Single-cell analysis of EphA clustering phenotypes to probe cancer cell heterogeneity

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The Eph family of receptor tyrosine kinases is crucial for assembly and maintenance of healthy tissues. Dysfunction in Eph signaling is causally associated with cancer progression. In breast cancer cells, dysregulated Eph signaling has been linked to alterations in receptor clustering abilities. Here, we implemented a single-cell assay and a scoring scheme to systematically probe the spatial organization of activated EphA receptors in multiple carcinoma cells. We show that cancer cells retain EphA clustering phenotype over several generations, and the degree of clustering reported for migration potential both at population and single-cell levels. Finally, using patient-derived cancer lines, we probed the evolution of EphA signaling in cell populations that underwent metastatic transformation and acquisition of drug resistance. Taken together, our scalable approach provides a reliable scoring scheme for EphA clustering that is consistent over multiple carcinomas and can assay heterogeneity of cancer cell populations in a cost- and time-effective manner.
Receptor Tyrosine Kinases (RTK) are critical for tissue homeostasis and their dysfunction is associated with cancer progression. Eph receptors, which comprise the largest family of RTK, interact with ephrin ligands expressed on the surfaces of neighboring cells. In particular, an increase in EphA activity has been implicated in many cancer types including 40% of breast cancers. This juxtacline receptor-ligand signaling creates significant opportunities for spatial and mechanical constraints at the cell–cell interface to become intermingled with signaling behavior. Recent studies on breast cancer cell lines reported that cytoskeleton remodeling and EphA2 signaling interact in a feedback loop to control the clustering of EphA2, which specifically binds to ephrinA1 ligands. Alterations in this feedback loop result in changes in receptor aggregate morphologies, which, in turn, lead to phenotypic changes in cell morphologies and behaviors. Seminal reports strongly suggest that the activities of EphA2, and of other Eph receptors, are regulated by their ability to spatially cluster in a cytoskeleton-dependent way. The cluster morphologies correlate with the invasion potentials of breast cancers and cell lines at the population level. Therefore, we hypothesized that developing a generic scoring scheme for individual cells could provide a robust phenotypic assay to ascertain the heterogeneity of cellular functional states within a cancer cell population, which could be compared across cancers of different origin. In particular, it could probe intra-tumor functional heterogeneity of EphA signaling in a cost- and time-effective manner. Such assays could also complement single-cell transcriptomic approaches by integrating gene expression profiles with essential information about cell behavior and cell-state in an unbiased way at the single-cell level.

In this paper, we developed a simple phenotypic assay based on a general scoring scheme to quantify EphA clustering induced by ephrinA1 ligation that we standardize amongst a variety of human cancer cell lines of different origins. We demonstrated that cluster morphology is an inheritable trait that propagates over cell divisions for several generations, and that it correlates with the migratory potential of single cells. Finally, we validated that our assay predicts alterations in EphA activation pathway in cancer cell populations.

Results

Working principles and work flow of the assay. In this assay, we used supported lipid bilayers functionalized with fluorescently-tagged (Alexa 568) ephrinA1 to allow for cytoskeleton-dependent clustering of EphA receptors. Upon ligand binding, the receptors oligomerize and get trans-phosphorylated. The mobile ligand/receptor pairs then aggregate into clusters driven by cytoskeleton rearrangements. Using single-cell image analysis, we quantified cluster morphologies based on scalar scores that reflected their degree of aggregation. The distribution of individual scores provided information about intra-tumor heterogeneity. Figure 1a summarizes the experimental scheme that is detailed in “Methods” section. In brief, a silicon gasket (20 × 20 × 0.3 mm), comprising of nine independent recording chambers (3 × 3 mm), was layered on a clean, hydrophilic glass coverslip (Supplementary Fig. 1a). We generated supported bilayers (96 mol% DOPE and 4% DOGS-NTA-Ni) by standard small unilamellar vesicle (SUV) deposition (“Methods” section) in millimeter-sized chambers. Polyhistidine–ephrinA1 labeled with Alexa 568 were conjugated with the NTA lipid moieties at saturating densities (4%). We optimized the size of the chambers and the spatial arrangement of the duplicates to achieve high reproducibility and to allow internal quality controls (Supplementary Figs. 1b, c). Following sedimentation, suspended cells (~100 cells per chamber) establish contact with the bilayer within 30–60 s (Fig. 1b). Upon establishment of contact, EphA receptors on the cell surfaces cluster and engage with the ligands freely diffusing on the bilayer (Fig. 1c). For all the cell types we tested, receptor aggregates reached a steady morphology within a few minutes to half an hour. Thereafter, these structures remained largely unaltered for up to 1 h. Our samples were fixed after 1 h to capture statistically significant differences in EphA clustering morphologies at steady state. Individual imaging of 150–200 cells typically took an hour. In this study, a manual implementation of the assay proved sufficient. Optimizing automated imaging and analysis of larger cell populations allowed by the scalable format of the assay (Fig. 1a–c) was left for further development.

EphA clustering in cancer cells of different origins. We first tested whether cells originating from different cancer types displayed variations in EphA clustering, as seen in breast cancer cell lines (Fig. 1d, e). Figure 2a displays the cluster morphologies obtained for seven ovarian (PEO1, SKOV3, OVCA420, OVCA429, A2780, HeyA8, and OVCA101), one lung (A549), one gastric (MKN28), and three breast cancer cell lines (MCF10A, MCF7, MDA MB231). Reflection interference contrast microscopy images confirmed that the EphA clusters localized in regions where cells established intimate contact with the bilayer (Fig. 2b).

Interestingly, the steady state variations in cluster morphologies between cell types resulted from differences in receptor–ligand radial transport (Fig. 2b–d and Supplementary Movies 1 and 2). Some cell types displayed small aggregates with very little cell-to-cell variability, whereas other cell types presented larger and nonhomogeneous cluster morphologies. For example, PEO1 cells presented small EphA clusters with limited mobility over 30 min of observation (Fig. 2c, left and kymograph in Fig. 2d, top). In contrast, EphA puncta in HeyA8 cells were quickly transported toward the inner regions of the cells (Fig. 2c, right), where they fused into aggregates. These aggregates further merged with other aggregates (kymograph in Fig. 2d, bottom) to form a series of larger clusters (Fig. 2b). In these experiments, we have also determined that for all cell types steady state was reached at 30 min after cell/bilayer contact and clustering morphologies remained unchanged from that point onwards. Thus, we performed all the following EphA experiments by fixing the cells at 60 min.

Scoring cluster morphologies. We hypothesized that some morphological features of the EphA clusters could reflect a general cell response to ephrinA1 binding that was independent of cancer type, whereas a different set of morphological features would be cancer-type specific. We thus probed 75 morphological descriptors of the cluster patterns for their specificity, using a combination of t-distributed stochastic neighbor embedding (t-SNE) algorithm and a machine-learning method (Random Forests) (see Supplementary Note, Supplementary Tables 1 and 2 and Supplementary Figs. 2 and 3). This unbiased study singled out the intensive descriptor $S_{EphA} = \rho \ln \rho$, where $\rho = \frac{I_{mean}}{I_{max}}$, is the ratio of the fluorescence intensity value $I$ of each pixel divided by the average fluorescence intensity $I_{max}$ under each cell (Supplementary Fig. 4). $S_{EphA}$ emerged as the best parameter to score the scattering degree of EphA clusters while ignoring their specific spatial arrangements that mostly reflected the cancer cell type of origin (Supplementary Fig. 4 for justifications). Morphologies with scattered small puncta resulted in low $S_{EphA}$.
scores while morphologies with large aggregates resulted in higher scores. Figure 3a shows the distribution of single-cell scores obtained for the epithelial-type PEO1 carcinoma cells and the mesenchymal-type HeyA8 cells. We linked individual scores to the corresponding cluster images. Whereas PEO1 cells showed homogenously limited clustering, HeyA8 cells displayed a high variability in cluster sizes, which correlated with high variability in $S_{\text{EphA}}$ scores. In this heterogeneous population, cells with scattered small clusters obtained low $S_{\text{EphA}}$ values (consistently with PEO1) and cells with large aggregated clusters obtained high $S_{\text{EphA}}$ values.

Quantitative assessment of $S_{\text{EphA}}$ distribution. We first tested the reproducibility of the determination of the $S_{\text{EphA}}$ score distribution over several cell types. Repeated measurements for the same cell population on different chips resulted in qualitatively identical salient features of the distributions (narrow vs. spread distribution). However, inter-chip variability limited the reproducibility of quantitative data. We noticed that with every chip, the experimental variations for all the tested cell lines were correlated. This suggested that quantitative changes arose from day-to-day fluctuations in the detection system rather than from cell-intrinsic noise. We alleviated these variations by implementing an
intrinsic normalization scheme. On every chip, we measured triplicates of MDA MB231 cells that were used as a reference cell line (Supplementary Fig. 1b). Supplementary Fig. 1c shows an example of the intra-chip and inter-chip variability for each triplicate in three independent chips. As a quality control measure for device preparation, we considered only those individual chips for which the MDA MB231 triplicates showed no significant differences ($p > 0.05$ using a Kruskal–Wallis nonparametric test with Dunn’s multiple comparison applicable to non-Gaussian distributions). When this criterion was not met (in less than 20% of chips), the entire chip was excluded from analysis. For each selected chip, we then combined the distribution obtained for the MDA MB231 triplicates into a single distribution and used its mean value to normalize all the distributions measured on the entire chip. Figure 3b illustrates the combined distribution of triplicates of MDA MB231 cells (breast cancer) and HN137m cells (patient-derived squamous head-and-neck cancer cell) run over six different chips. Pairwise comparison demonstrates the inter-chip variability observed experimentally. Due to the proportionality of the pairwise inter-chip variability, we could normalize all the scores for a given chip by the averaged value of the MDA MB231 distribution ($N > 150$ cells). Normalized values are represented as $\tilde{S}_{\text{EphA}}$, where $\tilde{S}_{\text{EphA}} = \frac{S_{\text{EphA}}}{S_{\text{EphA, MDA MB231}}}$. The use of a reference cell line is a simple strategy that provides dual advantage: a method for controlling chip quality and a scheme to reduce inter-chip variability below 5% (Fig. 3c).

**Heterogeneity of clustering within diverse cell populations.** We then used the above approach to quantify the distribution of
The intercellular variability in clustering morphologies is exempliﬁed for PEO1 (seven cells) and HeyA8 (eight cells) lines (Scale bar: 5 μm). The matching between clustering morphologies and SEphA scores is exempliﬁed for PEO1 (seven cells) and HeyA8 (eight cells) lines (Scale bar: 5 μm). Normalization of all distributions on individual chips by the average scores of triplicates of the reference cell line MDA MB231 abolishes signiﬁcant inter-chip variability. It allows quantitative comparison between chips and cell types. Each distribution is based on over 200 single-cell analysis. We established signiﬁcance using the two-sample Kolmogorov–Smirnov test that probes for changes in the shape of the distribution.

Fig. 3 Reproducible measurement of SEphA scores. a Distributions of SEphA scores for PEO1 and HeyA8 lines. Every PEO1 cell displays very scattered EphA scores, whereas more mesenchymal-like cell showed higher, widely-distributed SEphA scores. Independent of their mean values, the shape of the distributions differed between the cell lines (Fig. 4b). The combinatorial statistical differences between different populations are presented in Supplementary Table 3. While PEO1 cells predominantly displayed very low scores with a relatively long tail toward higher scores, A549 cells and MDA MB231 cells mostly displayed a symmetric distribution around their average scores. Kolmogorov–Smirnov tests revealed that considering distributions shapes together with their average values provided characteristics of each cell population with 90% accuracy (Supplementary Table 3). Taken together, the results shown in Figs. 3a and 4b suggest that the shapes of SEphA distribution, in addition to average scores, could be used to predict single-cell heterogeneity in each population, rather than being regarded as measurement noise.

EphA clustering ability is an inheritable trait. So far, we established that the distribution of EphA clustering could characterize the cell population. We then tested whether individual cell response was an intrinsic property of individual cells that could be retained even upon cell division. To this end, we adopted the following strategy. We used patient-derived primary cultures from head-and-neck tumor (speciﬁcally HN137p)22. We isolated and expanded 30 individual cells into clonal colonies for 10 days, following which we tested half of the cells from every colony with our assay. The other half was left to expand for another 10 days, and cells from every colony were tested again after 20 days. We then compared the distribution of scores for the original population (before cell isolation) with the distribution after 10 and 20 days for each clonal expansion. Nine colonies survived this screening process (Fig. 5a).

During the ﬁrst 10 days (seven to nine cell divisions), we monitored the expansion of clonal colonies using phase contrast microscopy. Some colonies displayed very homogeneous morphologies (e.g., colony 1) with well-deﬁned cell–cell contacts and smooth edges, a typical characteristic of epithelial-like cells. Other colonies, however, appeared more heterogeneous (e.g., colony 9) with large variations in their individual cell phenotypes, a lower degree of contact inhibition, and more diffuse edges (Fig. 5b).
dropped to 16.7% at 20 days demonstrating the cells with very low. Of note, reconstituted populations after 10 and 20 days lacked of the original population diversity from each clonal expansion.

Fig. 4 Measuring $S_{\text{EphA}}$ distributions across cell types. $S_{\text{EphA}}$ distributions for a panel of breast, ovarian, lung and gastric cancer cell lines. All cell types displayed cluster morphologies that could be quantitatively assessed in our assay, leading to significantly different average value ($N > 200$ for each line).

**Notably**, at 10 days after reconstitution, the colonies displayed homogeneous epithelial morphologies (e.g., colony 1 or 2) displayed $S_{\text{EphA}}$ scores significantly lower than their unsorted population average. Conversely, $S_{\text{EphA}}$ scores higher than the population average characterized the colonies with more irregular appearances (e.g., colonies 8 or 9). At day 10, each colony displayed a different average $S_{\text{EphA}}$ score (Fig. 5d, top). However, intra-colony variability was small as shown by the narrowly distributed scores around their respective average values (Fig. 5d, bottom). After 20 days of culture, the colonies evolved toward similar $S_{\text{EphA}}$ averages (Fig. 5d, top), which converged toward the average value measured for the entire population. Concomitantly, intra-colony variations increased substantially as shown by the larger coefficients of variation of $S_{\text{EphA}}$ distributions (Fig. 5d, bottom). The growth of clonal colonies limits the number of cells available for this analysis. However, confidence analysis (Supplementary Note Information and Supplementary Fig. 5) demonstrates that the distribution of scores obtained for each colony cannot be interpreted (with 99.5% confidence) as a random subset of the unsorted distribution, instead they must be attributed to the phenotypes of the initial clones. In addition, after 10 days of growth, 45.5% of the colonies significantly differed from each other (Kruskal–Wallis with Dunns posttest). This ratio dropped to 16.7% at 20 days demonstrating the "reconstitution" of the original population diversity from each clonal expansion. Of note, reconstituted populations after 10 and 20 days lacked cells with very low $S_{\text{EphA}}$ scores (below 0.2). Instead, an increasing fraction of cells presented $S_{\text{EphA}}$ larger than 0.6. This observation was consistent with the idea that expanding clonal colonies did not favor the growth of the most epithelial-like cellular types (only nine cells could be expanded out of the initial 30 cells, with most of them being quiescent or apoptotic). During proliferation, most epithelial phenotypes were not reconstituted, and cells appeared to drift toward the more mesenchymal cell types.

Altogether our results, suggest that EphA cluster morphology of a single cell is an inheritable trait that gradually diverges upon sequential divisions to progressively regenerate the initial population diversity.

$S_{\text{EphA}}$ correlates with cell migration potential. In breast cancer cells, different EphA activation levels were associated with different invasion potentials at the population level. This correlation was attributed to pathways modulating the actin cytoskeleton, that could induce EphA clustering as well as promote cell migration. We therefore tested how phenotypic characterizations of EphA clustering overlapped with other standard phenotypic assay, including the migratory potential of cancer cells.

Using microchip arrays, we first measured the transcriptional levels of all EphA receptors in all our cell lines. EphA2 proved to be the most abundantly expressed receptor in 8 out of the 11 cell lines (Supplementary Table 4). $S_{\text{EphA}}$ scores at the population level (Supplementary Fig. 6) did not correlate with the mRNA levels of EphA2 and also with all EphA pulled together. The correlation greatly improved when EphA2 protein expression levels (Supplementary Fig. 6b, c, Western blot) were considered. We then tested if the cellular phenotypes defined by our assay correlated with cancer hallmarks, epithelial mesenchymal...
The spreads of all distributions increase simultaneously, indicating a larger variability in cell behavior. Statistical analyses of populations (Kruskal-Wallis with Dunn’s posttest) and confidence test (Bootstrapping) are discussed in Supplementary Note.

Some of our cell lines clearly showed heterogeneity in migratory behaviors within the cell population as it often occurs in cancer cells with intermediate EMT score (Fig. 6a, right). We thus explored whether this heterogeneity was also reflected in EphA distribution patterns. In the three cell lines that we used—A549 (lung), SKOV3 (ovarian), HN137m (head-and-neck PDGC)—migration speeds were substantially heterogeneous, and, correspondingly, their EphA distributions were also scattered. We then used a modified wound-model assay to isolate the fast-migratory cells from the slow ones within each cell population and compared the EphA distributions between the different subpopulations (fast-migratory cells, slow migratory cells, unsorted populations) (Fig. 6b). We initially confined the cells on 300-micron islands on a PDMS sheet coated with fibronectin (see “Methods” section) and, after removal of the confinement, we allowed migration in 2D for 3 days. We then punched-out the central part of the colonies with a 750-micron punch, thereby sorting cells that had migrated more than 250 microns away from their initial position. In all cell types, fast migrating cells displayed transition scores (EMT) and their migration potential. Interestingly, SEphA did not correlate with any of the listed cancer hallmarks (Supplementary Table 5). We then computed EMT scores for all cell types according to Tan et al. by measuring the transcription levels of a list of 118 genes. Overall, EMT scores poorly correlated with SEphA (Fig. 6a, left; dash line, Pearson coefficient = 0.34). However, we obtained a much higher correlation by subdividing the cell lines into two groups using SEphA = 0.4 as threshold (Fig. 6a, left; dotted lines, Pearson coefficients = 0.83 for group with SEphA > 0.4, and Pearson coefficients = 0.99 for group with SEphA < 0.4). We then systematically performed wound-model experiments to assess the migration potentials of all cell lines. Interestingly, cell motility speed highly correlated with SEphA (Fig. 6a, right; Pearson coefficient = 0.89). Furthermore, we observed that groups of cells lines with equivalent EMT scores, but different SEphA score (colored ovals in Fig. 6a, left) had as a common discriminator their cell migration potential and cell lines in the group with SEphA > 0.4 were fast and SEphA < 0.4 were slow migratory. These results suggest that the strong phenotypic link between EphA clustering and migration potential established for breast cancer cell lines may remain valid for cancer cell lines of different origin.

Thirty individual cells were picked from the unsorted population of HN137p cells, out of which nine clonal colonies could grow for 20 days (a). Phase contrast images of the clonal colonies over time. Some colonies display a very homogeneous aspect (e.g., colony 1), some others show a noticeable spread in cell morphologies (e.g., colony 9) (Scale bar: 100 µm). b Close-up of colonies 1 and 9 after 20 days (Scale bar: 25 µm). c SEphA distributions for the unsorted cell population, for populations reconstituted from the nine colonies at 10 and 20 days, and for each individual colony at both time points. Note the absence of low scores in distributions converge toward the same distribution, indicating the genomic drift of the cellular states over time. d Convergence of the mean score for every colony between day 10 and day 20. The spreads of all distributions increase simultaneously, indicating a larger variability in cell behavior. Statistical analysis of populations (Kruskal-Wallis with Dunn’s posttest) and confidence test (Bootstrapping) are discussed in Supplementary Note.
EphA distribution with a higher average value and higher spread (Fig. 6c).

These results demonstrated the correlation between EphA clustering and migration potential at the population level across diverse cancer cell types. Furthermore, enrichment of cells with high EphA in subpopulations selected for their migration potential support the idea that EphA correlate with single-cell migration potential. EphA scores hence appear as a fast and reliable measurement of the migration potential of cells and heterogeneity of their invasion potential within a given population.

**Fig. 6 Correlation between EphA scores EMT score and migration speed.**

a Correlation of the average EphA with the EMT scores (based on 118 genes). The correlation is globally poor, but it largely increases (Pearson coefficient -0.85) if cell types are split into slow and fast migrating cells. The correlation between the average EphA with the mean mobility of cells in the migration front of the gap-closure assay for each population is relatively high (Pearson coefficient ~0.89).

b Schematic drawing of our separation assay based on 2D migration. Cells are plated on 300-μm islands obtained by protein printing. The cells are left to migrate away from the initial island. Using a 750-μm punch, we then separated the fast migrating cells that migrated more than 750 μm away from the initial island center, from the rest.

c EphA distributions for unsorted, slow and fast populations from A549 (lung), SKOV3 (ovarian), HN137m (head-and-neck) cancer cell types. Note that the fast (respectively slow) subpopulations systematically higher (respectively lower) EphA scores than the unsorted population. N > 200 cells.
**S**\(_{\text{EphA}}\) as a measure of evolution of disease. Finally, we tested whether our assay could follow changes in the cellular phenotypes during cancer progression in patient-derived models. We have previously reported the generation of multiple primary cancer cell lines from three different head-and-neck patient tumor samples\(^{22,27}\). These include HN137, HN120, and HN148 cell lines, that were generated from the primary site of tumors (HN120p, HN137p, HN148p) or the paired lymph node metastatic sites (HN120m, HN137m, HN148m). In addition, we also generated in vitro acquired-resistance models for cisplatin for those cell lines derived from the primary site (HN137pcr, HN120pcr)\(^{27}\). Extensive characterization of these cells demonstrated that they all presented very distinct transcriptomic profiles and migratory phenotypes\(^{22,27}\). We therefore tested whether these previously reported characteristics were reflected in the distribution of \(S_{\text{EphA}}\) for these distinct eight PDCs.

Figure 7a shows that for patient HN120, the \(S_{\text{EphA}}\) distributions from primary, metastatic and cisplatin resistant cell lines displayed increasing average values. This observation correlated with the increased migration potential observed across these cell lines\(^{22,27}\). In addition, HN120p and HN120pcr cells displayed very different \(S_{\text{EphA}}\) scores, corroborating the previous observation that HN120pcr cells are transcriptomically distinct from HN120p. This conclusion was also supported by comparing the \(S_{\text{EphA}}\) scores to the single-cell transcriptomic profiles (N > 100 cells) established based on the expression levels of 71 genes, which have been reported to be downstream of EphA2 signaling cascade\(^{15}\) (Supplementary Table 6). The transcriptional scores confirmed that all PDCs lines were distinct. Importantly, EphA2 expression profile of the HN120 lines ranked similar to and correlated with the matched \(S_{\text{EphA}}\) scores.

Our results from patient HN137 (Fig. 7b) showed that HN137p and HN137pcr cells did not display any significant differences in \(S_{\text{EphA}}\) scores. However, scores in HN137m cells were slightly higher than that observed in HN137p, in line with the observed invasion potential\(^{22,27}\). A similar trend was reflected in the transcriptomic scores. Patient HN148 (Fig. 7c) displayed equal scores in HN148p and HN148m cells, although the transcriptional scores were slightly smaller in HN148m cells than in HN148p cells. Furthermore, average population values of \(S_{\text{EphA}}\) scores and transcription score displayed strong correlation (Pearson coefficient = 0.93, \(p = 0.001\)), thereby indicating that our clustering score provides a reliable measure for EphA associated gene expression profile and phenotypes (Fig. 7d).

Overall, single-cell analysis of EphA clustering behavior and transcriptomic response in PDCs showed that our assay provided a reasonable assessment of the evolution of cell-states within these patient samples in a fast and cost-effective manner.

**Discussion**

Intra-tumor heterogeneity poses a major challenge in therapeutic outcomes for cancer treatment in the clinic. Different clonal or sub-clonal cell populations can exhibit differential response to a
given drug, and utilize distinct mechanisms to evolve therapy resistance, such as cell-state change like EMT or dormancy induced quiescence. Therefore, it is important to develop methodologies that can probe this heterogeneity to be able to identify novel genetic and therapeutic vulnerabilities. While existing single-cell genomic/transcriptomic platforms have been shown to be extremely powerful in defining intra-tumor heterogeneity and the identification of new cell types and cell states, methodologies that can uncover functional heterogeneity at the level of single cells remain largely under-explored. In our study, we show that EphA cluster morphologies, which are associated with distinct migratory state of cells, constitute a single-cell-inherited trait of that can uncover functional heterogeneity at the level of single cells. We developed a common analysis scheme to quantitatively correlate cluster morphologies with the intrinsic migration potentials of cells. In addition, results from this study extend to multiple cancer types our previous conclusions based on breast cancer cell lines that cluster morphologies are indicative of the degree of transcriptional activation of genes involved in EphA signaling cascade. This signaling pathway largely involves cytoskeleton regulatory genes, which can explain the correlation between cluster morphologies and migration properties. Oppositely the other gene pools that we tested (Supplementary Table 5) hardly involve cytoskeleton components. We thus believe that the absence of correlation with these pathways is additional evidence that our scoring approach targets EphA signaling pathways specifically. By using PDCs from various cancer types, our study reinforces the growing body of evidence that clustering properties of juxtarce receptors can be used to define the “phenotypic state” of single cells. We conclude that this assay provides a technically simple, cost-effective and effective strategy to follow the gradual buildup of intracellular heterogeneity either upon cell division, (Fig. 5) or across genomic drifts and drug selection, (Fig. 7). Such an approach is nonetheless restricted to ligands that foster cell adhesions and spreading on fluid bilayers. If EphB4 ligands have already been tested, the extension to other non-membrane tethered RTK need to be explored. In this case, alternative approaches such as coupling several ligands together, could be devised. The assay can also be performed in a 4 x 4 mm well that are the standard size of the compartments in a 384 well plate. We thus provided proof of concepts that the assay can be scaled up. However, this would entail the parallel formation of lipid bilayers in each well with quality control. We foresee that only automatically dispensing can provide sufficient reproducibility in lipid deposition and rinsing to ensure high enough quality bilayer. The reproducibility we achieved here relied on the normalization of each batch data by the scores of MDA MB231 cell line measured on the same day on the same chip. Using a reference line was handy to check the fluidity of the bilayer, to check the quality of the ligand and to perform a normalization of the acquisition conditions. One could argue that a physical calibration of the system (e.g., using fluorescent beads and Fluorescence Recovery After Photobleaching) would alleviate the constraints of using a reference cell line that might differ from lab to labs or drift over time. This process could indeed be implemented on a calibrated microscope equipped with a FRAP module. Such calibration could be performed systematically before each experiment. However, we found no significant drifts in MDA MB231 used no more than 12 passages from the commercial batch. They are also suited for calibration due to the large variability of the distribution of their scores as compared with epithelial cell lines such as PEO1. A systematic comparison of data acquired in different lab using our scheme remains to be done. Furthermore, the procedure to determine cell contours and cluster morphology can already be performed without any user assistance. It could thus be potentially incorporated in a detection pipeline provided that an automated imaging routine is developed.

Lastly, the rationale to develop parallelizable phenotypic assays stems from the observation that the levels of genes related to a certain pathway do not always directly correlate with the behavioral response of cells. It is therefore important to develop cost-effective and simple behavioral assays that allow phenotypic probing of response to an activated pathway. We believe that this work is a step forward in this direction.

**Methods**

**Preparation of SUV.** A clean, 5 ml round-bottom flask (PYREX) was rinsed with Milli-Q water and it was washed by vigorous sideways shaking to remove all traces of organic solvent. Thereafter, the flask was rinsed with Milli-Q water and placed in an 80 °C dry oven for at least 2 h. Meanwhile, a PDMS silicon foil of 0.25 mm thickness (STERNE SAS, France) was cut into square chips of 2 x 20 mm sizes and further subdivided into nine recording chambers (3 x 3 mm) using a cutting plotter (Graphtec Corporation, Japan). Importantly, one of the corners of the square was cut to break the double symmetry and allow easy identification of wells (see Fig. 1a and Supplementary Fig. 1a, b). As the PDMS foil was polymerized in between two plastic films, the PDMS films were cut free from dust and contaminants. Finally, PDMS cuts were mounted on the glass coverslips (22 × 22 mm) from high precision. The PDMS cuts were free from dust and contaminants. Finally, PDMS cuts were rinsed in ethanol solution and sonicated for 30 min, submerged in 1:1 isopropanol/Milli-Q water solution (isopropanol from Fisher Chemical) for at least 1 h. The lipid suspension was then transferred to a round-bottom polystyrene tube and sonicated for 20 min until the suspension became milky white. The liposomes suspension was then mixed in chloroform, the cholesterol suspension was evaporated on the membrane was washed using imaging buffer (150 mM NaCl, 30 mM KCL, 2 mM CaCl2, 2 mM MgCl2, 10 mM Hepes, 10
mM D-Glucose, pH 7.4). The device was then placed in a petri dish and submerged in 2 ml of imaging buffer. Devices were freshly prepared and long-term storage was avoided. Devices were kept covered in the dark at the room temperature while waiting for the cells to be seeded. The devices were acclimated inside a 37 °C incubator for at least 1 h before cell seeding.

Cell culture, dissociation, seeding, and fixation. All the cells used in this study were tested for mycoplasma using Mycoprobe Mycoplasma Detection kit from R&D system. Cells were passaged no more than 10 times from the initial stocks. The lung adenocarcinoma cell lines were obtained commercially or were kindly donated by Dr. Ken Yamaguchi from Kyoto University Graduate School of Medicine. The isolation and derivation of the PDCs are extensively described in Ref. 21. Briefly, tumors were minced and dissociated enzymatically using 4 mg/ml collagenase type IV (Thermo Fisher, cat. no. 17104019) in DMEM/F12, at 37 °C for 2 h. The cell suspensions were strained through 70 µm cell strainers (Falcon, cat. no. 352350), prior to pelleting and resuspension in RPMI (Thermo Fisher, cat. no. 61870036), supplemented with 10% foetal bovine serum (Biowest, cat. no. S181B) and 1% penicillin-streptomycin (Thermo Fisher, cat. no. 15140122). The identities of the cell lines were checked using the STR profile (IndexDNA) of the cell line to its original tumor. A standard cell culture incubator was set at 37 °C with 5% CO₂; was used to expand all cells. The cells were subcultured at 80% confluence in T25 flasks (Thermo Fisher Scientific) and reseeded into fresh flasks at ~30–50% confluence. Prior to the experiments, cells were grown at 50–70% confluence and dissociated with an enzyme-free dissociation buffer (Gibco). In the meantime, the device was kept submerged in imaging media in a 35 mm petri dish and was acclimated inside an incubator at 37 °C. After removing the imaging buffer from the petri dish, the chambers of the device were left filled with imaging media (~2 µl). Care was taken in this very critical step to avoid exposing the lipid bilayer to air-liquid interface at all times. Immediately after removal of the media, a 5 µl of cell suspension was directly deposited on top of each chamber of the device, forming small drops with high contact angle due to the hydrophobicity of the silicon-based device. Spacing between the chambers kept cells contained within their respective chambers. The three cell types labeled with different fluorescent markers (stably transfected GFP and RFP cell lines and one with Hoechst live nuclear stain) were used to ensure the absence of overspill of cells from adjacent chambers. Due to the low volume and height of the chambers, seeded cells nearly immediately contacted the bilayer.

About 200 µl of sterile Milli-Q water was pipetted along the walls of the petri dishes to provide extra humidity and prevent evaporation of water from the chambers. Petri dishes were covered with a secondary container (typically a 140 mm petri dish with humidifying pads soaked in sterile Milli-Q water) previously acclimated at 37 °C. The secondary container was then placed inside a 37 °C incubator for 30 min to allow EphA clustering to occur at nearly physiological conditions. After 30 min, clustering was stopped by PFA fixation: devices inside the 35 mm petri dish were flooded with 2 ml of 4% PFA solution (Thermoscientific). Cells were fixed at 37 °C for 15 min. Thereafter, devices were washed with 1× PBS solution for five times and incubated in 10 mM NH₄Cl solution for 10 min at the room temperature. Devices were then washed with 1× PBS solution for three times and mounted on glass slides using fluorescent mounting medium (Dako) and the coverslips were sealed using nail polish.

Microscopy and image analysis. Fixed samples were imaged using an IX81 microscope (Olympus) equipped with an ×100 oil-immersion objective (UAP200XIRF, NA 1.49, Olympus), X-Cite light source (Excitels Technologies), a CoolSNAP EZ CCD camera (Photronics) and an appropriate filter set to detect Alexa 568 conjugated to ephrinA1. Pairs of bright-field and fluorescence images were acquired for each field of view. The focus of the objective was adjusted to visualize the supported bilayer. Images were analyzed using a custom MATLAB code (can be provided upon request). The code performed bunch analysis of images found in the input folder. The analysis was divided into two separate steps: first, cell segmentation was performed by manually outlining cell borders on the confidence maps generated from the bright-field and corresponding fluorescence images. This process defines the regions of interest for subsequent analysis and generates a third masking image where the ROI are defined with ascending natural numbers. In the second step, image intensity distribution from the ROIs was analyzed and the image entropy for each cell quantified according to $S_{\text{EPH}} = -\int \rho \ln \rho \, d\sigma$, where $\rho = \frac{\rho_{\text{Eph}}}{{\rho_{\text{Eph}}}}$. For live-cell images, trajectory analysis shown in Fig. 2c has been performed using ImageJ manual tracking plugin.

Western blotting. Protein lysates (30 µg) from individual cell lines were firstly dissolved in SDS-PAGE loading buffer and loaded onto 10% SDS-PAGE gel. Using Bio-Rad Mini-PROTEAN electrophoresis system, protein samples were run at 110 V, until samples ran completely through the stacking gel and into the resolving gel. The voltage was then increased to 150 V, until dye front ran off the bottom of the gel. After protein separation by SDS-PAGE, proteins were transferred from gel to a PVDF membrane (Mercer, cat. no. IPV07852). Transfer was carried out for 45 min at 110 V in transfer buffer (cold room). Following this, the PVDF membrane was blocked with 5% skim milk powder in TBS-Tween 20 (0.1% v/v) at the room temperature for ~1 h. Anti-EphA2 (Cell Signaling, cat. no. 6997) and anti-beta actin (Fierce, cat. no. MA51739) were used as primary antibodies and diluted in blotting solution. The membrane was first probed with primary antibodies at 4 °C overnight. Anti-rabbit HRP (Invitrogen, cat. no. G21234) and anti-mouse HRP (Invitrogen, cat. no. G21040) were used as secondary antibodies and diluted in blocking solution. The membrane was then probed with secondary antibodies at the room temperature for 1 h. The membrane was washed three times for 10 min in TBS-Tween20 (0.1% v/v) on an orbital shaker after blocking with milk and after each probing step. Finally, the membrane was treated with chemiluminescent substrate (ThermoScientific, cat. no. 34080) and protein bands were detected by Bio-Rad Chemidoc imaging system.

Wound-model assay. Classical wound-model assays have been performed31,32. Briefly, a rectangular cut of PDMs was placed in the middle of a 6-well plate and let to adhere for 2 h at 37 °C before cell seeding was performed. Cancer cells were seeded around the PDMs cut and allowed to grow to full confluence inside a cell culture incubator for about 1 day. After removal of the PDMs cut, the 6-well plate was placed inside a BioStation CT (Nikon) and time lapses of cell migration by phase contrast were acquired for 12 h (time interval between image acquisitions was 15 min). Cell movement within the first 50 µm from the wound edge was analyzed using Particle Image Velocimetry in MATLAB.

Single-cell transcriptomic analysis. We previously generated single-cell gene expression matrix for 682 cells from PDCs from head-and-neck (HN) carcinomas from primary site (P), secondary/metastatic tumor (M), and primary with acquired cisplatin resistances (PCR) from three patients (patients 120, 137 and 148). Thus, PDC nomenclature is HN120P, HN120M, HN120pcr, HN137P, HN137M, HN137pcr, HN148P and HN148M. The references and the raw data to obtain the transcriptomic analysis are fully available from the references above. The expression levels were quantified as $E = \log_{2}(TPM + 1)$. We selected only the subset of genes potentially involved in the EphA1 signaling cascade (Supplementary Table 6)15. Potential markers with average expression levels lower than 1 across all the cells were excluded. We defined single-cell genomic score as the linear average of the expression levels of the remaining genes. Transcriptomic scores were normalized by average score of primary samples of each patient to account for individual genetic background. Few cells with low EphA2 score (<3) were discarded as outliers.

Statistics and reproducibility. Statistical analysis was performed using GraphPad Prism software. For all statistical analysis data from at least three biological repeats performed on separate days was used.

Data availability
Previously generated single-cell transcriptomics data from patient-derived cell lines are available in GEO, accession number GSE117872. Raw data behind the graphs are available in Supplementary Data 1. All original images, data, and MATLAB codes are available from the corresponding author upon reasonable request. All the data are currently stored in data storage facility at Mechanobiology Institute, Singapore.

Code availability
Custom made Matlab code for image analysis is available from the corresponding author upon reasonable request. Python codes used for image feature extraction and unbiased machine-learning method are available at Github: https://github.com/aneeshaethe/Ephrin_cluster_analysis. Data analysis, statistics, and presentation have been made using GraphPad Prism 6 Version 6.01.

Received: 19 September 2019; Accepted: 29 June 2020; Published online: 06 August 2020

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Acknowledgements

V.V. and J.T.G. acknowledge the support of the collaborative NRF CRP11-2012-02 grant, and MBI seed funds. A.R. and C.B. acknowledge financial support from seed funds of the Pontificia Universidad Católica de Chile. R.D. acknowledges funding partly from the National Medical Research Council (Singapore) (NMRC/CIRG/1434/2015).

Proof reading was kindly provided by Andrew Wong and Sruthi Jagannatha. The cell lines were kindly given by Dr. Ken Yamaguchi from Kyoto University Graduate School of Medicine. We would like to extend our gratitude to all patients and families involved in this study.

Author contributions

A.R., M.Z.M. performed the experiments. A.R., S.C., A.A., E.Y.C., C.B. performed the data analysis. A.S. devised the unsupervised clustering. S.C., A.A., N.G.I. produced the published maps and institutional affiliations.

Competing interests

The authors declare no competing interests.

Additional information

Supplementary information is available for this paper at https://doi.org/10.1038/s42003-020-01136-4.

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