1	H3K27me3 is a determinant of chemotolerance
2	in triple-negative breast cancer
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23 Summary

24

25 Triple-negative breast cancer is associated with the worst prognosis and the highest risk of 26 recurrence among all breast cancer subtypes¹. Residual disease, formed by cancer cells persistent to chemotherapy, remains one of the major clinical challenges towards full cure^{2,3}. There is now 27 28 consensus that non-genetic processes contribute to chemoresistance in various tumor types, 29 notably through the initial emergence of a reversible chemotolerant state^{4–6}. Understanding non-30 genetic tumor evolution stands now as a prerequisite for the design of relevant combinatorial 31 approaches to delay recurrence. Here we show that the repressive histone mark H3K27me3 is a 32 determinant of cell fate under chemotherapy exposure, monitoring epigenomes, transcriptomes 33 and lineage with single-cell resolution. We identify a reservoir of persister basal cells with EMT 34 markers and activated TGF- β pathway leading to multiple chemoresistance phenotypes. We 35 demonstrate that, in unchallenged cells, H3K27 methylation is a lock to the expression program of 36 persister cells. Promoters are primed with both H3K4me3 and H3K27me3, and removing H3K27me3 37 is sufficient for their transcriptional activation. Leveraging lineage barcoding, we show that 38 depleting H3K27me3 alters tumor cell fate under chemotherapy insult - a wider variety of tumor 39 cells tolerate chemotherapy. Our results highlight how chromatin landscapes shape the potential of 40 unchallenged cancer cells to respond to therapeutic stress.

41 **Text**

42 Emergence of resistance phenotypes from initially responding or partially-responding tumors has been 43 modeled as a multi-step process in cancer⁷. Initially, post drug insult, only a pool of persister cells - also 44 called drug-tolerant persister cells (DTPs) - manage to tolerate the cancer treatment and survive⁸. 45 These cells constitute a reservoir of cells from which drug-resistant cells, actively growing under cancer 46 treatment, will ultimately emerge⁸⁻¹⁰. In triple-negative breast cancer (TNBC), both genetic and 47 transcriptomic mechanisms have been proposed to drive cancer evolution towards chemoresistance, through combined selective and adaptive modes of evolution¹¹. The recent identification of a multi-48 49 clonal reversible drug-tolerant state post neo-adjuvant chemotherapy in patient-derived xenografts 50 (PDX)⁶ suggested that the earliest steps of chemoresistance in TNBC are not driven by genetic 51 alterations, but rather by non-genetic plasticity in multiple cancer cells. Similarly, in other cancer types, 52 persister states have been identified solely by changes in transcriptomic and epigenomic features in 53 response to targeted therapies or chemotherapies^{12–15}.

54 Genetic history of many cancer types has been extensively modelled thanks to both bulk and single-55 cell approaches^{11,16}. In contrast, little is known about the epigenomic heterogeneity and dynamics of 56 acquisition of epigenetic alterations. While recent studies have focused on the evolution of DNA 57 methylation^{17,18}- among the most stable epigenetic locks to gene expression, contribution to tumor 58 evolution of more versatile epigenetic modifications, has remained poorly understood. Single-cell 59 methods to map repressive and permissive histone modifications, key players of cellular plasticity, have emerged only recently^{19,20}, enabling the study of epigenomic diversity within biological systems. 60 61 Here, combining single-cell transcriptomics and epigenomics with lineage barcoding, we show that the 62 distribution of H3K27me3 – trimethylation at lysine 27 of histone H3 - is a key determinant of cell fate 63 upon chemotherapy exposure in TNBC, shaping expression programs and cell potential to tolerate chemotherapy. 64

Resistance to adjuvant chemotherapy for TNBC patients, post-surgery, cannot be easily studied as
biopsies are not routinely performed when the disease progresses. To circumvent these limitations,

67 we modelled, in vivo and in vitro, phenotypes of drug-response observed in patients. In vivo, we 68 treated a patient-derived xenograft (PDX) model, established from a patient with residual TNBC^{19,21}, 69 with Capecitabine, the standard of care for residual breast tumors. After the first round of 70 chemotherapy treatment, mice displayed a pathological complete response (pCR), but tumors 71 eventually recurred ('recurrent') and mice were treated again with chemotherapy, to which tumors 72 responded to various extents ('responder' or 'resistant') (Fig. 1a and Extended Fig. 1a-c). These 73 recurrent tumors potentially arose from persister cells, surviving initial chemotherapy treatment⁹. We 74 isolated 'persister' cells by pooling the fat pad from 6 mice with pCR (Extended Fig. 1d-f). We also 75 generated 'residual' tumors (n=2) to phenocopy a clinical situation of partial response (Fig. 1a and 76 Extended Fig. 1a and 1d) by administering a moderate dose of Capecitabine (270 mg/kg, half the usual 77 dose).

In vitro, we treated an initially chemosensitive TNBC cell line (MDA-MB-468), with the pro-drug of Capecitabine, Fluorouracil (5-FU)²², as Capecitabine is not metabolized *in vitro*. We drove independently three pools of cells to chemoresistance with prolonged 5-FU treatment (>15 weeks). After 3 weeks, only few cells survived drug insult (0.01% of the initial population) and started proliferating again under chemotherapy after 10-15 days (Fig. 1b and Extended Fig. 2a-b). Over 15 weeks, populations of resistant cells emerged, with an IC50 to 5-FU over 4-fold higher than chemonaive population and doubling times comparable to chemonaive cells (Extended Fig. 2c).

85 To characterize transcriptomic evolution of chemonaive cells towards chemotolerance and 86 subsequently chemoresistance, we performed single-cell RNA-seq (scRNA-seq) in both cell lines and 87 PDX models (Fig. 1c-d, Extended Fig. 1g-i, 2d-f). In vivo, scRNA-seq was mandatory to identify the rare 88 human persister cells among the vast majority of stromal mouse cells. Out of the fat pad, we isolated 89 n~3,480 persister cells, an average of 580 cells per mouse. Both in vivo and in vitro models, diverse cell 90 populations, with distinct expression programs (Fig.1c-d), originated from the pool of persister cells, 91 which recurrently activated the same set of pathways compared to chemonaive cells (Fig. 1c-e, 92 Extended Fig. 1g-i, Extended Fig. 2d-g). Originating from KRT5-expressing cancer cells, persister cells

93 recurrently activated sets of genes further establishing basal cell identity (Extended Fig. 1h-j), such as 94 *KRT14* (Fig. 1c-d). In addition, persister cells showed an activation of the TGF- β pathway with the 95 expression of multiple players including ligands and receptors: in vivo - TGFB2, INHBA, INHBB, TGFB2R 96 and TGFB3R (Fig.1c, Extended Fig.1h-i) - and in vitro - TGFB1, INHBA and TGFBR1 (Figure 1d, Extended 97 Fig. 2f). Compared to chemonaive cells, persister cells also showed an activation of genes associated 98 to the Epithelial-to-Mesenchymal Transition (EMT, Fig.1c-d, Extended Fig. 1h-i, 2e-f), such as FOXQ1, 99 a transcription factor previously identified to drive EMT^{23,24} in cancer, and *NNMT*, characteristic of the 100 metabolic changes that accompany EMT^{25–27}. TGF- β and EMT associated-genes have been shown as 101 markers of residual TNBC^{28,29} and were proposed as potential drivers of chemoresistance in lung³⁰, pancreatic³¹, breast³², and colorectal cancers^{33,34}. *In vivo*, we showed that persister and residual tumor 102 103 cells actually clustered together (cluster C1) and shared a common expression program, suggesting 104 similar mechanisms of chemotolerance independent of the residual burden (Extended Fig. 1g). Yet 105 persister cells displayed a decreased proliferation rate as attested by a higher number of cells in G0/G1 106 (Extended Fig. 1f and Extended Fig. 2b), in line with previous reports^{12,14,35}. *In vitro*, we identified two 107 clusters of persister cells (clusters C2 and C4), that differ by their expression of additional EMT markers 108 such as CDH2 (Fig. 1d) and TWIST1. Early individual persisters (day 33) solely belonged to cluster 109 C2/CDH2- whereas growing persisters could either belong to C2/CDH2- or C4/CDH2+ (Fig.1d and 110 Extended Fig. 2d). Overall, we identified both in vivo and in vitro a reservoir of persister basal cells with 111 EMT markers and activated TGF- β pathway evolving to multiple resistant phenotypes (Fig. 1c-d). We 112 pinpoint TGF- β and EMT pathway activation as the earliest common molecular events at the onset of 113 chemoresistance in TNBC, defining a common Achilles' heel, to target chemoresistant cells before 114 they phenotypically diversify.

To follow clonal evolution under therapeutic stress, we had initially introduced unique genetic barcodes in chemonaive MDA-MB468 cells prior to our experiments (Extended Fig. 3a). We leveraged our previous barcoding method³⁶ to allow robust detection of barcodes in scRNA-seq data, as shown by the high fraction of cells with a lineage barcode (Extended Fig. 3b). In addition, we verified that 119 barcodes frequencies detected in scRNA-seq data correlated with those detected in bulk, confirming 120 the sensitivity of barcode detection in scRNA-seq data (Extended Fig. 3c). Then comparing barcode 121 diversity at the persister stage, we observed that 5-FU and DMSO-treated cells display equivalent 122 lineage diversity (Extended Fig. 3d-e), showing that the persister state is multi-clonal. To test if the 123 lineages that persist are a random draw of the chemonaive population, we compared barcode 124 frequencies between the starting population and the 5FU or DMSO-treated cells. As cells after barcode 125 tagging and before chemotherapy treatment are growing, resulting in several cells with the same 126 barcodes, if surviving cells had no particular predisposition then they should resemble a random draw 127 of the chemonaive barcoded cells (day 0). This is what we indeed observed for DMSO-treated cells 128 when compared to random drawing (Extended Fig. 3f-g). However, barcode frequencies of the 5-FU 129 treated cell deviate from the random scenario (Extended Fig. 3f-g), indicating that some lineages 130 present in the chemonaive population have a predisposition to tolerate the treatment. This was 131 further confirmed by the comparison of barcode frequencies across persister states from independent 132 experiments showing that independent persister populations shared lineage barcodes (Extended Fig. 133 3h). To monitor clonal evolution within the different subgroups of persister cells, we compared the 134 barcode diversity within expression clusters (Fig. 1f). We found that lineage diversity decreases over 135 the course of treatment, eventually leading to few clones dominating the chemoresistant cluster (Fig. 136 1f). The diversity in the CDH2+ persister cluster was lower than in the CDH2- persister cluster (Fig. 1f), 137 suggesting that only rare persister cells switched to the CDH2+ persister state. By combining detection 138 of lineage barcode and expression programs at single-cell resolution, we demonstrated that a small 139 pre-disposed subset of cells tolerate chemotherapy, progressively transiting from a CDH2- to a CDH2+ 140 persister state, and eventually leading fewer cells to resist.

To hamper this chemo-driven clonal evolution of cancer cells, we next investigated the molecular basis of such rapid phenotypic evolution. Using whole-exome sequencing (Extended Fig. 4a), we first analyzed mutations, copy-number alterations (CNA) and related mutational signatures acquired by persister and resistant cell populations since the onset of 5-FU treatment ('chemonaive D0' 145 population). We could not identify any recurrent mutations across experiments (Extended Fig. 4b), or 146 any CNA (amplifications or homozygous deletions) or mutations affecting known driver genes of breast 147 cancer in any population¹⁶. Only a minor fraction of mutations found in persister cells were attributed 148 to 5-FU (mutational signature 17³⁷) in comparison to resistant cell populations where over 50% of acquired mutations are associated to 5-FU exposure (p<10⁻¹⁰, Extended Fig.4c), indicating that chemo-149 150 related mutations are acquired over a timeframe that is not compatible with the rapid phenotypic 151 evolution seen in persister cells. Finally, computing cancer cell fractions for each mutation, we 152 confirmed that persister populations are extensively multi-clonal compared to chemonaive cells 153 (Extended Fig. 4d), in line with the lineage barcoding results.

154 We next investigated changes in epigenomes during chemotherapy treatment. Using single-cell 155 profiling (scChIP-seq) of the repressive H3K27me3 epigenomic mark, we observed that H3K27me3 156 epigenomes faithfully captured the evolution of cell states with chemotherapy (Fig. 2a, Extended Fig. 157 5a-c). Persister cells shared a common H3K27me3 epigenome (Fig. 2a-c, cluster C1), in contrast to 158 resistant cells split in clusters C1 and C3. In comparison to chemonaive cells, cells from cluster C1 159 showed recurrent redistribution of H3K27me3 methylation, the highest changes (|log2FC|>2) 160 occurring specifically at transcription start sites (TSS) and gene bodies (Fig. 2d) and corresponding to a 161 loss of H3K27me3 enrichment (75 regions with log2FC<-2, and 2 regions with log2FC>2). This depletion 162 correlated with the highest changes in gene expression observed by scRNA-seq (Fig. 2e-f and Extended 163 Fig. 5d) and was associated to the transcriptional de-repression of 14% of persister genes (Fig. 2g) -164 genes overexpressed in persister versus chemonaive cells. In the chemonaive cells, two epigenomic 165 subclones were recurrently identified indicated an epigenomic heterogeneity in this population (n=3 166 experiments, C2 & C4, Fig. 2b and Extended Fig. 5e-f). In contrast to cells from cluster C4 (median 167 correlation score r=-0.34, no cells over r=0.20), a fraction of cells within cluster C2 shared epigenomic 168 similarities with cells from C1 (49/381 cells over r=0.20, Fig. 2c and Extended Fig. 5g-h), but remained 169 discernible from the pool of persister cells (no cells from C2 over median r of C1, see Methods). This 170 suggests that cells from C2 could fuel the persister population when exposed to chemotherapy, with

the need of chemo-induced chromatin changes to achieve tolerance. In addition to these two epigenomic subclones within the chemonaive cells, we also detected rare cells with a persister epigenomic signature, in only one of our three experiments (60/976 chemonaive-D60-#1 cells in cluster C1 – Extended Fig. 5f), suggesting that spontaneous transition to H3K27me3 persister state rarely occurs in the absence of chemotherapy.

176 To test whether H3K27me3 enrichment was the lock to the persister expression program in 177 chemonaive cells, we treated cancer cells with the EZH2 inhibitor (EZH2i) UNC1999³⁸, to deplete 178 H3K27me3 from cells (Extended Fig. 5i), in the absence of chemotherapy. EZH2i treatment 179 phenocopied persister state to chemotherapy as expression fold-change induced by EZH2i were 180 specifically correlated to those induced by chemotherapy exposure at early time points (Fig. 2h, 181 Extended Fig. 5j, r=0.71 versus r=0.31 with changes in resistant cells). Furthermore, we observed that 182 EZH2i was sufficient to lead to the activation of 62% of H3K27me3-enriched persister genes (15/24 183 genes), suggesting that H3K27me3 was the sole lock to their activation (Fig. 2i and Extended Fig. 5k-l). 184 EZH2i was also sufficient to lead to the over-expression of 61% of *persister* genes independently of any 185 H3K27me3 enrichment in chemonaive cells (86/144 genes), such as KRT14, suggesting that these genes 186 might be targets of H3K27me3-regulated persister genes.

187 As we observed H3K27me3 changes precisely at TSS, we further explored the evolution of chromatin 188 modifications at TSS, focusing on H3K4me3 - trimethylation at lysine 4 of histone H3 - a permissive 189 histone mark shown to accumulate over TSS with active transcription. In contrast to single-cell 190 H3K27me3 epigenomes which were sufficient to separate cell states along treatment (Fig 2a), 191 individual H3K4me3 epigenomes of chemonaive and persister cells were indiscernible (Fig. 3a and 192 Extended Fig. 6a-b). Comparing H3K4me3 single-cell tracks of chemonaive and persister cells (Fig. 3b 193 and Extended Fig. 6c), we observed sparse H3K4me3 enrichment at the TSS of persister genes already 194 in chemonaive cells, compared to house-keeping genes (Extended Fig. 6d). In individual persister cells, 195 H3K4me3 enrichment was significantly more synchronous at these TSS (Extended Fig. 6e, p=9.2 10⁻³) 196 than in chemonaive cells. Altogether, H3K4me3 and H3K27me3 could either accumulate on the same

TSS but in different chemonaive cells, or H3K4me3 could accumulate together with H3K27me3 in asubset of cells.

199 To test whether H3K4me3 could co-exist with H3K27me3 in the same individual cells prior to 200 chemotherapy exposure, we performed successive immunoprecipitation of H3K27me3 with H3K4me3 201 or H3K27me3 with isotype control (IgG). We detected n= 1,547 transcription start sites significantly 202 enriched in DNA immunoprecipitated with both H3K27me3 and H3K4me3, compared to the control 203 (H3K27me3/IgG) precipitated fraction (peak-ratio>0.15, q-value<1.0 10⁻³, Extended Fig. 6g). We found 204 that bivalent chromatin in chemonaive cells (D0) was detected at genes associated to basal and EMT 205 pathways, as well as various developmental pathways (e.g Hedgehog pathway) (Fig. 3c-d and Extended 206 Fig. 6f), and at the ligand of the TGF- β pathway, TGF- β 1 (Fig. 3c). The majority of K27-regulated 207 persister genes (18 out of 24) were found in a bivalent chromatin configuration in the chemonaive cell 208 population (Fig. 3c, Extended Fig. 6f). In vivo, we observed enrichment of H3K27me3 and H3K4me3 209 modifications measured independently at n=1,370 TSS, particularly at EMT-associated genes and TGF-210 β2 (Fig. 3e-f and Extended Fig. 6h-i), corroborating that *persister* genes are in a bivalent configuration 211 in chemonaive cells.

212 To further validate that H3K27me3 distribution regulates the emergence of persister cells, we next 213 modulated H3K27me3 distribution simultaneously to chemotherapy. We used both the EZH2 inhibitor 214 (EZH2i - UNC1999³⁸), and a KDM6A/B inhibitor (KDM6A/Bi - GSK-J4³⁹) to prevent demethylation of 215 H3K27me3 residues. Co-treating cells with one of these modulators together with 5-FU, we observed 216 opposite modulation of the ability of cells to tolerate the chemotherapy. EZH2i increased the number 217 of persister cells, whereas KDM6A/Bi led to a decrease in the number of persisters at day 21 (Fig. 4a-218 b). At day 60, KDM6A/Bi further completely abolished the growth of persister cells under 5-FU, 219 whereas it has no effect on chemonaive cancer cells (Fig. 4c and Extended Fig. 7a). These results were 220 confirmed in a second TNBC cell line HCC38, albeit to a lesser extent for EZH2i (Extended Fig. 7b-c). We 221 then tested how EZH2i affected cell fate of persister cells. Comparing the number of unique barcodes 222 present in the two clusters under chemotherapy pressure with or without EZH2i (Extended Fig. 3a),

223 we observed that co-treating cells with EZH2i and 5-FU increases the number of lineage barcodes in 224 persister cells, and to a large extent in C4/CDH2+ cluster (Fig. 4d-f). These results showed that under 225 EZH2i cancer cells have an increased potential to reach persister state, and a mesenchymal CDH2+ 226 state. Overall, inhibiting the removal of methyl groups at H3K27 prevented cells from reaching 227 chemotolerance and resistance, demonstrating that some cancer cells need to actively demethylate 228 H3K27me3 residues to reach persister state and to proliferate under chemotherapy exposure. 229 Conversely, depleting H3K27me3 from chemonaive cells not only launched a persister-like expression 230 program, but it also enhanced chemotolerance. These results were consistent with a mechanism 231 where persister genes would be repressed by H3K27me3 in chemonaive cells, and primed with 232 stochastic H3K4me3 in a subset of cells – loss of H3K27me3 being the key to activation.

233 In conclusion, our study shows that the transition to persister state in TNBC is dependent on the 234 control of H3K27me3 distribution. We propose that combining chemotherapy with histone 235 demethylase inhibitors at the onset of chemotherapy exposure will decrease the pool of persister cells, 236 and thereby decrease recurrence. We demonstrate a role for both H3K27me3 writer and erasers in 237 regulating the phenotypic switch from chemonaive to chemopersister state, highlighting the 238 instrumental role of repressive histone landscapes as determinants of cell fate. Several studies had 239 started to interrogate which epigenetic modifiers could regulate expression programs of persister or 240 resistant cells, but studying them in isolation^{5,12,14,40,41}. We also show that the *persister* expression 241 program is primed in chemonaive cells with a stochastic presence of H3K4me3 but is repressed by 242 H3K27me3. In other words, genes are ready to be activated, H3K27me3 remaining the only lock to 243 activation. Persisters are cells without H3K27me3 or the one releasing the H3K27me3 lock, or a mixture 244 of both phenomena. We observe epigenomic priming of signaling pathways known to participate to 245 drug resistance in TNBC⁴², including Hedgehog, WNT (Extended Fig. 2f), TGF- β and ATP-binding 246 cassette drug transporters pathways (Extended Fig. 2f). Such epigenomic priming is reminiscent of developmental bivalency priming mechanisms⁴³ found in stem cells prior to differentiation and 247 248 appears key for the rapid activation of the genes upon therapeutic stress. Remains to be understood,

- 249 how only a fraction of bivalent genes are targeted by gene activation upon chemotherapy exposure,
- 250 whether tolerance to different drugs triggers the activation of the same set of bivalent genes, and

251 whether such mechanisms could be shared across cancer types. Determining the precise addressing

252 mechanisms that target H3K27me3 and H3K4me3 writers and readers to TSS of bivalent genes in

253 cancer cells, will be instrumental in the future to identify dedicated co-factors which could serve as

- alternative therapeutic targets to restrict the epigenetic plasticity of cancer cells.
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346 Legends

347 Figure 1: Identification of a pool of basal persister cells in TNBC in vivo and in vitro. a. (Up) Graph of 348 the relative tumor volumes (RTV, mm3) over time (days). Colored growth curves correspond to tumors 349 which have been further studied by scRNA-seq. Black arrows indicate the start of the second round of 350 Capecitabine treatment for the corresponding mice. (Down) Phenotypes and cell numbers are 351 indicated, with the number of mice used to collect samples in brackets. **b.** (Up) Graph representation 352 of the cell proliferation of triple negative breast cancer cell line MDA-MB-468 (MM468) treated with 353 5-FU (green for persister cells, and orange lines for resistant cells) or with DMSO (chemonaive - grey 354 lines). Each dot corresponds to an independent experiment. (Down) Schematic view of the 355 experimental design. Experience replicate numbers and passages of the cells at D0 are indicated. c. 356 UMAP representation of PDX scRNA-seq datasets, colored according to sample of origin (first panel) 357 or gene expression signal for differentially expressed genes between persister cells and chemonaive 358 tumor cells (remaining panels), log2FC and adjusted p-values are indicated above the graph. d. UMAP 359 representation of MDA-MB-468 cells scRNA-seq datasets, colored according to the sample ID (first 360 panel) or gene expression signal for differentially expressed genes between persister cells and 361 chemonaive cells (*KRT14* and *TGFB1* panels) or for a differentially expressed gene between the two 362 persisters clusters, *i.e.* clusters C4 vs C2 (CDH2 panel). Chemonaive population (in grey) corresponds 363 to DMSO-DO-#1. e. Barplot displaying the top 5 pathways activated in persister cells both in vivo and 364 in vitro from MSigDB c2_curated Breast, c2_KEGG, c7_Hallmark and c5_GO annotations. x-axis 365 corresponds to -log10 adjusted p-values for PDX. f. (Left) UMAP representation of scRNA-seq as in 1d, 366 cells are colored according to lineage barcode. (Right) Histogram of the lineage barcodes diversity 367 detected in the scRNA-seq data within Louvain partition clusters, and across samples. Colors 368 correspond to sample ID as in 1d.

369

370 Figure 2: H3K27me3 represses the persister expression program in chemonaive cells. a. UMAP 371 representation of scChIP-seq H3K27me3 datasets, cells are colored according to the sample of origin. 372 Chemonaive samples correspond to DMSO-treated cells, persister and resistant samples correspond 373 to 5-FU-treated cells, days of treatment are indicated. b. Same as in a. with cells colored according to 374 epigenomic clusters. c. Graph representation of the cell to cell inter-correlation between clusters C1, 375 C2 or C4 and the cluster C1. **d.** Genomic association between H3K27me3 peaks and gene annotation. 376 Full bars indicate adjusted p-value<1.0 10^{-2} . Empty bars indicate non-significant adjusted p-values. 377 "PC" indicates protein coding gene **e.** Repartition of H3K27me3 depleted peaks within re-expression 378 quantiles in persister cells. f. Cumulative scH3K27me3 profiles over TGFB1 and FOXQ1 in chemonaive 379 and persister cells (D33). Log2FC and adjusted p-value correspond to differential analysis comparison of cells from cluster C1 versus clusters C2 + C4. **g.** Pie chart displaying the fraction of persister genes potentially regulated by H3K27me3 in MM468 cells. **h.** Dot plot representing log2 expression foldchange induced by 5-FU or EZH2i at D33 versus D0. Correlation scores and associated p-value are indicated. **i.** Bulk H3K27me3 chromatin profiles for *TGFB1* and *FOXQ1* in cells treated with DMSO, 5-FU or EZH2i at D33.

385

386 Figure 3: Epigenomes of chemonaive cells are primed with co-accumulation of H3K27me3 and 387 H3K4me3. a. UMAP representation of scChIP-seq H3K4me3 datasets, cells are colored according to 388 the sample of origin. **b.** Cumulative scH3K4me3 enrichment profiles over TGFB1 and FOXQ1 in 389 chemonaive cells (D0) and persister cells (D60). c. Bulk H3K27me3->H3K4me3 and H3K27me3->IgG 390 chromatin profiles of TGFB1 and FOXQ1 in the chemonaive population (D0). Enrichment tracks show 391 enrichment over IgG control with associated odd ratio and adjusted p-value. **d.** Barplot displaying the 392 top 5 pathways (as in Fig. 1e) enriched in genes detected with combined H3K27me3/H3K4me3 in the 393 chemonaive MDA-MB-468 cell population (D0). e. Density plot representing cumulative scH3K27me3 394 and scH3K4me3 log2 enrichment at TSS in two independent cell populations within chemonaive PDX. 395 f. Cumulative scH3K27me3 and scH3K4me3 profiles over TGFB2 and ELN in the chemonaive PDX 396 tumor.

397

398 Figure 4: EZH2 inhibition modulates cell fate upon chemotherapy exposure. a. Representative 399 pictures of MDA-MB-468 cells treated 21 days with 5-FU alone or in combination with an inhibitor of 400 EZH2 (UNC1999) or an inhibitor of KDM6A/B proteins (GSKJ4). b. Histogram representing the number 401 of cells after treatment with 5-FU alone, 5-FU and EZH2i or KDM6i over 21 days, relative to the number 402 of cells at D0. (n=3, Mean ± sd). c. Colony forming assay of MDA-MB-468 co-treated with DMSO or 5-403 FU and indicated concentrations of KDM6A/Bi for 60 days. The data corresponds to 1 of 3 biological 404 replicates. d. UMAP representation of scRNA-seq datasets, colored according to the sample ID. e. As 405 in 4d. cells colored according to lineage barcode. f. Histogram of the diversity of lineage barcodes 406 detected in scRNA-seg data in each louvain partition cluster obtained from the chemonaive 407 population, 5-FU or EZH2i persister cells (D33). Colors correspond to sample ID as in 4d.

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- 410

411 Methods

412 PDXs. Female Swiss nude mice were purchased from Charles River Laboratories and maintained under 413 specific-pathogen-free conditions. Mouse care and housing were in accordance with institutional 414 guidelines and the rules of the French Ethics Committee (project authorization no. 02163.02). In this 415 study, we used a xenograft model generated from a residual triple-negative breast cancer post-416 neoadjuvant chemotherapy (HBCx95) established previously at Curie Institute with informed consent 417 from the patient^{44,45}. Five mice were not treated and kept as controls and twenty-seven mice were 418 treated orally with Capecitabine (Xeloda; Roche Laboratories) at a dose of 540 mg/kg, 5 d/week for 6 419 to 14 weeks. Relative tumor volumes (mm3) were measured as described previously²¹. Latency was 420 defined as the number of days between the observation of a complete response (Tumor size < 10421 mm3) after the first round of Capecitabine treatment, and the detection of a recurrent tumor (Tumor 422 size > 10 mm3). Eight mice were sacrificed after the first round of chemotherapy to study residual (2 423 mice) and persister (6 mice) human tumor cells. Seven mice with recurrent tumors (tumor volume 424 between 200 and 600 mm3) were treated for a second round of Capecitabine. The GraphPad PRISM 9 425 was used for statistics in Extended Fig.1b. The results represent the mean ± sd and statistical analysis 426 was performed using two-tailed Mann-Whitney test.

427 Before downstream analysis (scChIP-seq, scRNA-seq), control and Capecitabine treated tumors were 428 digested for 2 h at 37 °C with a cocktail of Collagenase I (Roche, Ref: 11088793001) and Hyaluronidase 429 (Sigma-Aldrich, Ref: H3506). Cells were then individualized at 37°C using a cocktail of 0.25% Trypsin-430 Versen (Thermo Fisher Scientific, Ref: 15040-033), Dispase II (Sigma-Aldrich, Ref: D4693) and DNase I (Roche, Ref: 11284932001) as described previously⁴⁶. Then, eBioscience red blood cell lysis buffer 431 432 (Thermo Fisher Scientific, Ref: 00-4333-57) was added to the cell suspension to remove red blood cells. 433 To increase the viability of the final cell suspension, dead cells were removed using the Dead Cell 434 Removal Kit (Miltenyi Biotec, Ref:130-090-101).

435

436 Cell lines, culture conditions and drug treatments. MDA-MB-468 cells were cultured in DMEM (Gibco-437 BRL, Ref: 11966025), supplemented with 10% heat-inactivated fetal calf serum (Gibco-BRL, Ref: 10270-438 106). HCC38 cell lines were cultured in RPMI 1640 (Gibco-BRL, Ref: 11875085), supplemented with 439 10% heat-inactivated fetal calf serum. All cell lines were cultured in a humidified 5% CO2 atmosphere 440 at 37 °C, and were tested as mycoplasma negative. GSKJ4 (KDM6A/B inhibitor, Sigma, Ref: SML0701), 441 GSKJ5 (GSK-J4 inactive isomer, Abcam, Ref: ab144397) and UNC1999 (EZH2 inhibitor, Abcam, Ref: 442 ab146152) were used at indicated concentrations. Cells were treated with 5 μM of 5-FU (Sigma, Ref: 443 F6627) alone or in combination with KDM6A/Bi or EZH2i for indicated days.

445 Colony forming assay. TBNC cells were plated in 6 multi-well plates at a density of 200,000 cells per
446 well and treated with the indicated drugs for 60 days (MDA-MB-468) or 50 days (HCC38). Treated
447 plates were monitored for growth using a microscope. At the time of maximum foci formation, colony

448 formation was evaluated after a staining with 0.5% Crystal Violet (Sigma, Ref: C3886).

449

450 Cell proliferation, doubling time and IC50. MDA-MB-468 and HCC38 cells were stained with Trypan
451 Blue (Invitrogen, Ref: T10282) exclusion test, and counted using a Countess automated cell counter
452 (Invitrogen, Ref: C10228) at indicated time of treatment (Fig.4b and Extended Fig.7b).

453 Doubling time (Extended Fig.2c) was calculated using this formula:

454 "DoublingTime = duration*log(2)/(log(Final Concentration)-log(Initial Concentration))"
455 For chemonaive condition and resistant condition, cell numbers were evaluated on cell population

456 during 10 days (n=3). For persister condition, cells were counted manually under the microscope at 457 day 13 and day 30 of treatment. Doubling time of 5-FU growing persister cells was studied from single 458 cell to confluent colony by assaying cell number every 4 days during 27 days (n= 6 single cells) (Figure 459 1b).

MDA-MB-468 chemonaive and chemoresistant cells were plated in 96 multi-well plates at a density of
10,000 cells per well and treated with increased concentration of 5-FU (1μM to 0.5M) for 72h. Cell
cytotoxicity was assayed with XTT kit (Sigma, Ref: 11465015001) and IC50 was calculated as the
concentration of 5-FU that is required to obtain 50% of cell viability (Extended Fig. 2c).

The GraphPad PRISM 9 was used for statistics and the results represent the mean ± sd of three independent experiments. Statistical analysis was performed using the Bonferroni test for multiple comparisons between samples (Fig.4b and Extended Fig.2c-right) or one-tailed Mann-Whitney test for the comparison between two conditions (Extended Fig. 2c-left and Extended Fig.7b).

468

469 Western blotting. DMSO- and EZH2i-treated cells (D33-#8) were lysed at 95°C for 10 minutes in 470 Laemmli buffer (50 mM Tris-HCl [pH 6.8], 2% SDS, 5% glycerol, 2 mM DTT, 2.5 mM EDTA, 2.5 mM EGTA, 471 4 mM Sodium Orthovanadate, 20 mM Sodium Fluoride, protease inhibitors, phosphatase inhibitors) 472 and proteins concentrations were measured using a Pierce BCA protein Assay Kit (Thermo Fisher 473 Scientific, Ref: 23225/23227). 10 µg of proteins were then separated on a 4-15% Mini-PROTEAN TGX 474 Stain-Free Gel (Bio-Rad, Ref: 4568085) at 160V. After transfer, the membrane was exposed to UV light 475 (Bio-Rad, ChemiDoc MP) and the image was further used for total protein quantification. The 476 membrane was then blocked for 1 h at room temperature in PBS pH 7.4 containing 0.1% Tween-20 477 and 1% milk (Regilait). Incubation anti-H3K27me3 (Dilution: 1:2000, Cell Signaling, Ref: 9733) primary 478 antibodies diluted in PBS pH 7.4, 0.1% Tween-20 were performed at 4°C overnight. Following 2 h 479 incubation at room temperature with an anti-rabbit peroxidase-conjugated secondary antibody

480 (Dilution: 1:10000, Thermo Fisher Scientific, Ref: 31460) diluted in PBS pH 7.4, 0.1% Tween-20,
481 antibody-specific labeling bands were revealed (Bio-Rad, ChemiDoc MP) using a SuperSignal West Pico
482 PLUS Chemiluminescent Substrate (Thermo Fisher Scientific, Ref: 34579).

483

Lentivirus packaging and cell transduction. Lentivirus was produced by transfecting the barcode plasmids pRRL-CMV-GFP-BCv2AscI and p8.9-QV and pVSVG into HEK293T cells as previously described³⁶. MDA-MB-468 cells from ATCC were infected at passage 11 with lentivirus produced from the barcode library (pRRL-CMV-GFP-BCv2AscI) which includes 18206 different barcodes of 20bp of a random stretch, at a low multiplicity of infection (MOI 0.1) to minimize the number of cells marked by multiple barcodes. Three weeks after transduction, cells were sorted for GFP expression to select cells with barcode insertion, and used for drug treatment.

491

Single-cell RNA-seq. For each single cell suspension (DMSO-D0-#1, 5-FU-D33-#1, 5-FU-D214-#1, 5-FUD67-#2, 5-FU-D171-#2, 5-FU-D50-#3, 5-FU-D77-#3 and 5-FU-D202-#3), approximately 3,000 cells were
loaded on a Chromium Single Cell Controller Instrument (Chromium Single Cell 3'v3, 10X Genomics,
Ref: PN-1000075) according to the manufacturer's instructions. Samples and libraries were prepared
according to the manufacturer's instructions. Libraries were sequenced on a NovaSeq 6000 (Illumina)
in PE8-8-91 with a coverage of 50,000 reads/cell.

498

Bulk lineage barcode library preparation and sequencing. Lineage barcodes are recovered by isolating genomic DNA from cells of interest (NucleoSpin Tissue, Mini kit for DNA from cells and tissue, Macherey Nagel, Ref: 740952.50). From the isolated genomic DNA, barcodes are amplified with three nested PCR steps as decribed in³⁶. In short, after a first specific PCR for the common region of the lineage barcodes, the amplified material was prepared for sequencing by addition of the illumina sequencing adaptaters and indexing and purification. Sequencing was done in order to obtain 50 reads, on average, per barcoded cell.

506

507 **Bulk ChIP-seq.** ChIP experiments were performed as previously described¹⁹ on 3x10⁶ MDA-MB-468 508 cells (DMSO-D67-#2, DMSO-D77-#3, DMSO-D113-#4, 5-FU-D67-#2, 5-FU-D77-#3, 5-FU-D113-#4) using 509 an anti-H3K27me3 antibody (Cell Signaling Technology, Ref: 9733 - C36B11). Sequencing libraries were 510 prepared using the NEBNext Ultra II DNA Library Prep Kit (NEB, Ref: E7645S) according to the 511 manufacturer's instructions. Libraries were sequenced on a NovaSeq 6000 (Illumina) in SE50 mode. 512

513 Single-cell ChIP-seq. Cells (DMSO-D60-#1, DMSO-D77-#3, DMSO-D131-#5, 5-FU-D33-#1, 5-FU-D67-#2,
 514 5-FU-D171-#2, 5-FU-D147-#3, 5-FU-D131-#6) were labeled by 15 min incubation with 1 μM CFSE

(CellTrace CFSE, ThermoFisher Scientific, Ref: C34554). Cells were then resuspended in PBS
supplemented with 30% Percoll, 0.1% Pluronic F68, 25 mM Hepes pH 7.4 and 50 mM NaCl. Cell
encapsulation, bead encapsulation and 1:1 droplet fusion was performed as previously described¹⁹.
Immunoprecipitation, DNA amplification and library were performed as in¹⁹. Libraries were sequenced
on a NovaSeq 6000 (Illumina) in PE100, with 4 dark cycles on Read 2, with a coverage of 100,000
reads/cell.

521

522 Quantitative chromatin profiling with chromatin indexing. Chromatin isolation, indexing, 523 immunoprecipitation and library preparation was adapted from⁴⁷. Briefly, 50,000 MDA-MB-468 were 524 lysed and digested with MNase for 20min at 37°C in the following buffer: 46mM Tris-HCl pH 7.4, 525 0.154M NaCl, 0.1% Triton, 0.1% NaDoc, 4.65mM CaCl2, 0.47x Protease Inhibitor Cocktail (Roche, Ref: 526 11873580001) and 0.09u/uL MNase (Thermo Scientific, Ref: EN0181). Fragmented nucleosomes were 527 then ligated for at least 24h at 16°C to double-stranded barcoded adapters containing 8bp barcodes 528 to combine samples: Pac1-T7-Read2-8bpBarcode-linker-Pac1. Next, 5 indexed chromatin samples 529 (DMSO, 5-FU, UNC, 5-FU + UNC, GSK-J4) were pooled, each containing a different 8-bp barcode, to 530 perform anti-H3K27me3 ChIP (Cell Signaling, Ref: 9733 - C36B11) on 250,000 cells in total in each pool. ChIP and DNA amplification was carried out as for scChIP-seq¹⁹ and a sequencing library was produced 531 532 for both IP and input pools and sequenced on NovaSeq 6000 (Illumina) in PE100 mode.

533

534 Sequential ChIP-seq. Primary ChIP experiments were performed as described previously¹⁹ on 30x 10⁶ 535 untreated chemonaive MDA-MB-468 cells using the anti-H3K27me3 antibody. After washes, samples 536 were eluted twice at 37°C for 15 min under agitation in an elution buffer (50mM Tris-Hcl pH8, 5mM 537 EDTA, 20mM DTT, 1% SDS) as in⁴⁸. Samples were diluted 10 times to decrease SDS and DTT 538 concentration. 10% of the eluted chromatin was kept as primary ChIP. Secondary ChIP, re-ChIP, was 539 performed overnight on the rest of the primary immuno-precipitated chromatin using an anti-540 H3K4me3 antibody (Cell signaling, Ref: #9751) or using an anti-IgG antibody (Cell signaling, Ref: #3900) 541 as a control, to determine the background level of the re-ChIP experiment. After washes, samples were 542 eluted twice at 65°C for 15 min under agitation in 0.1M NaHCO3 and 1% SDS as in⁴⁸. After reverse 543 crosslinking and DNA clean-up, 3 to 15 ng of immunoprecipitated DNA were used to prepare the 544 sequencing libraries using the NEBNext Ultra II DNA Library Prep Kit (NEB, Ref: E7645S) according to 545 the manufacturer's instructions. Libraries were sequenced on a NovaSeq 6000 (Illumina) in SE100 546 mode.

547

548 Whole exome sequencing. Genomic DNA from samples (DMSO-D0, DMSO-D147-#3, DMSO-D171-#5,
549 DMSO-D131-#6, 5-FU-D67-#2, 5-FU-D153-#2, 5-FU-D50-#3, 5-FU-D147-#3, 5-FU-D171-#5 and 5-FU-

- 550 D131-#6) were extracted with NucleoSpin Tissue, Mini kit for DNA from cells and tissue (Macherey
- 551 Nagel, Ref: 740952.50) and sequenced on a NovaSeq 6000 (Illumina) with a 100X depth.
- 552

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- 566
- 567 Fundings.
- 568 This work was supported by the ATIP Avenir program, by Plan Cancer and by the SiRIC-Curie program
- 569 SiRIC Grants #INCa-DGOS-4654 and #INCa-DGOS-Inserm_12554 (to C.V.). NGS was performed by the
- 570 ICGex platform of the Institut Curie. The work was supported by an ATIP-Avenir grant from CNRS and
- 571 Bettencourt-Schueller Foundation (to L.P.), by the *Labex CelTisPhyBio* (ANR-11-LABX-0038 to L.P.) and
- by a starting ERC grant from the H2020 program (758170-Microbar to L.P.).
- 573

574 Author contributions.

- 575 JM, AD, CL, LB, SBT, AE and AT performed experiments. scChIP-seq experiments were conducted
- 576 together with SF and KG. PDX experiments were performed by EM, LS and AD. MB and SB performed
- 577 sequencing. PP and CV performed omics data analysis. Lineage barcoding data were analyzed by AML,
- 578 CV and LP. Whole exome sequencing data were analyzed together with EL. CV, LP and JM conceived
- 579 and designed experiments. CV, JM, PP and LP wrote the manuscript with input from all authors.
- 580

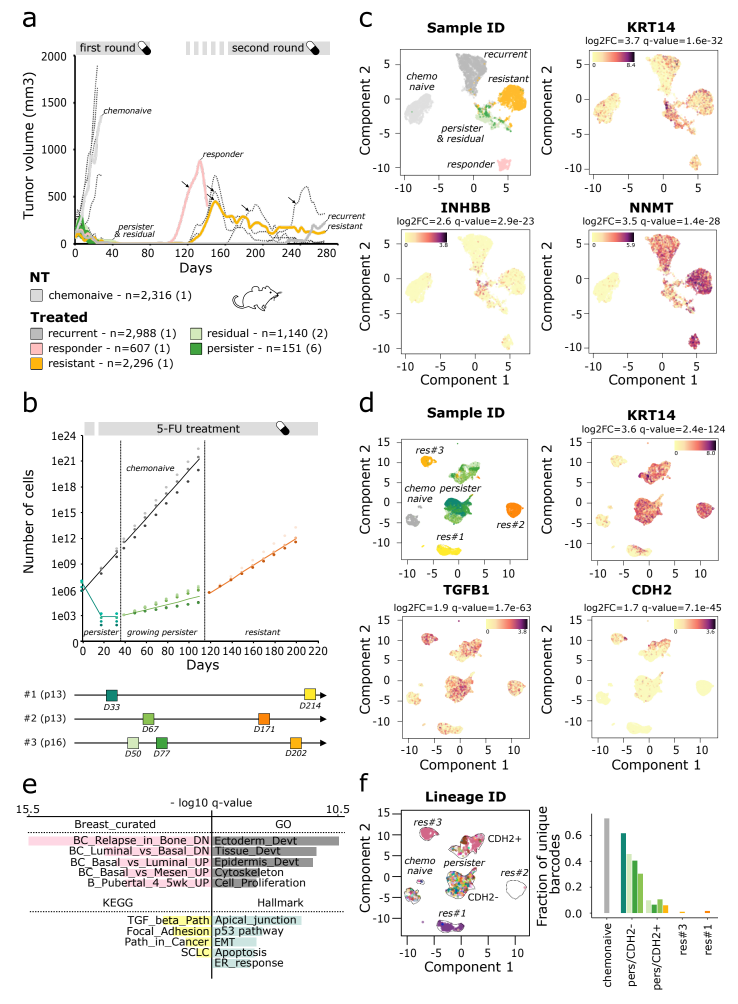
581 **Competing interest declaration**.

- 582 The authors declare no financial competing interest.
- 583

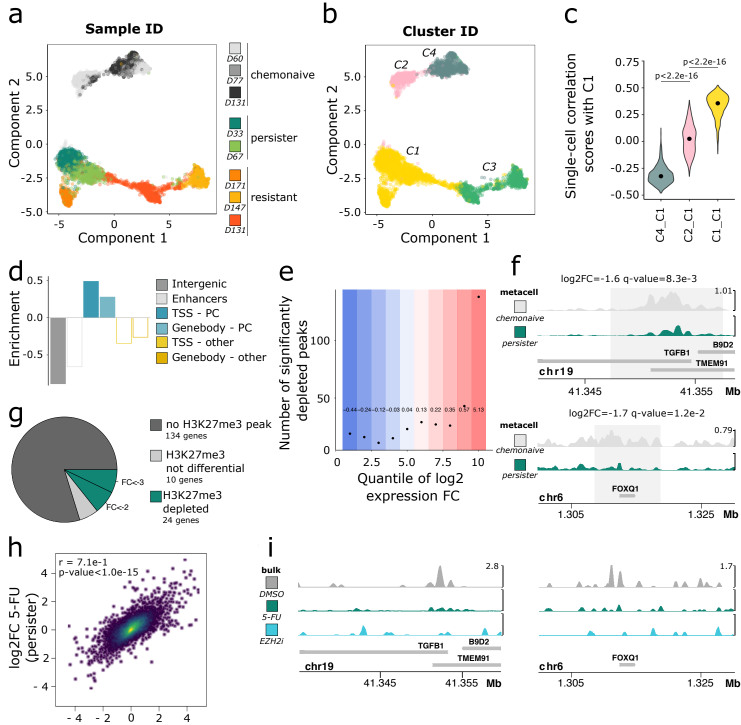
584 Additional Information.

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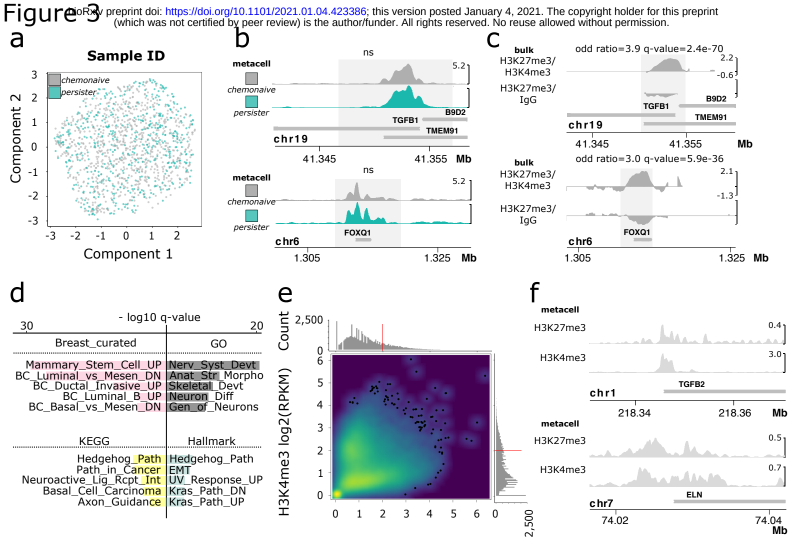
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log2FC EZH2i



H3K27me3 log2(RPKM) Count

