelial cell migration (59). Furthermore, the unique observation in T47D rather 876 than MCF7 cell line may be in part explained by the lower basal migratory property and 877 basal Wnt activation in the T47D cell line, which might allow additional gain of function. 878 MCF7 WT cells showed approximately four-fold higher wound closure ratio than T47D 879 at 72 hours (Fig. 5B and Supplementary Fig. S16A) and furthermore it expresses 880 truncated mutant version of LRP5 (60), which confers constitutive Wnt signaling

881 activation. In vivo experiments revealed enhanced metastasis in the MCF7-Y537S but 882 not D538G model. This discrepancy with the *in vitro* data could possibly be explained 883 by the longer distant metastatic latency requirement of D538G cells in vivo, consistent 884 with a recent study using overexpression cell models (14). Alternatively, it is also 885 plausible that Y537S mutant cells exhibit stronger in vivo outgrowth at metastatic sites. 886 Further in vivo metastatic experiments in the absence and presence of E2 are 887 warranted to delineate the reason. These data support strong allele and context 888 dependent effects of the ESR1 mutation on metastatic phenotypes, in line with context 889 dependent effects on transcriptome, cistromes and accessible genome in ESR1 mutant 890 cells (11,12,14,19). Of note, previous efforts using multiple cell line models with ESR1 891 mutations elucidated several congruent molecular and functional alterations associated 892 with endocrine resistance (14,15,54), suggesting that mechanisms underlying 893 metastasis of ESR1 mutant clones exhibit a higher degree of heterogeneity. This is also 894 supported by clinical data: the recent BOLERO2 trial showed significant differences in 895 overall survival and everolimus response between Y537S and D538G mutations (9). and results from the recent PALOMA3 trial suggest a potential Palbociclib resistance 896 897 uniquely gained in tumors bearing the Y537S mutation (61). Given our model are limited 898 to MCF7 and T47D cells, there's a pressing need to establish additional ESR1 mutant 899 models with different background to follow-up on our observation and to perform further 900 pre-clinical investigations. Taken together, these proof-of-concept studies are setting 901 the stage for a more contextual and personalized therapeutic targeting strategy in ESR1 902 mutant breast cancer.

903 Of note, our comprehensive clinical investigation from five different cohorts (N=996) 904 suggest that ESR1 mutations more common in distant compared to local recurrences. 905 which we propose is due to gain-of-function of ESR1 mutant clones ie those cells are 906 more equipped to escape from the local-regional microenvironment. However, there are 907 some limitations to our study. First, it is challenging to differentiate local recurrences 908 from secondary primary tumors, limiting our analysis. Second, in some of the cohorts 909 we observed significant differences in number of lines of therapy and time to recurrence 910 comparing patients i) with ESR1 WT vs mutant tumors, and ii) with local and distance 911 recurrences. Although these analyses are limited by different numbers of cases, and

data that are available, nevertheless, they suggest that lack of *ESR1* mutation in local
recurrences could at least in part be also due to differences in prior therapies as
previously reported (6). Of note, a recent study identified hotspot *ESR1* mutations in 15
out of 41 (36%) of local-regional ER+ recurrences albeit at significantly lower mutation
allele frequencies (62). Given our data presented in this study, together with prior data
(14-16), we propose that *ESR1* mutations can facilitate metastatic spread although they
might not be sufficient to function as genetic drivers for such events.

919 Lastly, we also sought to address the ER regulatory mechanisms involved in induction 920 of candidate metastatic driver genes utilizing ChIP-seq technology. Interestingly, none 921 of the metastatic candidate genes in ESR1 mutant cells gained proximal ER binding 922 sites. This could be a result of our stringent hormone deprivation protocol resulting in 923 depletion of weaker binding events, and thus less sensitive binding site readouts. This 924 idea is supported by ChIP-seq data from Harrod et al. (20), which shows stronger ER 925 binding sites around DSC2, DSG2 and TIMP3 gene loci in MCF7-Y537S cells. Our 926 data, however, clearly shows that ER mutant cells display changes in indirect gene 927 regulation, resulting in metastatic phenotypes. This observation is due to non-canonical 928 ER action on chromatin structure remodeling, which was alternatively validated from our 929 ATAC-seg and FOXA1 ChIP-seg data. We propose that mutant ER reprograms FOXA1, 930 resulting in redistribution of FOXA1 binding to specific enhancers controlling the key 931 migratory driver gene(s). It's also likely that mutant ER can impact FOXA1 occupancy 932 by cooperating with other known epigenetic regulators such as GATA3 (63). In addition, 933 *ESR1* mutations might alter the expression of several important histone modifiers such 934 as KDM5B and KMT2C which showed expression changes in *ESR1* mutant cells. 935 Alteration of histone writers or erasers may reshape global H3K4 methylation and thus 936 differentially recruit FOXA1 (64). These mechanisms warrant future investigation. In 937 addition, several recent studies uncovered the promising role of androgen receptor (AR) 938 in *ESR1* mutant tumors and cell models (18,65), and additional studies are warranted to 939 study de novo interplay between FOXA1, AR and mutant ER. 940 Overall, our study serves as a timely and important preclinical report uncovering 941 mechanistic insights into ESR1 mutations that can pave the way towards personalized

942 treatment of patients with advanced metastatic breast cancer.

943

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1130 Table1

Cohorts	Site of	Total	ESR1	ESR1	Fisher's
	Recurrence	Number	WT	Mutant	Exact p
METAMORPH/POG570/	Distant	877	722	155 (18%)	0.0006
MSKCC/IEO Merged			(82%)		
	Local	44	44	0 (0%)	
			(100%)		
WCRC/Charite	Distant	48	36 (75%)	12 (25%)	0.0031
	Local	27	27	0 (0%)	
			(100%)		

1131

1132 Table Legend

Table 1. Significant enrichment of *ESR1* mutations in distant compared to local
recurrences.

1135 Upper panel: Data from 877 distant metastatic and 44 local recurrence cases were

merged from three cohorts (METAMORPH, 39 distant/9 local; POG570, 86 distant/14

local; MSKCC, 716 distant/8 local; IEO, 36 distant/13 local). ESR1 mutation status was

1138 previously identified by whole exome sequencing (METAMORPH), whole genome

sequencing (POG570) or target panel DNA sequencing (MSKCC, IEO). Lower panel: 48

1140 distant ER positive metastases and 27 local ER positive recurrences were obtained

1141 from the WCRC and Charite cohorts. Genomic DNA (gDNA) was isolated from either

1142 FFPE or frozen tumor tissues, and subjected to droplet digital PCR (ddPCR) detection

1143 with specific probes against Y537S, Y537C, Y537N and D538G hotspot point mutations

1144 (cDNA rather than gDNA was used for 3 of the local recurrent samples). Hotspot ESR1

1145 mutation incidences between distant metastatic and local recurrent samples in both

1146 panels were compared using a Fisher's exact test.

1147

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1148 Figure legends

1149

1150 Figure 1. Transcriptomic landscape of *ESR1* mutant metastatic breast cancers.

- 1151 A. Schematic overview of transcriptomic analysis of four ER+ metastatic breast cancer
- 1152 cohorts.
- 1153 B. Box plots representing the enrichment levels of "Estrogen Response Early" and
- 1154 "Estrogen Response Late" signatures in *ESR1* mutant versus *ESR1* WT metastatic
- 1155 tumors in each cohort. (WCRC, 46 *ESR1* WT/8 mutant; MET500, 34 *ESR1* WT/12
- 1156 ESR1 mutant; DFCI, 98 ESR1 WT/32 mutant; POG570, 68 ESR1 WT/18 mutant). Four
- 1157 quantiles are shown in each plot. Mann-Whitney U test was used to compare the
- enrichment of the signatures in WT and mutant tumors. (* p<0.05, ** p<0.01)
- 1159 C. Volcano plots representing the differentially expressing genes (DE genes) in ESR1
- 1160 mutant tumors versus WT tumors in the three metastatic breast cancer cohorts. DE
- genes were selected using the cutoff of FDR<0.1 and |log₂FC|>1.5. Genes that were
- 1162 upregulated or downregulated were labelled in red and blue respectively with
- 1163 corresponding counts.
- 1164 D. Dot plots showing the top 5 altered cellular and molecular functional categories
- derived from DE genes analysis using Ingenuity Pathway Analysis software. Specific
- 1166 sub-functions within overarching categories are presented as individual dots.
- 1167 Consistently altered pathways across all four cohorts are indicated in red.
- 1168 E. Stacked bar plot showing the distribution of 14 hotspot *ESR1* mutations identified in
- 1169 six independent cohorts using unbiased DNA sequencing approaches. Specific sample
- 1170 numbers were indicated in the plots. Variants with percentages above 1% were labelled
- 1171 on the top of each bar.
- 1172 F. Scatterplot representing enrichment level distribution of 50 hallmark gene sets in 10
- 1173 Y537S and 8 D538G metastatic tumors (after being normalized against 98 WT
- 1174 counterparts) from the DFCI cohort. Top enriched pathways from each quartile are
- 1175 labelled.
- 1176
- 1177 Figure 2. *ESR1* mutant cells exhibit stronger cell-cell adhesion.

1178 A. Representative images of day 6 hormone deprived MCF7 and T47D spheroids 1179 seeded in 6-well ultra-low attachment (ULA) plates. Images were taken under 1.25x 1180 magnification. Representative experiment from three independent repeats is shown. 1181 B. Bar plot representing day 7 cell numbers of MCF7 or T47D WT and ESR1 mutant 1182 cells seeded into flat bottom ULA plates. Cell abundance were quantified using Celltiter 1183 Glo. Fluorescence readouts were corrected to background measurements. Each bar 1184 represents mean ± SD with 10 (MCF7) or 6 (T47D) biological replicates. Representative 1185 experiment from six independent repeats is shown. Dunnett's test was used between WT and each mutant. (** p<0.01) 1186

1187 C. Left panel: A calcein labelled cell-cell adhesion assay was performed in MCF7 WT 1188 and mutant cells. Adhesion ratios were calculated by dividing the remaining cells after 1189 each wash to the initial readout from unwashed wells. A pairwise two-way ANOVA 1190 between WT and each mutant was utilized. Each point represents mean ± SD with five 1191 biological replicates. Representative experiment from 17 independent repeats is shown. 1192 Right panel: Adhesion ratios after three washes were extracted from 17 independent 1193 experiments displayed as mean ± SEM. Dunnett's test was used to compare between 1194 WT and each mutant. (* p<0.05, ** p<0.01) 1195 D. Line plot representing the aggregation ratio of MCF7 cells seeded into round bottom 1196 ULA plates. Cell aggregation processes were followed by the IncuCyte living imaging

1197 system every hour. Spheroid areas were normalized to time 0. Each dot represents

1198 mean ± SD with eight biological replicates. Representative images after 3 hours of

aggregation are shown across the top panel. Images were captured under 10x

1200 magnification. Representative experiment from five independent repeats is shown. A

pairwise two-way ANOVA between WT and each mutant was utilized. (** p<0.01)

1202 E. Representative images of MCF7 cell cluster status after two hours of flow under

1203 physiological shear stress produced by the ibidi microfluidic system. Images were taken

1204 under 10x magnification. A regional 2x zoom in is presented on the top of each image.

1205 Representative experiment from three independent repeats is shown.

1206 F. Bar graph representing the percentage of MCF7 cells in a cluster based on the

1207 quantification of cluster and single cell numbers from 12 representative images per

1208 group. Each bar represents mean ± SD. Cell cluster ratios after 2 hours of flow were

- 1209 further normalized to time 0 to correct for baseline pre-existing clusters. Representative
- 1210 experiment from three independent repeats is shown. Dunnett's test was used between
- 1211 WT and mutant cells. (** p<0.01)
- 1212 G. Bar plots showing the cluster size distribution of MCF7 cells after normalization to
- 1213 time 0. Each bar represents mean ± SD from 12 representative images per group.
- 1214 Representative experiment from three independent repeats is shown. Dunnett's test
- 1215 was used between WT and each mutant cell type within the same cluster size category.
- 1216 (** p<0.01).
- 1217 H. Schematic overview of short-term *in vivo* circulating tumor cell evaluation
- 1218 experimental procedure.
- 1219 I. Left panel: Representative images of two-cell clusters (WT) and a multicellular cluster
- 1220 (Y537S). Images were taken under 40x magnification. Right panel: Stacked bar chart
- 1221 representing the distribution of cancer cells in each cluster type. This experiment was
- 1222 performed once. Fisher's exact test was applied to test whether multicellular clusters
- were enriched in *ESR1* mutant cells. (** p<0.01)
- 1224 J. Left panel: Representative images of a WT and Y537S two cell cluster. Lines
- 1225 connecting the two nuclei centers were indicated. Images were taken under 40x
- 1226 magnification. Right panel: Dot plot represents the inter-nuclei distance of all two-cell
- 1227 clusters in MCF7 WT and mutant cells. Measured distances were normalized to the
- 1228 average radius of both cells of this cluster size to avoid cell size bias. This experiment
- 1229 was performed once. Mann-Whitney U test was performed between WT and each
- 1230 mutant cell. (** p<0.01)
- 1231 K. Schematic overview of *in vivo* metastatic evaluation of *ESR1* mutant cells introduced
- 1232 via tail vein injections.
- 1233 L. Representative H&E staining images the tumorous portion of MCF7-Y537S induced
- 1234 macro-metastatic (macro-met) tumors from 3 different mice. This experiment was
- 1235 performed once. Images were taken under 20x magnification.
- 1236 M. Left panel: Dot plots showing the number of macro-met per mouse from MCF7 ESR1
- 1237 WT and mutant cells-injected mice. Pairwise Mann-Whitney U test was used to
- 1238 compare the macro-met numbers in each mutant group to WT cell-injected groups.
- 1239 Right panel: Quantification of lung micro-met areas based on human specific CK19

- 1240 staining quantification. This experiment was performed once. Pairwise Mann-Whitney U
- 1241 test was applied for statistical analysis. (WT, n=7; Y537S, n=6; D538G, n=7) (* p<0.05)
- 1242 N. Representative images of micro-metastatic loci on the lung sections of T47D-ESR1
- 1243 mutant cell-injected mice. Images were taken under 10x magnification. Metastatic loci
- were indicated with white arrow. This experiment was p once. (WT, n=7; Y537S, n=6;
- 1245 D538G, n=7) (Blue: nuclei; Red: CK8+18; Green: Human specific CK19)
- 1246 O. Left panel: Dot plots showing the macro-metastatic counts per mouse from T47D
- 1247 *ESR1* mutant-injected mice. Pairwise Mann-Whitney U test was used to compare the
- 1248 macro-met numbers in each mutant group to WT cell-injected groups. Right panel:
- 1249 Quantification of lung micro-met areas based on CK19 staining and was performed in a
- 1250 blind manner. This experiment was performed once. Pairwise Mann-Whitney U test was
- 1251 applied for statistical analysis. (N=1, * p<0.05)
- 1252 P. Representative images of CTCs clusters detected through the CellSearch Platform
- 1253 after EpCAM dependent enrichment (Pink: nuclei, Green: CK8/CK18/CK 19). Image
- 1254 resolution and magnification were achieved in accordance with the CellSearch Platform.
- 1255 Q. Mosaic plot showing the association between *ESR1* genotype status and clustered
- 1256 CTCs. A significant positive association was observed by Fisher's exact test between
- 1257 *ESR1* mutations and high clustered CTCs (clustered CTCs > 4).
- 1258 R. Kaplan Meier plot representing the impact of clustered CTCs in terms of Overall
- 1259 Survival (OS). Patients with clustered CTCs > 4 experienced the worse prognosis in
- 1260 terms of OS both with respect to those without clusters (both stage IV indolent and
- stage IV aggressive) and those with clusters but with \leq 4 clustered CTCs (P < 0.0001).
- 1262 Patients at risk are reported at each time point. Log rank test was to compare the
- 1263 survival curves of the two patient subsets.
- 1264

Figure 3. Desmosome and gap junction adhesome reprogramming confers enhanced adhesive properties in *ESR1* mutant cells.

- 1267 A. Gene Set Variation Analysis (GSVA) scores of desmosome and gap junction gene
- 1268 sets enrichment in MCF7 and T47D ESR1 mutant vs WT cell RNA-seq data sets. Each
- 1269 cell type has four biological replicates. Dunnett's test was used to test the significance
- 1270 between WT and mutant cell lines. (** p<0.01)

1271 B. Heatmaps showing all desmosome and gap junction component genes in MCF7 and

- 1272 T47D *ESR1* mutant cells. Data were extracted from RNA-sequencing results with four
- 1273 biological replicates. Color scale represents the Log2 fold changes in each mutant
- 1274 normalized to WT counterparts using the log₂(TPM+1) expression matrix. Genes with
- 1275 counts=0 in more than one replicate in each cell type were filtered out of analysis.
- 1276 Genes with a log₂FC>1.2 and a p<0.05 in at least one group are labelled in red.
- 1277 C. Western blot validation of the expression level of DSG2, DSC1, PKP1, Cx43 and
- 1278 Cx26 in MCF7 WT and *ESR1* mutant cells after hormone deprivation. Tubulin was
- blotted as a loading control. Representative blots from three independent repeats wasshown for each protein.
- 1281 D. qRT-PCR validation of selected altered candidate desmosome and gap junction
- 1282 genes in MCF7 *ESR1* mutant cells. $\Delta\Delta$ Ct method was used to analyze relative mRNA
- 1283 fold changes normalized to WT cells and *RPLP0* levels were measured as an internal
- 1284 control. Each bar represents mean ± SD with biological triplicates. This experiment was
 1285 a representative from four independent repeats. Dunnett's test was used to compare the
- 1286 gene expression between WT and each mutant. (* p<0.05, ** p<0.01)
- 1287 E. Representative images of immunofluorescence staining showing the distribution of
- desmoglein 2 (DSG2) in MCF7 WT and ESR1 mutant cells. Images were taken under
- 1289 20x magnification. A 2x zoom in of each image is presented. Right lower panel: *DSG2*
- 1290 signal intensities were quantified and normalized to cell numbers in each image. Data
- 1291 from 20 regions within the collected images were combined from four independent
- 1292 experiments. Mean ± SD is presented in each plot. Dunnett's test was used to test the
- significance between WT and mutant cells. (** p<0.01)
- 1294 F. Box plots representing GSVA scores of the enrichment of the top desmosome and
- 1295 gap junction candidate genes (genes with log₂FC>2 in at least one mutant line) in
- 1296 patient matched primary-metastatic paired samples. Delta GSVA score of each sample
- 1297 was calculated by subtracting the scores of primary tumors from the matched metastatic
- 1298 tumors. Four quantiles are shown in each plot. Mann-Whitney U test was performed to
- 1299 compare the Delta GSVA scores between *ESR1* WT (n=44) and mutation (n=7)
- 1300 harboring tumors. (* p<0.05)

1301 G & J. Representative images of cell cluster status after two hours of flow under 1302 physiological shear stress in the ibidi microfluidic system, with or without 300µM of the 1303 desmosomal blocking peptide (G) or 100µM of carbenoxolone (J) treatment. Images 1304 were taken under 10x magnification. This experiment was a representative from two 1305 (desmosome peptide treatment) and three (CBX treatment) independent repeats. 1306 H & K. Bar graphs representing the T0 normalized percentage of cells in cluster status 1307 after quantification of cluster and single cell numbers under each treatment. Each bar 1308 represents mean ± SD quantified from 12 images per group. This experiment was a 1309 representative from two (desmosome peptide treatment) and three (CBX treatment) 1310 independent repeats. Student's t test was used to examine the effects of treatment 1311 between each group's cluster ratio. (** p<0.01) 1312 I & L. Bar graphs representing the T0 normalized 2 cell and greater than 5 cell cluster 1313 percentages under each treatment. Each bar represents mean ± SD quantified from 12

images per group. This experiment was a representative from two (desmosome peptide

1315 treatment) and three (CBX treatment) independent repeats. Pairwise student's t test

1316 was used to examine the effects of treatment between each group's cluster ratio. (**

1317 p<0.01)

1318 M. Bar graphs representing gRT-PCR measurement of DSC1, DSC2, GJA1, GJB2 and 1319 GJB5 mRNA levels in MCF7 WT and ESR1 mutant cells following siRNA knockdown of 1320 ESR1 for 7 days. $\Delta\Delta$ Ct method was used to analyze relative mRNA fold changes 1321 normalized to WT cells and RPLP0 levels were measured as an internal control. Each 1322 bar represents mean ± SD with three biological replicates. Representative experiment 1323 from three independent repeats is displayed. Student's t test was used to compare the 1324 gene expression between scramble and knockdown groups of each cell type. (* p<0.05, 1325 ** p<0.01)

1326 N & O. Western blot validation of the expression level of ER, Cx43 and cFOS in MCF7

1327 WT and *ESR1* mutant cells after seven days of *ESR1* knockdown (N) or three days of

1328 20µM T-5224 treatment (O). Tubulin was blotted as a loading control. Representative

1329 blot from three (N) and five (O) independent repeats is displayed.

1330 P. Screen shot of H3K27ac and H3K4me2 binding peaks at proximity to genomic DSC1

and DSG1 loci in MCF7 parental cells. ChIP-seq data were visualized at WashU

- 1332 Genome Browser based on public available data set from ENCODE (H3K4me2:
- 1333 ENCSR875KOJ ; H3K27ac: ENCSR752UOD). Y axis represents the binding intensity of
- each ChIP-seq data set. Selected peaks for ChIP-qPCR assessment in Q wereindicated.
- 1336 Q. Bar graph showing the fold enrichment levels of the two active histone modification
- 1337 markers at the two selected peaks around *DSC1* and *DSG1* gene loci illustrated in P.
- 1338 Each bar represents mean ± SD from biological triplicates. Fold enrichment levels were
- 1339 calculated by normalizing to IgG controls and further normalized to WT levels. This
- 1340 experiment is representative from two independent repeats. Dunnett's test was used
- 1341 within each group. (N=2, * p<0.05, ** p<0.01)
- 1342

Figure 4. *ESR1* mutant cells show diminished ECM adhesion and enhanced invasion via an altered *TIMP3*-MMP axis.

- 1345 A. Gene set enrichment plots showing the comparison of enrichment levels of the
- 1346 "KEGG ECM Receptor Interaction" gene set (MSigDB, M7098) between WT and mutant
- 1347 tumors in DFCI cohort. (98 *ESR1* WT and 32 mutant tumors)
- 1348 B. Heatmap representation of adhesion ratio on 7 ECM components performed with
- 1349 MCF7 and T47D *ESR1* WT and mutant cells. Adhesion ratio of each condition with
- 1350 biological quadruplicates was quantified by dividing the number of remining cells after
- 1351 washing to the original total cells plated. All data was further normalized to WT cells
- 1352 within each cell line. This experiment was performed once. Dunnett's test was applied to
- 1353 each condition of each cell line. (* p<0.05, **p<0.01)
- 1354 C. Representative images *ESR1* WT and mutant cells remaining on collagen I after
- three PBS washes. Images were taken using 4x magnification. Experiment displayed isrepresentative from three independent repeats.
- 1357 D. Quantification of adhesion ratios on collagen I in each cell type. Bar graphs represent
- 1358 the mean ± SD with four biological replicates in each group. Dunnett's test was utilized
- 1359 within each cell line to compare WT and mutant adhesion ratios. Experiment displayed
- 1360 is representative from 12 (MCF7) and 11 (T47D) independent repeats. (* p<0.05, **
- 1361 p<0.01)

- 1362 E. Volcano plots showing the alterations of 84 ECM adhesion genes in all mutant cell
- 1363 types in a pairwise comparison to the WT counterparts. Genes were pre-filtered with an
- 1364 average Ct<35 in at least one group. An FDR<0.1 was considered as a significantly
- 1365 altered gene in *ESR1* mutant cells. Overlapping downregulated (blue) or upregulated
- 1366 (red) genes between the two mutants of each cell line were further highlighted, with
- 1367 gene name labels for the top targets. Top changed genes in each T47D mutant cells
- 1368 were labelled in green. This experiment was performed once.
- F. Venn diagrams showing the consistently differentially expressed genes between thetwo mutant variants within each cell line. *TIMP3* was highlighted as the only overlapping
- 1371 gene in all four *ESR1* mutant cell types.
- 1372 G. qRT-PCR validation of *TIMP3* expression in WT and *ESR1* mutant cells. Ct values
- 1373 were normalized to *RPLP0* and further normalized to WT cells. Bar graphs represent
- 1374 the mean ± SD with biological triplicates in each group. Representative experiment from
- 1375 seven independent repeats is shown. Dunnett's test was utilized within each cell line. (*
 1376 p<0.05, ** p<0.01)
- 1377 H. Western blot validation of *TIMP3* from whole cell lysates after hormone deprivation.
- 1378 Tubulin was used as a loading control. Representative experiment from six independent1379 repeats is shown.
- 1380 I & J. Quantification of adhesion ratios on collagen I in each mutant variant following
- 1381 transfection of pcDNA empty vector or *TIMP3* plasmids in MCF7 (I) and T47D (J) cell
- 1382 models. Bar graphs represent the mean ± SD from 5 (MCF7) and 7 (T47D) biological
- 1383 replicates. Representative experiment from four independent repeats is shown.
- 1384 Student's t test was used to compare the empty vector and *TIMP3* overexpressing
- 1385 groups. (* p<0.05, ** p<0.01)
- 1386 K & L. Graphical view of pan-MMP FRET kinetic assay. MMPs in MCF7 (K) and T47D
- 1387 (L) cell lysates were pre-activated and mixed with MMP substrates. Fluorescence was
- 1388 measured in a time course manner and normalized to T0 baseline and further
- 1389 normalized to WT cell readouts. Each point represents the mean ± SD value from three
- 1390 biological replicates. Representative experiment from four independent repeats is
- 1391 shown. Pairwise two-way ANOVA between WT and each mutant cell type was
- 1392 performed. (* p<0.05, ** p<0.01)

- 1393 M. Top panel: Representative images of the spheroid-based collagen invasion assay in
- 1394 ESR1 WT and mutant cell models. MCF7 and T47D spheroids were mixed in collagen I
- 1395 for 4 and 6 days, respectively. Bright field images were taken accordingly with 10x
- 1396 magnification. Bottom panel: Quantification of invasive areas within images. Invasive
- 1397 areas were calculated by subtracting each original spheroid area from the
- 1398 corresponding endpoint total area. Each bar represents mean ± SD with 10 biological
- 1399 replicates. Experiments displayed are representative from three independent repeats
- 1400 from each cell line. Dunnett's test was used to compare the difference between WT and
- 1401 mutant cells. (* p<0.05, ** p<0.01)
- 1402 N. Representative images of the spheroid-based collagen invasion assay with 10 μ M of
- 1403 Marimastat treatment in MCF7 (Top panel) and T47D (Lower panel) cell models for 4
- 1404 and 6 days, respectively. Images were taken under 10x magnification. Experiment was
- 1405 performed with 20 biological replicates for once.
- 1406 O. Quantification of corresponding invasive areas from N. Student's t test was used to
- 1407 compare the effects of Marimastat treatment to vehicle control. (** p<0.01)
- 1408

Figure 5. *De novo* FOXA1-mediated Wnt pathway activation enhances migratory
property of the T47D-D538G cells.

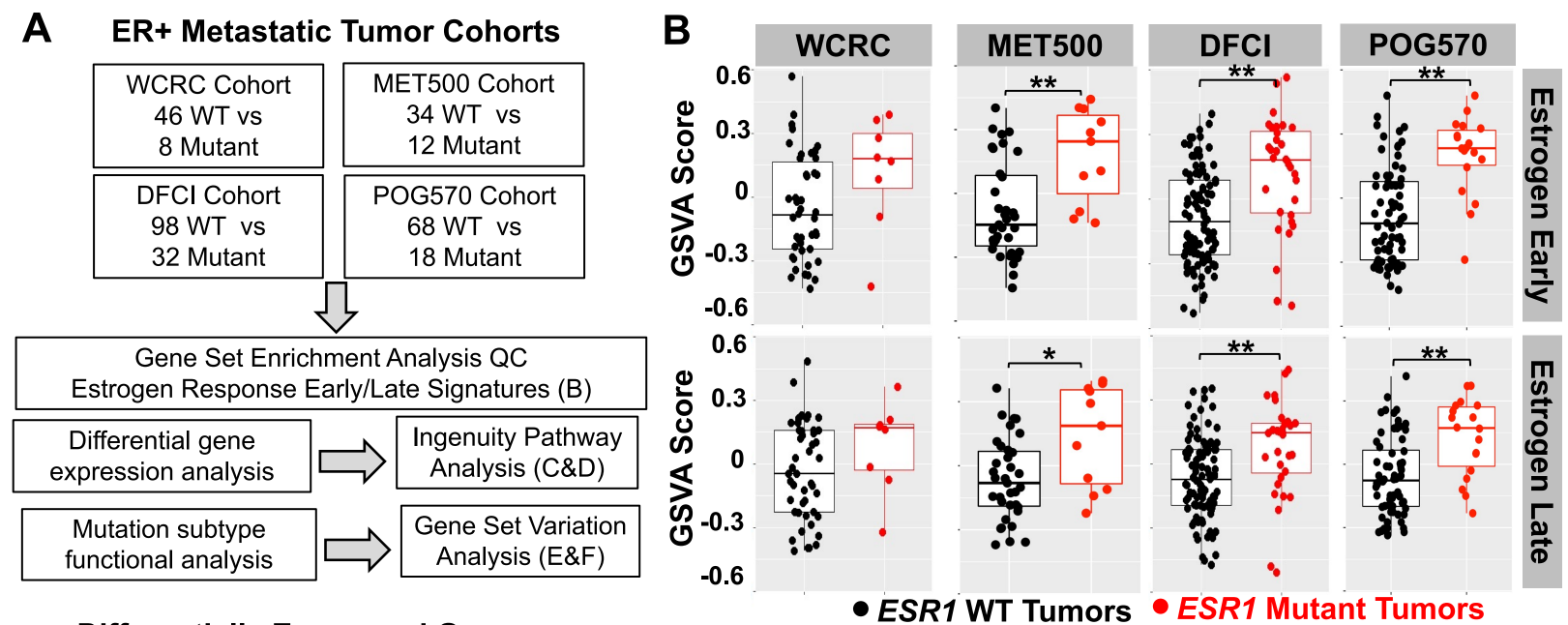
- 1411 A & B. Representative images (A) and quantification (B) of wound scratch assay of
- 1412 T47D WT and ESR1 mutant cells performed using IncuCyte living imaging system over
- 1413 72 hours. The migratory region normalized to T0 are labelled in blue. Images were
- 1414 taken under 10x magnification. Cell migration rates were quantified based on relative
- 1415 wound densities with 8 biological replicates. Representative experiment from 11
- 1416 independent repeats is shown. Pairwise two-way ANOVA between WT and each mutant
- 1417 was performed. (** p<0.01)
- 1418 C. Representative magnified images of the migratory edge of each group in wound
- 1419 scratch assays in A.
- 1420 D & E. Representative images (D) and quantification (E) of spheroid collective migration
- 1421 assays in T47D mutant cells. T47D cells were initially seeded into round bottom ULA
- 1422 plates to form spheroids, which were then transferred onto collagen I coated plates.
- 1423 Collective migration was measured after 4 days. The migratory edge of each spheroid is

- 1424 circled with a white line. Migratory distances were calculated based on the mean radius
- 1425 of each spheroid normalized to corresponding original areas. Representative
- 1426 experiment from three independent repeats is shown. Dunnett's test was used for

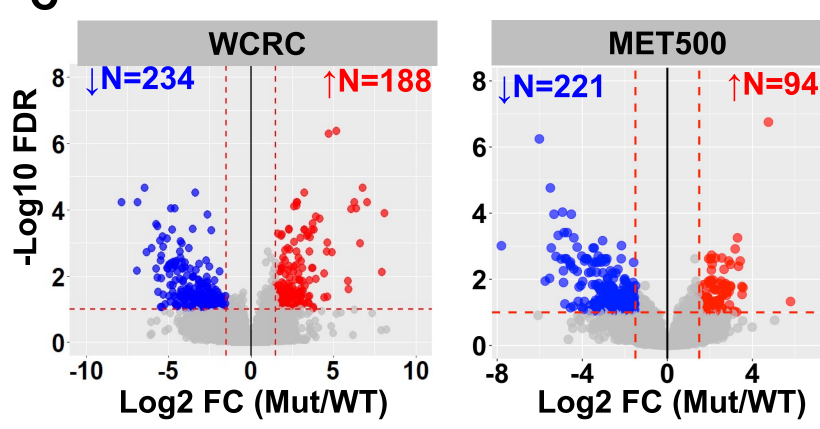
1427 statistical analysis. (** p<0.01)

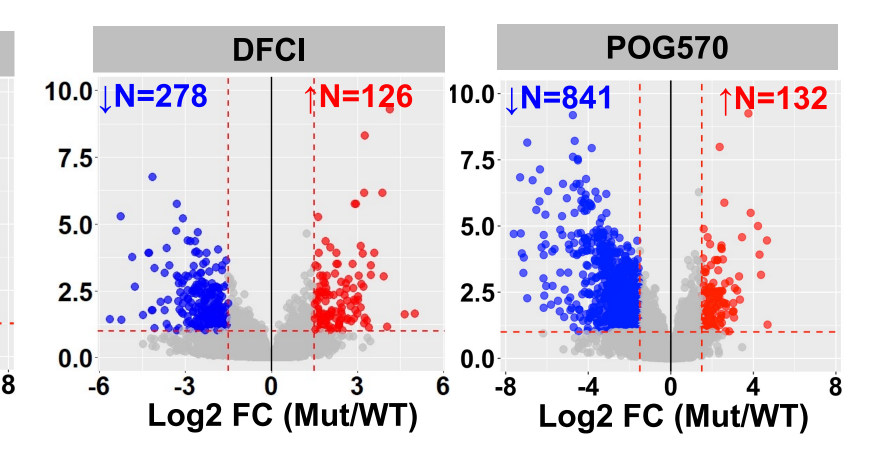
- 1428 F. Dot plots representing the enrichment distribution of the 50 MSigDB curated Hallmark
- 1429 gene sets in T47D-Y537S and T47D-D538G models normalized to WT cells.
- 1430 Significantly enriched gene sets (FDR<0.25) are highlighted in red, with names labeled
- 1431 in the venn diagram plot on the right panel. Gene sets enriched in Y537S and D538G
- 1432 cell models are in green and blue circles respectively.
- 1433 G. Immunoblot detection of β-catenin, phospho-GSK3β (Ser9), phospho-GSK3α
- 1434 (Ser21) total GSK3β and total GSK3α levels in T47D WT and mutant cells after
- 1435 hormone deprivation. Tubulin was blotted as a loading control. Representative blots
- 1436 from three independent repeats is displayed for each protein.
- 1437 H. Quantification of IncuCyte wound scratch assay with or without 5µM LGK974
- 1438 treatment for 72 hours. The migratory region normalized to T0 are labelled in blue.
- 1439 Images were taken under 10x magnification. Cell migration rates were quantified based
- 1440 on relative wound densities with eight biological replicates. Representative experiment
- 1441 from three independent repeats is shown. Pairwise two-way ANOVA between WT and
- 1442 each mutant was performed. (** p<0.01)
- 1443 I. IncuCyte migration assay with combination treatment of four different doses of
- 1444 LGK974 and Fulvestrant in T47D-D538G cells. Inhibition rates were calculated using
- 1445 the wound density at 48 hours normalized to vehicle control with values labelled using
- 1446 color scales in the heatmap. Positive Bliss scores are considered a synergistic
- 1447 combination. Representative experiment from three independent repeats is shown.
- 1448 J. Dot plot representing the fold changes of all Wnt signaling component genes in both
- 1449 T47D ESR1 mutant cell models normalized to WT cells. The blue dotted frame
- highlights the unique T47D-D538G enriched genes as well as genes that are enriched
- in both mutants, but with a larger magnitude of enrichment in the T47D-D538G cells.
- 1452 K & L. Immunoblot validation of Fulvestrant-induced ER degradation (K) and FOXA1
- 1453 knockdown (L). Cell lysates were subjected to ER and FOXA1 detection. Tubulin was
- 1454 blotted as a loading control. These validation experiments were performed once.

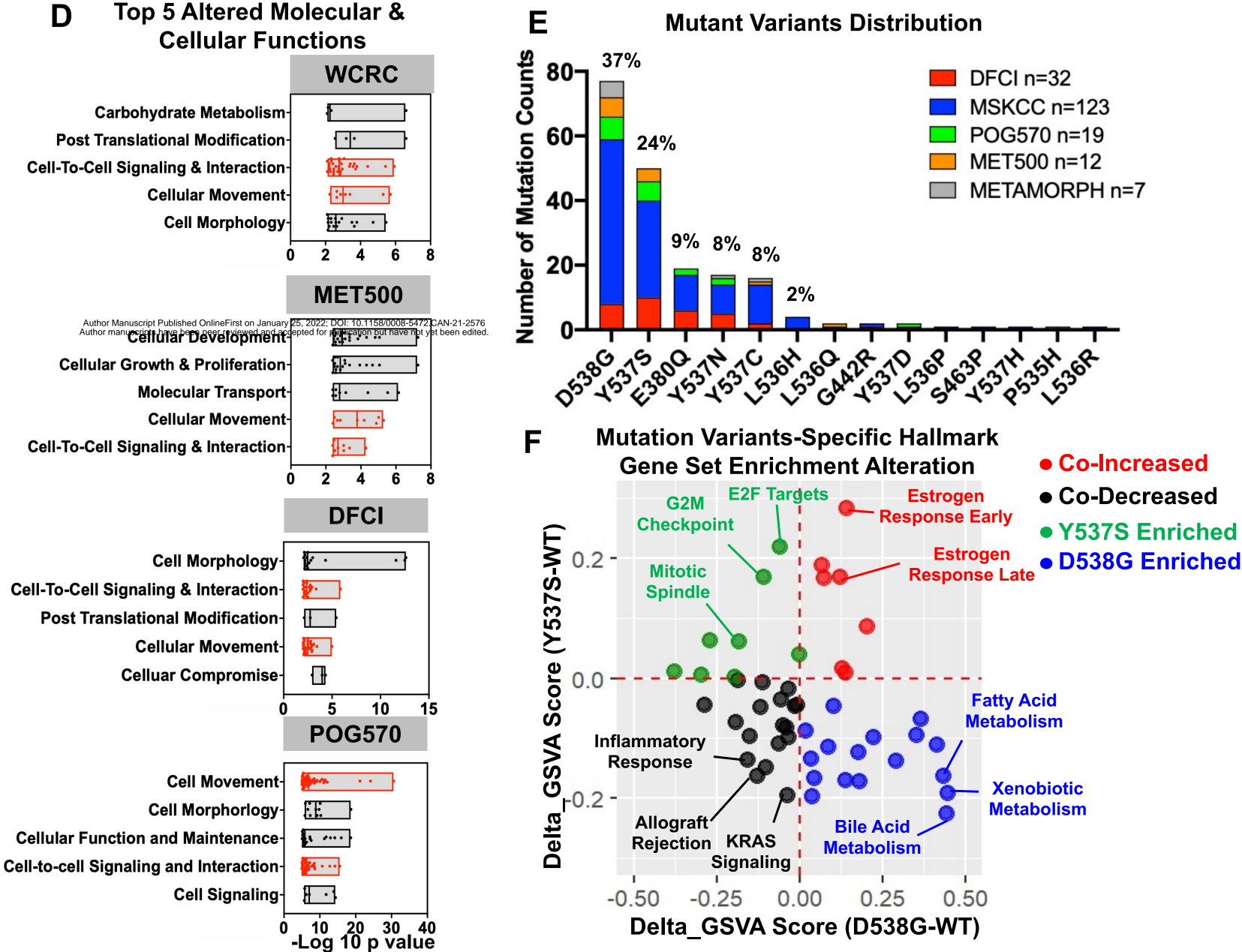
- 1455 M & N. Wound scratch assay in T47D-D538G and WT cells with 1µM of Fulvestrant
- 1456 treatment (M) or knockdown of FOXA1 (N) for 72 hours. Cell migration rates were
- 1457 quantified based on wound closure density. For fulvestrant treatment, data were merged
- 1458 from 3 (WT) or 6 (D538G) independent experiments. For FOXA1 knockdown,
- 1459 representative result from three independent repeats is displayed. Pairwise two-way
- 1460 ANOVA between siScramble/siFOXA1 or vehicle/Fulvestrant conditions in each cell
- 1461 type was performed. (* p<0.05, ** p<0.01)
- O. PCA plot showing the FOXA1 peak distribution of T47D WT, WT+E2, T47D-Y537Sand T47D-D538G groups.
- 1464 P. Heatmaps representing the comparison of FOXA1 binding intensities in T47D-D538G
- 1465 mutants to FOXA1 binding in WT cells. Displayed in a horizontal window of ± 2kb from
- 1466 the peak center. The pairwise comparison between WT and mutant samples was
- 1467 performed to calculate the fold change (FC) of intensities. Binding sites were sub-
- 1468 classified into sites with increased intensity (FC>2), decreased intensity (FC<-2), and
- 1469 non-changed intensity (-2<FC<2). Percentages of each subgroup are labelled on the
- heatmaps.
- 1471 Q. Bar charts showing the percentage of ATAC peaks overlapping (black) or not
- 1472 overlapping (grey) with FOXA1 binding sites in T47D-WT, T47D-Y537S and T47D-
- 1473 D538G cells.
- 1474 R. Venn diagram showing the intersection of genes annotated from dually gained ATAC
- 1475 and FOXA1 peaks (±3kb of TSS with 200kb of the peak flank) and RNA-seq
- 1476 differentially expressed non-canonical ligand-independent genes (gene with |fold
- 1477 change|>2, FDR<0.005 in D538G vs WT excluding genes with |fold change|>1.5,
- 1478 FDR<0.01 in WT+E2 vs WT groups). *TCF4* is highlighted.
- 1479 S. Wound scratch assay in T47D-WT and T47D-D538G cells with or without prior
- 1480 transfection of a dominant negative *TCF4* plasmid for 72 hours. Pairwise two-way
- 1481 ANOVA between vehicle and treatment conditions was performed. Data from one
- 1482 representative experiment of three independent experiments (each with six biological
- 1483 repeats) is shown. (** p<0.01)
- 1484
- 1485



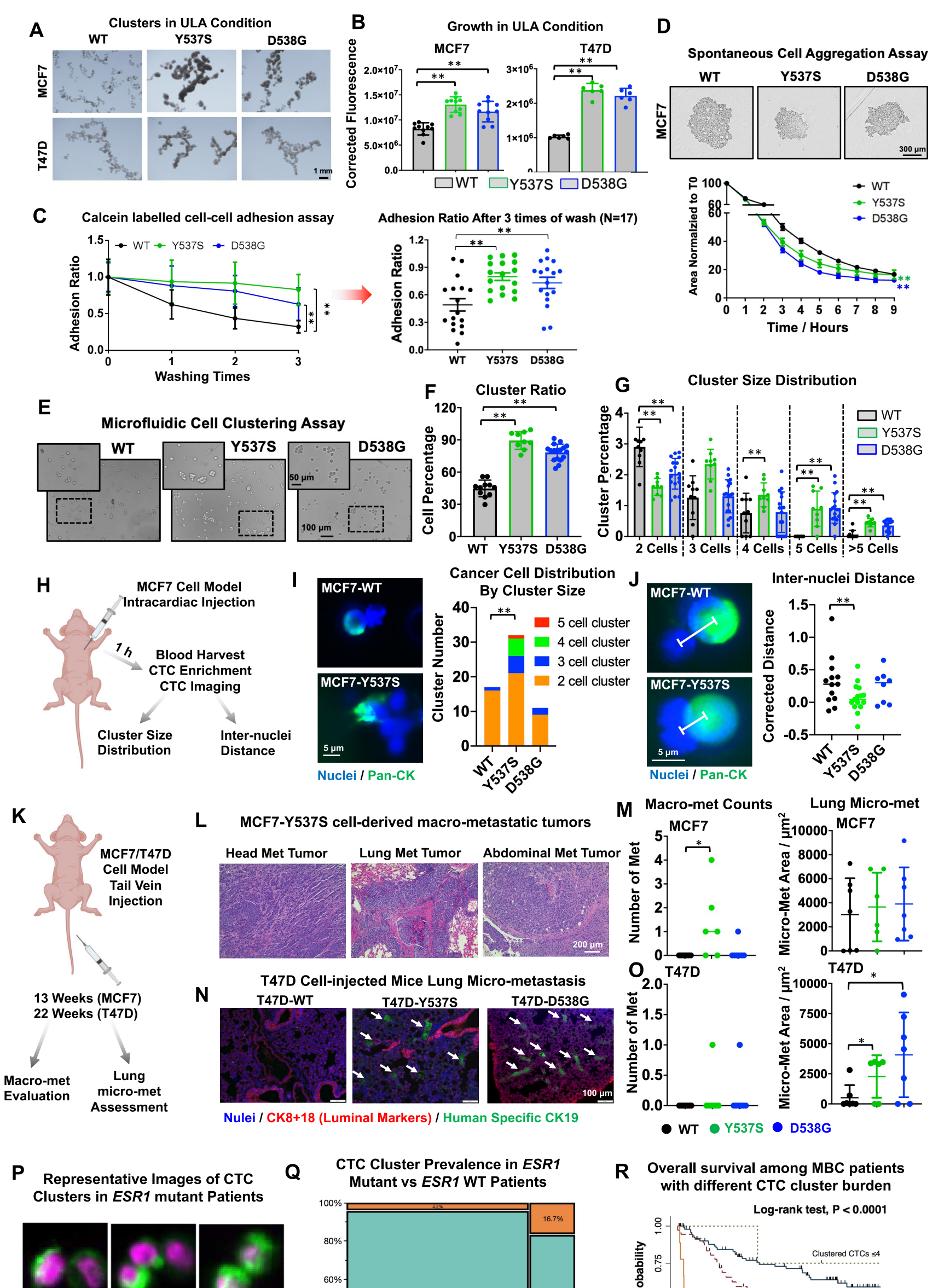
Differentially Expressed Genes С





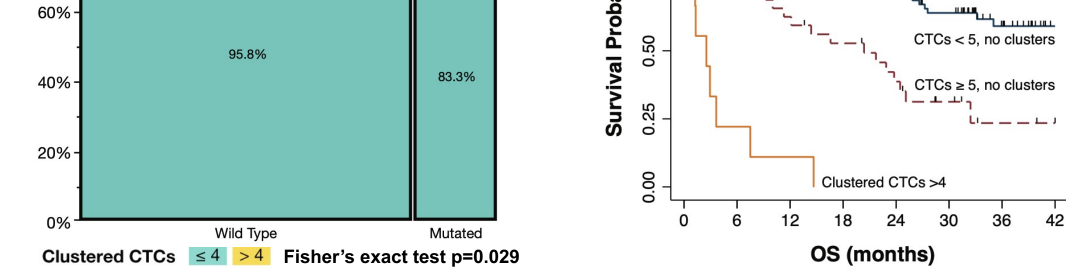


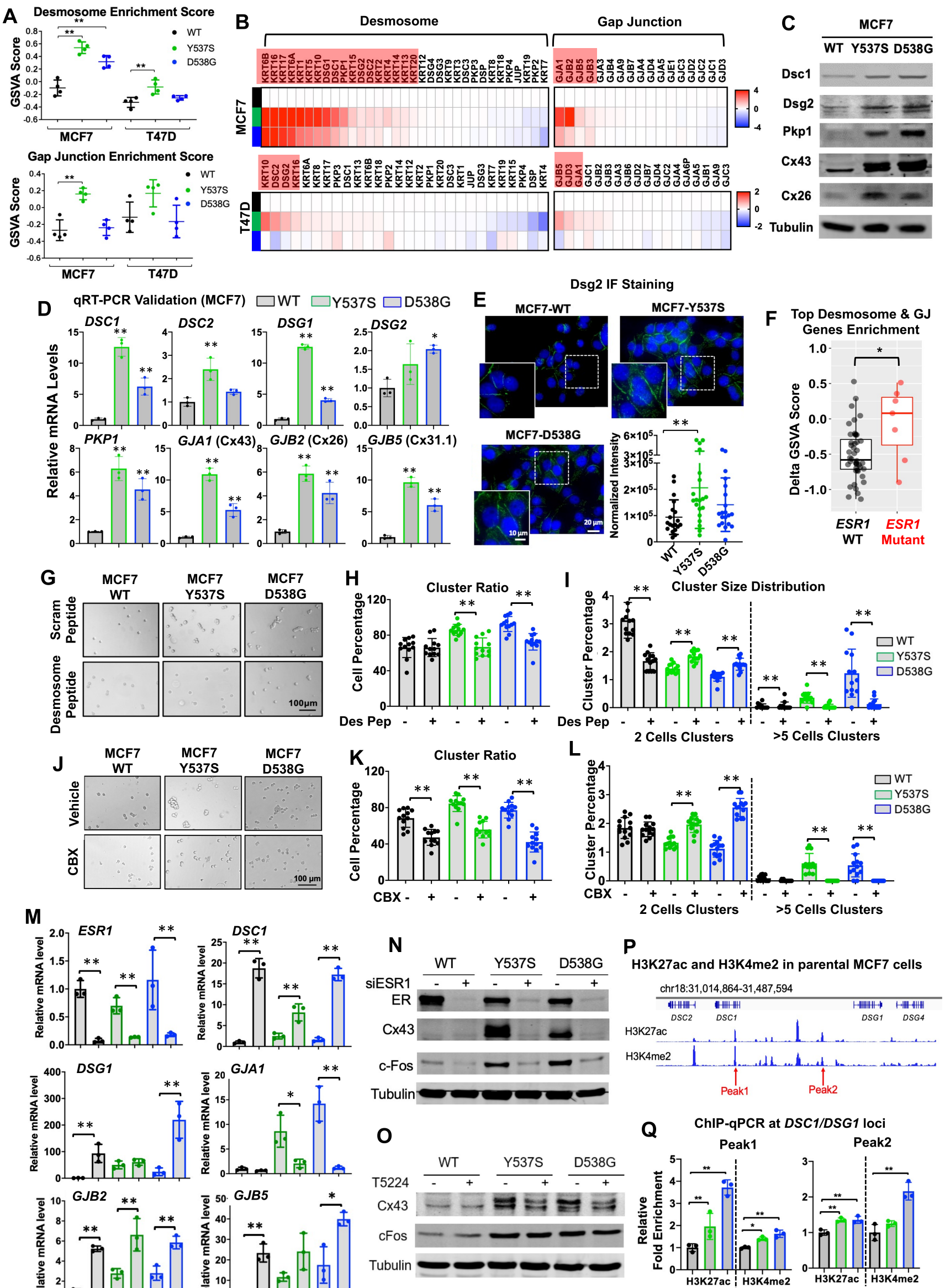
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Green: Ck8/18/19; Pink: Nuclei

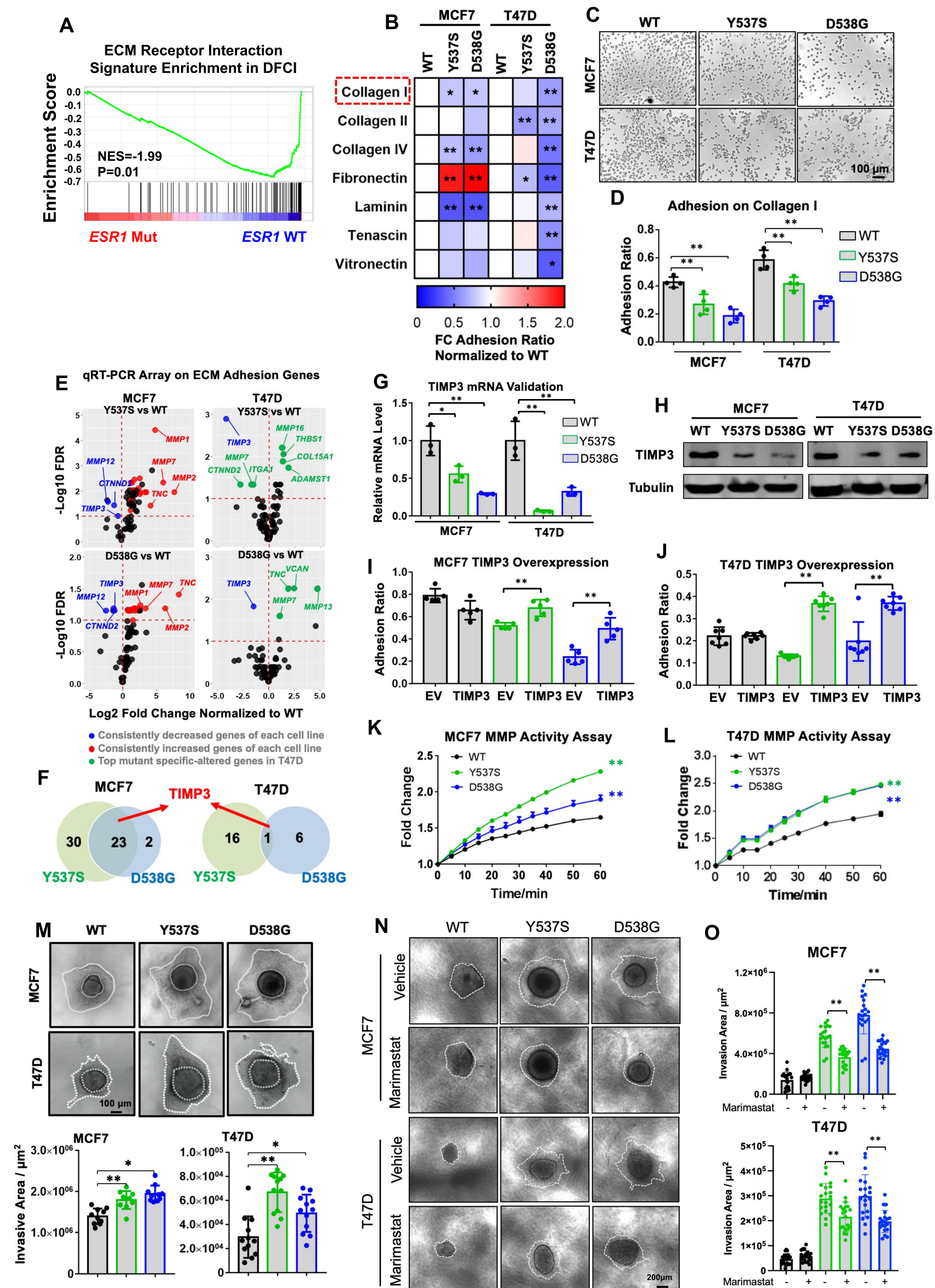






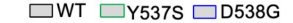


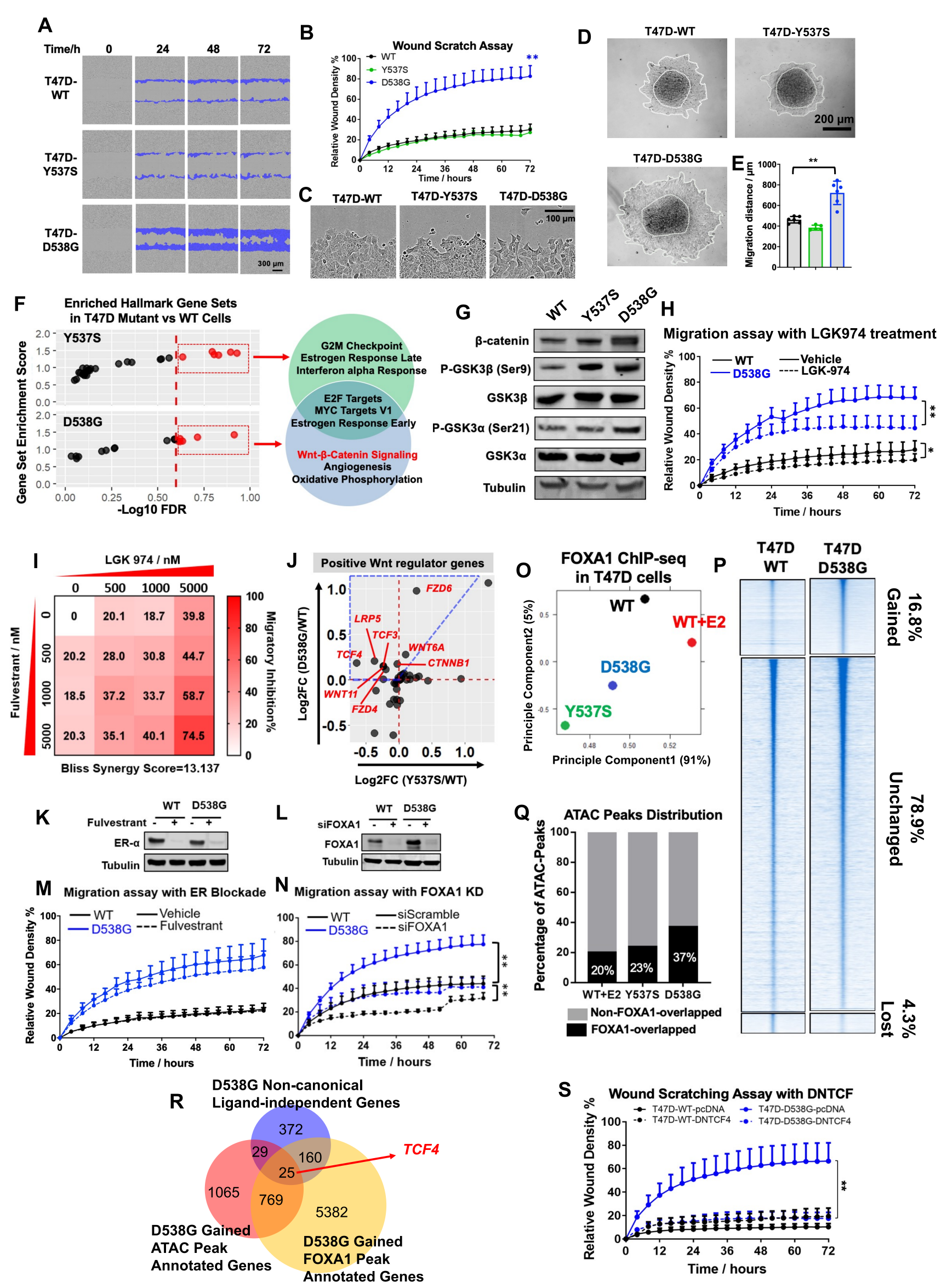
















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Hotspot ESR1 mutations are multimodal and contextual modulators of breast cancer metastasis

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