Single-cell western blotting

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To measure cell-to-cell variation in protein-mediated functions, we developed an approach to conduct ~103 concurrent single-cell western blots (scWesterns) in ~4 h. A microscope slide supporting a 30-µm-thick photoactive polyacrylamide gel enables western blotting: settling of single cells into microwells, lysis in situ, gel electrophoresis, photoinitiated blotting to immobilize proteins and antibody probing. We applied this scWestern method to monitor single-cell differentiation of rat neural stem cells and responses to mitogen stimulation. The scWestern quantified target proteins even with off-target antibody binding, multiplexed to 11 protein targets per single cell with detection thresholds of <30,000 molecules, and supported analyses of low starting cell numbers (~200) when integrated with FACS. The scWestern overcomes limitations of antibody fidelity and sensitivity in other single-cell protein analysis methods and constitutes a versatile tool for the study of complex cell populations at single-cell resolution.

Heterogeneity is inherent in cellular processes including stem cell differentiation^{1,2}, development³, cancer^{4,5}, pharmaceutical efficacy⁶ and immune response⁷. Owing in large part to recent technological advances, genomic and transcriptomic studies of cell-to-cell heterogeneity are flourishing^{4,8}. However, recent singlecell and population-wide studies comparing transcriptomes to proteomes in microorganismal and mammalian cells found only mild correlations between mRNA and protein expression^{9–11}. Therefore, to fully understand diverse and often rare behaviors in complex cell populations, researchers need analytical tools that are optimized for protein analysis of many cells, offer single-cell resolution, provide quantitative and highly specific detection of target proteins, and do not employ labels that may perturb protein and cell function¹².

Single-cell proteome-wide studies are currently limited to readouts from synthetic fluorescent-protein fusion libraries^{9,11}, which, though illuminating, are challenging to generate and can potentially perturb protein function. Single-cell protein immunoassays (for example, flow cytometry⁷ and immunocytochemistry (ICC)¹³) have proved immensely important for assessing cell-to-cell heterogeneity, yet existing methods depend

on analyte discrimination with antibody probes that often have limited specificity. This dependence on antibody probe quality restricts assay performance, as cross-reactivity can create misleading background signals that are difficult to correct for ^{14–16}, even with careful controls ^{17,18}. This vulnerability broadly affects antibody-based assays (such as ELISAs and protein microarrays¹⁴). The widely used western blot is less affected by antibody cross-reactivity because proteins are first separated by molecular mass (via electrophoresis) before the antibody probing step, thereby enabling clear discrimination between on-target and off-target signals, even in complex backgrounds such as cell lysates^{19,20}. However, the cell population averaging required by existing blotting methods masks the rich single-cell behaviors found in complex populations^{20,21}. Although microwestern arrays afford remarkable target multiplexing and throughput, lysate pooled from ~10³ cells is required for each electrophoresis assay (~250 ng of protein)²⁰. Capillary and microfluidic designs reduce mass demands, but their form factors are not readily scalable to the thousands of concurrent electrophoresis assays required to measure variation within a population of single-cells²¹.

We address the need for high-specificity protein assays capable of measuring cell-to-cell heterogeneity within complex populations of cells by introducing the scWestern method. Specifically, a scalable open-microwell array architecture permits simultaneous assays of \sim 2,000 individual cells in <4 h. We applied the scWestern to study variability in stem cell signaling and differentiation responses to homogeneous *in vitro* stimuli.

RESULTS

Development and characterization of scWesterns

scWestern analysis employs a microscope slide coated with a thin photoactive polyacrylamide (PA) gel²¹ micropatterned with an array of 6,720 microwells (**Fig. 1**). The microwells (20 μm in diameter) are patterned during polymerization of a 30- μm -thick PA gel against a silicon wafer studded with SU-8 microposts (**Fig. 1a**). To allow for concurrent western analysis of thousands of single cells, the scWestern integrates all key western blotting steps (**Fig. 1b,c**) in a dense array format.

Three fundamental design principles underpin the scWestern. First, we address the scWestern globally in terms of fluidic, optical

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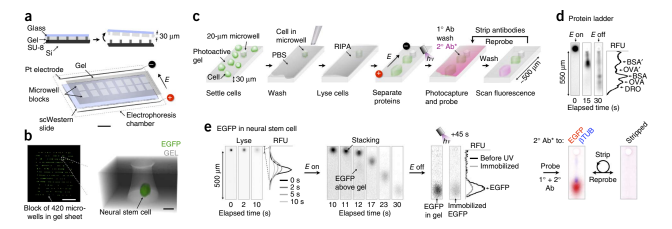


Figure 1 | Single-cell western blotting. (a) The scWestern array consists of thousands of microwells (20 μm in diameter, 30 μm deep) patterned in a 30-μm-thick photoactive polyacrylamide gel seated on a glass microscope slide. The array is comprised of 16 blocks of 14 × 30 microwells (6,720 in total) cast against an SU-8 photoresist master fabricated by soft lithography. E, electric field. Scale bar, 10 mm. (b) Left, wide-field micrograph of a microwell block containing 15-µm fluorescent microspheres. Scale bar, 2 mm. Right, confocal micrograph of a live EGFP-expressing NSC settled in a rhodamine-tagged gel (GEL). Scale bar, 10 µm. (c) Open-gel scWestern analysis is a 4-h, six-stage assay comprising cell settling, chemical lysis with a denaturing RIPA buffer, PAGE, UV-initiated protein immobilization onto the gel (hv, photon energy), diffusion-driven antibody probing (i.e., primary and fluorescently labeled secondary antibody probes: 1° Ab and 2° Ab*) and fluorescence imaging. (d) PAGE resolves five fluorescently labeled proteins in a 550-µm separation distance (DRO, Dronpa, 27 kDa; OVA, ovalbumin, 45 kDa; BSA, bovine serum albumin, 66 kDa; OVA', OVA dimer, 90 kDa; BSA', BSA dimer, 132 kDa). (e) scWestern analysis of EGFP and βTUB from a single NSC. RFU, relative fluorescence units. Distinct fluorescent dyes on each 2° Ab* enable multiplexed target analysis (EGFP, Alexa Fluor 488-labeled 2° Ab*; BTUB, Alexa Fluor 555-labeled 2° Ab*). Chemical stripping and reprobing allows multiplexed scWestern analysis.

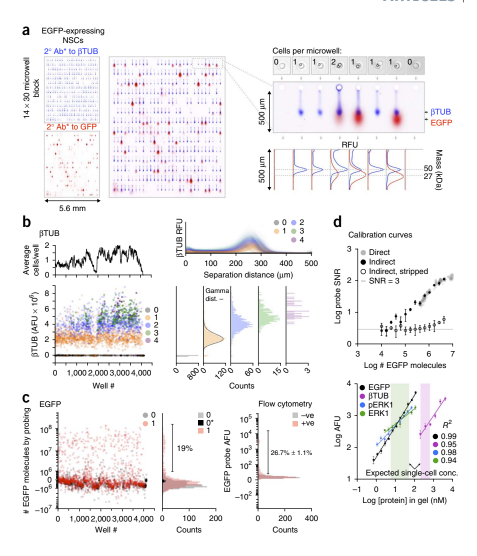
and electrical interfacing. Global interfacing yields highly parallel analyses by eliminating independent hardware access to each of the thousands of microwells. Initially, a cell suspension is seeded into microwells via passive gravity-driven cell settling, resulting in capture of 0-4 cells per microwell in 5-10 min. For neural stem cell (NSC) densities of 1,000-1,800 cells per mm² slide area $(2 \times 10^6 - 3.5 \times 10^6 \text{ cells in total})$, we observed single cells in 40-50%of microwells (Supplementary Fig. 1). Notably, FACS can be integrated with scWesterns to analyze subpopulations of ~200 cells with single-cell resolution (Supplementary Fig. 2), enabling analyses of rare or precious cells. Next, we perform buffer exchange to a denaturing radioimmunoprecipitation assay (RIPA) buffer that lyses cells in the microwells in 2.6 \pm 1.5 s (\pm s.d., n = 6 cells; **Supplementary Video 1**), solubilizing intracellular proteins while providing a suitable conductivity for subsequent electrophoresis. Protein diffusion from cells occurred within ~10 s of lysis (Fig. 1e). Simulations suggest that diffusion of cell contents from microwells is responsible for the moderate protein losses of $40.2\% \pm 3.6\%$ observed during lysis buffer introduction (\pm s.d., n = 3 microwells from three separate slides; Supplementary Note 1 and Supplementary Fig. 3). Future innovation in microwell enclosure methods or the use of higher-viscosity lysis buffers may reduce these losses.

As a second design principle, we achieve a high-density scWestern array by optimizing for short-separation-distance PA gel electrophoresis (PAGE). To initiate electrophoresis after cell lysis, we apply an electric field across the submerged scWestern slide, electrophoresing proteins through the microwell walls and into the thin PA gel sheet. To characterize this process, we assayed a ladder of purified fluorescently labeled proteins (27–132 kDa; **Fig. 1d**) that partition into microwells (Supplementary Note 2 and **Supplementary Fig. 4**). Under our denaturing, nonreducing PAGE conditions, we (i) observed stacking of purified proteins during electromigration into the bulk PA gel, (ii) verified a log-linear relationship between protein molecular mass and migration distance in scWestern separations, as anticipated for SDS-PAGE^{21,22} $(R^2 = 0.97;$ Supplementary Fig. 5) and (iii) resolved covalent protein dimers (Supplementary Fig. 6). Moderate PAGE performance was achieved, with molecular mass differences of 51% \pm 1.6% (\pm s.d., n=3 separations) resolvable in \sim 500- μ m separation lengths and 30-s separation times. In general, we observed agreement between scWestern separations and conventional western blotting (Supplementary Note 3). For comparison, a microwestern array setup (integrated with robotic bulk-cell lysate printing) offers similar resolving power but requires 18-fold longer separation distances (9 mm; ref. 20).

The third scWestern design principle harnesses small characteristic lengths for reaction (protein immobilization) and transport (antibody probing). Following PAGE, protein immobilization relies on a benzophenone methacrylamide co-monomer crosslinked into the PA gel. We measured protein photocapture in the gel at 27.5% \pm 2.9% of EGFP from EGFP-expressing NSCs after brief (45 s) gel exposure to UV light (\pm s.d., n = 6 single cells from experiments on 4 separate days; Fig. 1e). Photoimmobilization benefits from small diffusion lengths between proteins and benzophenone moieties within the PA gel²¹. Probing of the separated, immobilized proteins is performed by sequential diffusion of primary and secondary antibodies into the thin PA gel layer, taking advantage of the short 30-µm characteristic transport length (Supplementary Fig. 7 and Supplementary Note 2). scWestern antibody consumption is comparable to that of conventional western blotting and ICC, with potential for additional optimization (Supplementary Note 2).

Analysis of multiple protein targets is crucial to understanding cell functions such as signal transduction¹. Our scWesterns are organized into 16 assay 'blocks' of 420 microwells each, a layout that allows application of different antibody solutions

Figure 2 | scWestern blotting of NSCs. (a) 420 concurrent scWesterns of EGFP-expressing NSCs for βTUB (Alexa Fluor 647-labeled secondary antibody (2° Ab*)) and EGFP (Alexa Fluor 555-labeled 2° Ab*). Bright-field imaging determines the number of cells per microwell. RFU, relative fluorescence units. (b) Top right, scWestern fluorescence for 4,128 separations by cells per microwell. Left, area under the curve for βTUB with (above) running average of cells per microwell (window size, 30 microwells). Microwells are indexed from left to right of the array. AFU, arbitrary fluorescence units. Bottom right, fit of fluorescence distribution for single cells to a gamma distribution (dist.) stemming from Poissonian mRNA production and exponentially distributed protein burst sizes (Online Methods). (c) EGFP fluorescence for one- and zero-cells-per-microwell blots compared to flow cytometry of fixed NSCs (+ve, EGFP transfected; -ve, untransfected). Note arcsinh-transformed scales (Online Methods). Technical noise was estimated from scWesterns with zero cells per microwell in a sparsely cell-seeded region (0*, separations 4,100-4,128; Supplementary Note 5). The fraction of EGFP+ cells is mean \pm s.d. for n=3. (d) SNR estimates for determining the limit of detection at 27,000 molecules using purified EGFP through direct and indirect methods (see main text; mean \pm s.d., n = 3 regions of interest per dot blot). Bottom, linear indirect calibration curves for purified standards (mean \pm s.d., n = 3 regions of interest per dot blot) span physiologically relevant βTUB and ERK concentrations (concentration in a probed band estimated from in-cell concentrations, see shaded regions; colors correspond to calibration curves 10,40).



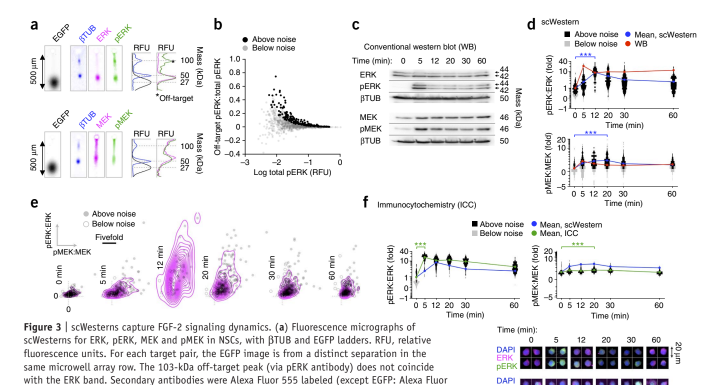
to different blocks. After probing, imaging with a fluorescence microarray scanner yields scWesterns of up to 48 targets per array (3-plex target quantitation for 16 microwell blocks). To further advance multiplexed analyte detection, we adopted serial stripping of antibodies using a strongly denaturing buffer. Using 11 antibody probe sets during nine stripping and reprobing rounds; nine unique targets were successfully detected in the same cell by scWestern blotting and validated by conventional western blotting (Supplementary Fig. 8). Stripping successfully removed antibody probes from scWestern slides, leading to greater-thantenfold reductions in the signal-to-noise ratios (SNRs) compared to initial SNRs (**Fig. 1e** and **Supplementary Fig. 9**). Furthermore, reprobing after the first stripping round led to full recovery of initial probe signals, and signal recoveries of 50% were typical even after nine stripping and reprobing rounds (Supplementary Figs. 8b and 9). Robust signal recovery is likely enabled by stable, covalent protein immobilization, in contrast to the relatively poor recovery observed in conventional platforms that use noncovalent blotting²³. As a result, scWestern slides can be stored for long-term archiving and reanalysis of single-cell separations.

Quantitative performance and calibration of scWesterns

We sought to assay cellular signaling and differentiation in stem cells, which often exhibit diverse behaviors in response to

homogeneous stimuli^{1,2}, using scWesterns. Initially we applied the scWestern to NSCs transduced with a retroviral vector encoding EGFP, using 12 blocks of a single slide (**Fig. 2a**). 4,128 separations of a possible 5,040 (82%) passed semiautomated gating on dust particles and gel defects. Additionally, of those, 1,608 separations (39%) came from single cells, on the basis of bright-field microwell occupancy determination (cells per microwell); and the microwell occupancy running average ranged between 0 and 2.1 cells per microwell with a mean of 1.1 cells per microwell (**Fig. 2b**). Automated occupancy scoring was used for all other data sets to identify single-cell-per-microwell separations (Online Methods); and in all cases, large numbers of microwells housed single cells.

Two protein targets, EGFP and β -tubulin (β TUB), were probed on the same scWestern slide, and the resulting probed band intensities were correlated with microwell occupancy (**Fig. 2a,b** and **Supplementary Fig. 10**). We observed a monotonic but nonlinear relationship between total fluorescence of the β TUB band and microwell occupancy (**Fig. 2b**), likely due to cell size–related bias for microwells with more than one cell (**Supplementary Note 4**). The β TUB fluorescence distribution for single-cell separations was well described by a gamma distribution derived from a stochastic kinetic model of transcription and translation in a homogeneous population of dividing cells²⁴. Our scWestern



(b) Ratio of off-target pERK to total fluorescence for 1,117 scWesterns at time points from d and e.
(c) Conventional western blots (20 ng/ml FGF-2), cropped to show regions of interest (full-length blots in **Supplementary Fig. 16**). (d) Fold change of pERK and pMEK relative to total ERK and MEK, respectively, with signals below technical noise indicated. Note arcsinh-transformed scales. Overlay from conventional western blot (WB) densitometry. ***P < 0.001, Mann-Whitney. n = 186, 186, 57, 236, 278 and 208 scWesterns for time points of 0, 5, 12, 20, 30 and 60 min, respectively. (e) Fold change from d with spatial density contours. (f) ICC coprobing for pERK-ERK and pMEK-MEK pairs; Alexa Fluor 555-labeled secondary phospho-antibodies and Alexa Fluor 647-labeled secondary total antibodies. ***P < 0.001, Mann-Whitney. pERK:ERK n = 160, 115, 186, 158, 172 and 197 cells, and pMEK:MEK n = 184, 216, 220, 223, 223 and 270 cells, for time points 0, 5, 12, 20, 30 and 60 min, respectively.

analysis of β TUB thus agrees with gamma-distributed single-cell protein expression profiles reported for fluorescent protein fusion libraries in *Escherichia coli*^{9,25} and mammalian cells²⁶.

488), in order: pERK, ERK and EGFP coprobe, βTUB, pMEK and MEK, with stripping between probings.

When benchmarked against flow cytometry, we observed 19% and 26.7% \pm 1.1% of the NSCs to be EGFP⁺ (i.e., probed band signals above technical noise) by scWestern analysis and flow cytometry, respectively (\pm s.d., n=3 technical replicates; **Fig. 2c**). The dynamic ranges were comparable (**Supplementary Note 5**).

To determine the linearity and sensitivity of scWestern fluorescence readouts, we undertook 'direct' calibration of EGFP and 'indirect' calibration of both EGFP and BTUB, as well as phosphorylated and total levels of the signaling protein ERK (Fig. 2d). Direct calibration correlates the number of purified EGFP molecules in a coverglass-enclosed microwell separation to probe fluorescence after immunoprobing, whereas the indirect method uses a partition-coefficient measurement to infer the number of molecules in a dot-blotted scWestern band (Supplementary Note 6 and Supplementary Figs. 11–13). The calibration results agreed for EGFP (Fig. 2d) and, together with the indirect calibration data, suggested a linear dynamic range of 1.3-2.2 orders of magnitude, from a limit of detection of 45 zeptomoles (27,000 molecules, comparable to fluorescence cytometry detection limits of 10³–10⁷ molecules²⁷). This detection limit matches an ideal noise threshold of 25,000 molecules set by the fluorescence microarray scanner to within 10%, is 45-fold more sensitive than microwestern arrays²⁰ and is 3.2-fold more sensitive than microfluidic western blotting²¹

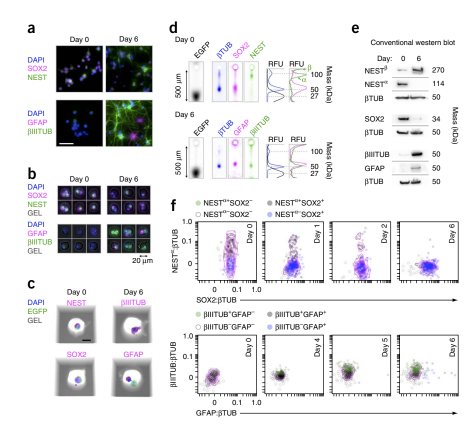
(**Supplementary Note 5**). For context, a median protein copy number of 50,000 has been reported for murine fibroblasts¹⁰, indicating that >50% of the mammalian proteome may be accessible via scWesterns (given the availability of suitable antibodies).

Heterogeneity in signaling after FGF-2 stimulation of NSCs

A defining property of stem cells is self-renewal, or maintenance of an immature state²⁸. Fibroblast growth factor 2 (FGF-2) is a mitogen and self-renewal signal for adult hippocampal NSCs^{28,29}, acting via the FGFR-1 receptor to activate the Ras-MAPK, p38 MAP and PI3K-Akt pathways²⁹. Signals are transmitted in MAPK cascades by sequential phosphorylation of downstream kinases including MEK and ERK. We applied scWesterns to study MAPK signaling dynamics within single NSCs that were starved of FGF-2, suspended, settled into scWestern microwells and stimulated with FGF-2 over a 60-min time course (**Fig. 3**). We first probed for phosphorylated ERK1/2 (pERK) and MEK1/2 (pMEK), after which we reprobed for total ERK1/2 and MEK1/2 (**Fig. 3a** and **Supplementary Fig. 14**). In parallel, β TUB and EGFP probing allowed estimation of molecular mass, and all targets were within 10% of their nominal masses.

scWestern analysis for pERK reported two proteins reactive to the anti-pERK antibody: the expected 38.8 ± 1.0 kDa protein, along with a heavier one of 103 ± 3 kDa (\pm s.d., n=3 separations). We hypothesize that the 103-kDa protein arose from off-target antibody probing because its pERK and ERK antibody fluorescence signals did not correspond (**Fig. 3a**). This off-target band

Figure 4 | scWesterns track NSC lineage commitment during differentiation. (a) ICC micrographs of mixed NSC differentiation cultures at days 0 and 6 for stem cell (nestin, NEST; SOX2) and differentiation markers (BIII-tubulin, BIIITUB; glial fibrillary acidic protein, GFAP). Scale bar, 50 μm. (b) Micrographs of NSCs in scWestern microwells, fixed and stained as in a. (c) Confocal images of fixed and stained stem cells (NEST+SOX2+), neurons (βIIITUB+) and astrocytes (GFAP+) in a rhodamine-tagged gel (GEL). Scale bar, 10 μm. (d) Inverted fluorescence micrographs of scWesterns. SOX2 (Alexa Fluor 555-labeled secondary antibody) and NEST (α and β isoforms; Alexa Fluor 488) were coprobed in separate blocks from GFAP (Alexa Fluor 555) and BIIITUB (Alexa Fluor 488); both block sets were stripped and coprobed for BTUB (Alexa Fluor 555) and EGFP (Alexa Fluor 488). Image sets from each day are the same separation, except EGFP images, which are from the same microwell array row as the corresponding image set. RFU, relative fluorescence units. (e) Cropped conventional western blots at differentiation days 0 and 6 (full-length blots in Supplementary Fig. 25). (f) scWestern fluorescence normalized by βTUB (arbitrary units). Note arcsinh-transformed scales. Spatial density is indicated by contours.



scWestern blot NSC marker sample sizes: n = 189, 353, 175 and 274 at 0, 1, 2 and 6 days, respectively. Differentiation marker sample sizes: n = 178, 253, 303 and 280 at 0, 4, 5 and 6 days, respectively. Data are from one of two biological replicates performed.

may be ERK5 (~80–100 kDa), as ERK5 has high sequence homology with ERK1/2 and because pERK1/2 antibodies have been previously shown to cross-react with pERK5 (ref. 30). The off-target band for pERK exhibited considerable cell-to-cell variability, did not correlate with on-target pERK signal and would have contributed up to 52% (with an average of 13%) of the overall pERK signal in unstimulated cells if not electrophoretically resolved from specific signal (**Fig. 3b** and **Supplementary Fig. 15**). Off-target probe binding can substantially influence existing single-cell immunofluorescence assays (such as ICC and flow cytometry) unless complex target-specific knockdown experiments are performed^{16,17}. In contrast, scWestern analysis is intrinsically well suited to identifying and discarding off-target probing signals.

Both scWesterns and conventional western blotting revealed dynamic, transient ERK and MEK phosphorylation responses (**Fig. 3c–e** and **Supplementary Fig. 16**), but the scWestern enabled analysis of statistical differences (**Supplementary Note 7**). Maximal pMEK:MEK phosphorylation levels agreed quantitatively, with ~3.5-fold increases compared to levels at time point 0 (for single-cell data, fold change is relative to a mean fold change of 1 at time point 0). A larger maximum fold change in the pERK: ERK ratio was observed, consistent with signal amplification in the MAPK phosphorylation cascade³¹.

We next compared scWesterns to a conventional single-cell technique: high-throughput ICC (Fig. 3f and Supplementary Figs. 17–20). pERK:ERK responses by ICC were similar to those measured by scWestern and conventional western assays, whereas pMEK:MEK responses by ICC were strongly attenuated with a maximum mean fold change of <2 (Supplementary Note 7). We

attribute the lower apparent response to nonspecific signal from spurious nuclear localization of the pMEK antibody (a common artifact in ICC^{16}), which obscures the subtle phosphorylation dynamics measurable by scWestern analysis.

Unlike conventional western blots, scWesterns quantified highly variable NSC responses to external stimuli. MEK was activated within 5 min in response to FGF-2, followed closely by ERK, as expected (Fig. 3e). However, a considerable spread in the MEK signal was observed, which was strongly amplified into a broad distribution in ERK activation at 12 min, followed by a transient decay in phosphorylation of both enzymes. This propagation of variation from MEK to ERK upon pathway activation is reflected in interquartile ranges of 3.7 and 7.3 fold-change units at 12 min for pMEK:MEK and pERK:ERK, respectively. The pERK:ERK distributions were skewed toward higher phosphorylation levels for the 0- and 60-min time points owing to the presence of rare activated cells (Fig. 3e and Supplementary Fig. 21). This rare activated state may arise from constitutive signaling or transient FGF-independent excursions from baseline phosphorylation states³².

Heterogeneity in NSC differentiation

In addition to self-renewal, a second hallmark of NSCs is multipotent differentiation from an immature state (markers SOX2⁺ and nestin, NEST⁺) into multiple lineages, such as astrocytes (glial fibrillary acidic protein, GFAP⁺) and neurons (β III-tubulin, β IIITUB⁺)¹³. As with many stem cells, exposure of NSCs to uniform culture conditions can drive stochastic differentiation^{13,32}. We applied the scWestern to study immature NSCs and their

differentiation over a 6-d period under mixed differentiation conditions that yielded both astrocytes and neurons (Fig. 4). Every 24 h, differentiating NSCs were settled into microwells (Fig. 4a-c) and analyzed. ICC in scWestern microwells did not suggest cell type or shape bias upon transfer of differentiated cells to microwells (Supplementary Table 1). The scWesterns successfully reported single bands for SOX2 (43.3 \pm 1.9 kDa), β IIITUB (47.2 \pm 0.7 kDa), and GFAP (54.0 \pm 1.0 kDa, \pm s.d., n = 3 separations; Fig. 4d and Supplementary Fig. 22). Each target protein was within 30% of its expected mass (Supplementary Note 8).

NEST is an intermediate filament protein hypothesized to regulate structural dynamics and cytoplasmic trafficking within neural stem and progenitor cells undergoing rapid rounds of division³³. Intermediate filament proteins often undergo regulation by alternative mRNA splicing, producing diverse isoforms that affect cell responses to stress and modulate intracellular signaling³⁴. In agreement with literature reports^{35,36}, NEST exhibited low- and high-molecular mass bands by scWestern analysis that we denote NEST $^{\alpha}$ (95.7 \pm 3.5 kDa, \pm s.d., n = 3 separations) and NEST^{β} (retained near the microwell edge), respectively (Fig. 4d and Supplementary Fig. 22). NEST $^{\beta}$ fully penetrated the scWestern gel for longer separation distances, indicating that this species is not an insoluble form of NEST (Supplementary Fig. 23).

We further scrutinized NEST α and NEST β by conventional western and scWestern analysis with a second antibody (rat-401) against an epitope known to be excised by alternative splicing in a third, 46-kDa, isoform, Nes-S (ref. 37). As for Nes-S, NEST α was not detected by the rat-401 antibody (Supplementary Fig. 24), suggesting that NEST α may be an alternatively spliced (or otherwise truncated) form of NEST $^{\beta}$ distinct from Nes-S. Intriguingly, NEST $^{\beta}$ was present at all time points over the 6-d experiment, whereas NEST α was variably expressed between cells and sharply downregulated during differentiation. Indeed, contributions of NEST $^{\beta}$ unrelated to proliferative capacity may account for the apparent promiscuity in NEST expression observed in various mature neural cells by ICC³⁸ (including in our data, Fig. 4a). NEST also exhibited two bands in conventional western blotting (114 and 270 kDa, respectively; Fig. 4e and Supplementary Fig. 25), though the extensive cell-to-cell variation in NEST $^{\alpha}$ expression was not detectable with a conventional western.

Consistent with progressive conversion of NSCs to differentiated lineages, conventional western blotting confirmed greaterthan-tenfold reductions in NEST $^{\alpha}$ (but not NEST $^{\beta}$) and SOX2, with accompanying greater-than-tenfold increases in BIIITUB and GFAP over the 6-d period (Fig. 4e). Likewise, both culture-plate and in-microwell ICC showed a corresponding reduction in total NEST+ NSCs, from ~90% to 40% of all cells. Similar overall trends were observed for scWestern data, normalized to constitutively expressed BTUB (Fig. 4f and Supplementary Fig. 26). On day 6, scWesterns put the fractions of committed βIIITUB+ neurons and GFAP+ astrocytes at 53% and 7.1%, respectively, matching ICC to within 15% (Supplementary Table 1).

Notably, scWesterns revealed high cell-to-cell marker expression variability, including profound increases in GFAP expression in the relatively rare astrocyte population over the course of differentiation, spanning a range of 46-fold on day 6. Single-cell expression levels of NEST $^{\alpha}$ at day 0 spanned a range of 22-fold relative to its technical-noise threshold, and the proportion

of cells expressing NEST^α dropped from 53% to 2% between days 0 and 6.

Additionally, scWestern blots successfully resolved off-target antibody signal of approximately equal magnitude to specific signal for MASH1 (ASCL1, a 34-kDa transcription factor involved in neuronal fate commitment) in late-passage, undifferentiated NSCs, as corroborated by conventional western blotting (Supplementary Fig. 27). These data confirm the ability of scWestern assays to accurately capture population expression dynamics by combining the single-cell capabilities of ICC with the molecular mass specificity of conventional western blotting.

DISCUSSION

scWesterns are a single-cell protein analysis technique capable of quantitative, multiplexed and at-the-bench operation, offering an avenue to advance our understanding of cell-to-cell variation in protein-mediated cell functions. Given the often mediocre performance of antibodies as probes^{16,39}, advances in assay specificity are necessary to discriminate between legitimate and spurious protein signals. Western blotting offers high protein specificity, as the technique reports both target molecular mass (via protein electrophoresis) and probe binding (via subsequent antibody probing), not simply probe binding¹⁸. scWesterns bring this specificity to the analysis of single cells, pointing toward a rich, graded heterogeneity in stem cell signaling trajectories. Furthermore, by reporting molecular mass as well as antibody binding, the scWestern could, we showed, identify two putative nestin isoforms and suggest that one (NEST α) better reflects the exit of NSCs from the immature state. In contrast, antibody-binding assays (ICC or flow cytometry) struggle to distinguish such isoforms. Clonal lineage tracing—aided by scWestern analyses—may enable further mechanistic insights into the functions of NEST isoforms²⁶.

More broadly, we envision a role for scWesterns in applications that integrate upstream functional or morphological screens, quantify cell-to-cell response to pharmaceutical agents (including rare circulating tumor cells⁵) and advance affinity-reagent performance by easing library screens.

METHODS

Methods and any associated references are available in the online version of the paper.

Note: Any Supplementary Information and Source Data files are available in the online version of the paper.

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AUTHOR CONTRIBUTIONS

A.J.H. and A.E.H. designed the scWestern. A.J.H., A.E.H., D.P.S. and D.V.S. designed experiments. A.J.H., Z.X. and C.-C.K. performed scWesterns

and calibration experiments. D.P.S. performed NSC culture, stimulation and differentiation; conventional western blots; flow cytometry; and plate-based immunocytochemistry. A.J.H. and D.P.S. performed in-microwell ICC. A.J.H., D.P.S. and Z.X. performed confocal microscopy. Z.X. and A.J.H. designed software and performed analysis for cell/microwell scoring, immunoprobing quality control and fluorescence quantitation. Z.X. performed fluid dynamics simulations. Validation data were analyzed by A.J.H., D.P.S. and Z.X. All authors wrote the manuscript.

COMPETING FINANCIAL INTERESTS

The authors declare competing financial interests: details are available in the online version of the paper.

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ONLINE METHODS

Cell culture. Neural stem cells (NSCs) were isolated from the hippocampi of adult female Fisher 344 rats 28 and cultured on tissue culture-treated polystyrene plates coated with $10~\mu g/ml$ polyornithine (P3655, Sigma-Aldrich) and $5~\mu g/ml$ laminin (23017-015, Life Technologies). NSCs were found to be Mycoplasma free within 12 months both before and following use in experiments. NSCs were cultured in 1:1 DMEM-F12 (11039-021, Life Technologies) supplemented with N-2 (17502-048, Life Technologies) and 20 ng/ml recombinant human FGF-2 (100-18, PeproTech) and were subcultured at 80% confluency using Accutase (A11105-01, Life Technologies) for cell detachment.

EGFP-expressing NSC cell lines were created through stable retroviral infection. The retroviral vector pCLPIT-GFP⁴¹ was packaged⁴², and purified virus was titered on NPCs. High-expressing EGFP NSCs were infected at a multiplicity of infection of 3 (MOI = 3) and analyzed in **Figure 2**, whereas low-expressing EGFP NSCs were infected at MOI = 0.5 and used in all other studies. Stable cell lines were obtained through selection in medium containing 0.3 μ g/ml puromycin for 72 h (P8833, Sigma-Aldrich).

EGFP-expressing NSCs for scWestern EGFP expression studies were cultured as described for uninfected NSCs. For scWestern signaling studies, EGFP-expressing NSCs were FGF-2 starved for 16 h. Cells were detached with Accutase, and suspensions analyzed by scWesterns (see the section "scWestern" below). EGFP-expressing NSCs for scWestern differentiation studies were cultured in DMEM-F12-N2 supplemented with 0.5 ng/ml FGF-2, 1 µM retinoic acid (RA, BML-GR100, Enzo Life Sciences) and 1% FBS (FBS, SH3008803, ThermoFisher Scientific) for 0–6 d. Cells were detached with trypsin-EDTA after the desired differentiation time (25-053-Cl, Corning Cellgro) and analyzed (N.B., cells were not differentiated within microwells; see "scWestern").

Proteins and reagents. 15- μ m fluorescent polystyrene microspheres were from Life Technologies (F-8844). Alexa Fluor 488–labeled purified ovalbumin and bovine serum albumin were also from Life Technologies (O34781, A13100). Purified standards for scWestern calibration were β -tubulin from bovine brain (TL238, Cytoskeleton); recombinant EGFP, His tagged (4999-100, BioVision); and recombinant human pERK1 (ab116536, Abcam). Aliquots of these purified standards were labeled with Alexa Fluor 568 using a protein labeling kit according to vendor instructions (A-10238, Life Technologies) for the determination of partition coefficients in indirect calibration experiments (see "scWestern calibration").

Purified His-tagged Dronpa was expressed in Rosetta competent cells transformed using a pET His6 to bacco etch virus (TEV) ligase independent cloning (LIC) vector, 2BT (EMD Millipore), grown in 2YT medium at 37 °C to an $\rm OD_{600}$ of 0.5, induced with 0.5 mM IPTG and grown for an additional 2.5 h at 37 °C before harvesting. Cells were pelleted by centrifugation for 15 min at 4 °C, and the pellets were resuspended in Nickel buffer A supplemented with protease inhibitors (25 mM HEPES, pH 7.5, 400 mM NaCl, 10% glycerol, 20 mM imidazole, 1 µg/ml leupeptin and pepstatin, 0.5 mM PMSF). Cells were lysed using an Avestin C3 homogenizer at a pressure of 15,000 p.s.i. Cell debris was pelleted at 15,000 r.p.m. for 30 min. The clarified lysate was loaded onto a 5-ml HisTrap FF Crude column (GE Healthcare), and unbound material was washed out with Nickel buffer A. Bound protein was eluted with a 10CV gradient up to 400 mM imidazole in Nickel buffer A. Absorption of the eluting material was monitored at 503 nm as well as at 280 nm to aid in pooling the target protein. Fractions containing Dronpa were pooled and desalted into IEX buffer A (50 mM sodium phosphate, pH 6.5). Desalted protein was loaded onto a 5-ml SP HP ion-exchange column (GE Healthcare), and unbound material was washed out with IEX buffer A. Bound material was eluted with a 20CV gradient up to 1 M NaCl in IEX buffer A. Fractions containing Dronpa were pooled and assayed for aggregation by analytical size-exclusion chromatography on a Superdex 200 5/150 column (GE Healthcare) equilibrated in 25 mM HEPES, 400 mM NaCl, 10% glycerol, 1 mM DTT. Samples were finally desalted into storage buffer (50 mM sodium phosphate, pH 6.5, 150 mM NaCl, 10% glycerol, 1 mM DTT).

N-[3-[(4-benzoylphenyl)formamido]propyl] methacrylamide (BPMAC) was synthesized in-house via the reaction of the succinimidyl ester of 4-benzoylbenzoic acid with N-(3-aminopropyl)methacrylamide hydrochloride in the presence of catalytic triethylamine according to standard protocols^{21,43}.

Fabrication of microwell scWestern substrates. SU-8 microposts were fabricated on mechanical-grade silicon wafers by standard soft lithography methods. SU-8 2025 photoresist (Y111069, MicroChem) was spun to a thickness of (typically) 30 µm according to manufacturer guidelines and exposed to 365-nm UV light at ${\sim}40~\text{mW/cm}^2$ for 12 s under a Mylar mask printed with 20- μm circular features at 20,000 d.p.i. (CAD/Art Services). The features were arranged in a square configuration with a pitch of 500 μm in the direction of separations and 190 µm in the transverse direction (a pitch of 700 μ m yielded separation lengths sufficient for NEST^{β} to fully enter the scWestern gel). 2×8 blocks of 14×30 features (6,720 total) were spaced 9 mm apart to match the dimensions of a 2×8 -well microarray hybridization cassette (AHC1X16, ArrayIt). 1-mm-thick rails spanning the length of the micropost array at a spacing of 24 mm were also patterned to support glass substrates at the height of the microposts. Uniformity of features was verified by optical profilometry after exposure and development using SU-8 developer solution (Y020100, MicroChem). The measured feature heights and diameters within a micropost block were $30.30 \pm 0.15 \,\mu\text{m}$ (±s.d., n = 4 microposts) and $20.52 \pm 0.68 \,\mu\text{m}$ (\pm s.d., n = 4 microposts) for respective nominal dimensions of 30 µm and 20 µm. Between-block CVs in the height and diameter measurements for blocks spaced across the full length of the array were 1.1% and 5.2%, respectively (n = 3 microposts). Wafers were silanized by vapor-deposition of 2 ml of the hydrophobic silane dichlorodimethylsilane (DCDMS, 440272, Sigma-Aldrich) for 1 h in vacuo, washed thoroughly with deionized (DI) water, and dried under a nitrogen stream immediately before use. Silanized wafers were robust to reuse after rinsing with DI water in excess of 20 times with moderate delamination of micropost structures.

Plain glass microscope slides (48300-047, VWR) were silanized to establish a self-assembled surface monolayer of methacrylate functional groups according to standard protocols⁴⁴. Silanized slides were placed facedown onto micropost wafers and manually aligned to the SU-8 rail and micropost features. Gel precursor solutions were 8%T (w/v total acrylamides), 2.7%C (w/w of the cross-linker *N*,*N*-methylenebisacrylamide) from a 30%T, 2.7%C stock (A3699, Sigma-Aldrich); 3 mM BPMAC from a

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100 mM stock in DMSO, 0.1% SDS (161-0301, Bio-Rad), 0.1% Triton X-100 (BP151, Fisher), 0.0006% riboflavin 5' phosphate (F1392, Sigma-Aldrich), 0.015% ammonium persulfate (APS, A3678, Sigma-Aldrich), and 0.05% tetramethylethylenediamine (TEMED, T9281, Sigma-Aldrich) in 75 mM Tris buffer titrated with HCl to a pH of 8.8. For confocal imaging of cells in rhodamine-tagged scWestern gels, the precursor included the fluorescent monomer methacryloxyethyl thiocarbamoyl rhodamine B (23591, Polysciences) at 3 µM from a 100 µM stock in DMSO. The precursor mixture was sonicated and degassed (Aquasonic 50D, VWR) for 1 min in vacuo immediately before the addition of detergents (SDS, Triton) and polymerization initiators (riboflavin, APS, TEMED). The precursor was then injected into the gap between the glass slide and silicon wafer using a standard 200-µl pipette. After allowing ~30 s for precursor to wick through the gap, the slide was exposed to blue light for 7.5 min at 470 lx (advanced light meter, 840022, Sper Scientific) from a collimated 470-nm LED (M470L2-C1, Thor Labs) mounted at a 45° angle above the slide. Polymerization was allowed to continue for an additional 11 min. Gel-fabricated glass slides were wetted at their edges using 2 ml of phosphate-buffered saline (PBS), pH 7.4, (21-040, Corning) and carefully levered from wafers using a razor blade. Fabricated slides could be stored at 4 °C in PBS for up to 2 weeks before use without loss of protein separation or photocapture properties.

scWestern. Fabricated slides were removed from PBS, and excess liquid was drained to a corner by gravity and absorbed using a Kimwipe (Kimberly-Clark). 1-2 ml of cell suspension was applied evenly across the surface of the slide and allowed to settle on a flat surface within a 100 mm × 100 mm Petri dish. Settling times varied from 5 to 30 min, with microwell occupancy monitored by bright-field microscopy until single-cell occupancies of roughly 40-50% were achieved. Intermittent, gentle movement of the Petri dish every 2-5 min for 10 s was sufficient to ensure cell access to microwells through cell rolling on the gel surface. After settling, slides were lifted to a 10-20° angle from one of the short edges to remove excess cell medium, and cells on the surface of the slide were removed by gentle pipetting of 4 or 5 1-ml aliquots of PBS to the raised edge of the slide surface, with excess buffer removed from the lower edge by vacuum. Slides were placed flat and prepared for cell counting by applying 1 ml of PBS onto the slide. A second plain glass slide was applied to the PBS layer from one short edge to the other to prevent entrapment of bubbles and lowered to form a 'sandwich' of slides. Microwells within the sandwich were imaged by bright-field microscopy at 4× magnification (Olympus UPlanFLN; numerical aperture (NA), 0.13) using 50-ms exposure times at 1×1 -pixel binning and a preset position list to guide a mechanical stage (Olympus IX71 inverted fluorescence microscope equipped with iXon+ EMCCD camera, Andor; motorized stage, ASI; and shuttered mercury lamp light source, X-cite, Lumen Dynamics; controlled by MetaMorph software, Molecular Devices). All 6,720 features could be imaged in ~4 min.

After cell counting, the top glass slide was removed from the sandwich by sliding gently across the gel layer. The scWestern slide with settled cells was then immediately transferred to a custom $60~\text{mm} \times 100~\text{mm}$ horizontal electrophoresis chamber fabricated from 3-mm-thick Perspex plastic. Platinum wire

electrodes (0.5-mm diameter, 267228, Sigma-Aldrich) were placed along the long edge of the chamber and interfaced with alligator clips to a standard electrophoresis power supply (Model 250/2.5, Bio-Rad). Slides were temporarily adhered to the bottom face of the chamber using petroleum jelly. 10 ml of a denaturing RIPA lysis/electrophoresis buffer consisting of 0.5% SDS, 0.1% v/v Triton X-100, 0.25% sodium deoxycholate (D6750, Sigma-Aldrich) in 12.5 mM Tris, 96 mM glycine, pH 8.3, (0.5× from a 10× stock, 161-0734, Bio-Rad) was poured over the slide to lyse cells. This buffer was supplemented with 1 mM sodium fluoride and sodium orthovanadate for phosphoprotein analyses (S7920 and 450243, Sigma-Aldrich). The RIPA buffer provides denaturing but nonreducing conditions, as reduction typically requires heating in the presence of a reducing agent for time scales longer than that of protein diffusion from microwells²¹. Lysis proceeded for 10 s with the electric field off, followed by application of 200 V (E =40 V/cm) for ~30 s. Separations from single EGFP-expressing NSCs were monitored in real time at 10× magnification (UPlanFLN; 0.3-NA objective) using a filter set optimized for EGFP (XF100-3, Omega Optical), 4×4 camera binning, 250-ms exposure time. Following separations, slides were immediately exposed for 45 s from above using a UV mercury arc lamp (Lightningcure LC5, Hamamatsu) directed through a Lumatec series 380 liquid light guide with inline UV filter (300- to 380-nm bandpass, XF1001, Omega Optical) suspended approximately 10 cm above the slide with UV power at the slide surface of ~40 mW/cm² (320-400 nm UV meter; C6080-365, Hamamatsu).

Following separation and photocapture of cell contents, slides were washed using 10 ml of the denaturing RIPA buffer, followed by 10 ml of TBST (100 mM Tris titrated to pH 7.5 with HCl, 150 mM NaCl, 0.1% Tween 20, 9480, EMD Millipore), each for 10 min. Slides could be stored before successful immunoprobing for at least 1 week at 4 °C in TBST.

In FGF-2 stimulation experiments, cells were stimulated between cell-per-microwell counting and lysis/electrophoresis steps by applying 1 ml of 20 ng/ml FGF-2-spiked culture medium to the slide surface for the desired stimulation time.

Purified-protein scWesterns. Purified proteins were assayed using a similar protocol to that for single cells. Gel slides were incubated with purified proteins in denaturing RIPA buffer for 30 min, submerged in fresh denaturing RIPA for 5 s and 'sandwiched' with a second glass slide to trap proteins within the gel layer. The glass slide sandwich was subjected to electrophoresis, UV-mediated protein capture, washing and probing as in single-cell assays; the top glass layer was removed after the capture step.

scWestern probing, imaging, and stripping. Slides were probed with primary and fluorescently labeled secondary antibodies by diffusive delivery in 2×8 -well microarray hybridization cassettes (AHC1X16, ArrayIt).

Primary antibodies with fold dilutions employed for single-cell, purified protein, and calibration assays (unless otherwise noted) were rabbit anti-ovalbumin (1:20, ab1221, Abcam), goat anti-GFP (1:20, ab6673, Abcam), rabbit anti- β -tubulin (1:20, ab6046, Abcam), rabbit anti-pERK1/2 (1:40, Thr202/Tyr204, 4370, Cell Signaling), rabbit anti-ERK1/2 (1:20, 4695, Cell Signaling), mouse anti-ERK1/2 ("ERK #2," 1:20, 4696, Cell Signaling), rabbit

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anti-pMEK1/2 (1:40, Ser217/Ser221, 9154, Cell Signaling), rabbit anti-MEK1/2 (1:20, 9126, Cell Signaling), goat anti-SOX2 (1:20, sc-17320, Santa Cruz Biotechnology), mouse anti-nestin ("NEST," 1:20, 611658, BD Biosciences), mouse anti-nestin ("NEST #2," 1:20, MAB353, clone: rat-401, EMD Millipore), goat anti-EphB4 (1:20, AF446, R&D Systems), mouse anti-MASH1 (1:20, 556604, BD Biosciences), mouse anti-SRC (1:20, 05-184, EMD Millipore), goat anti-GFAP (1:20, ab53554, Abcam), mouse anti-βIII-tubulin (1:20, T8578, Sigma-Aldrich). Secondary antibodies were Alexa Fluor 488-, 555-, or 647-labeled donkey anti-mouse, rabbit or goat IgG from Life Technologies (A31571, A31573, A21447, A31570, A31572, A21432, A21202, A21206, A11055), except for the probing of ovalbumin in Supplementary Figure 7, which used Alexa Fluor 568-labeled goat anti-rabbit IgG (A-11011, Life Technologies). All were used at the same dilution factor as that of the corresponding primary antibody.

Each block of separations was incubated at room temperature with 40 µl of primary antibody solution diluted in TBST supplemented with 2% bovine serum albumin (BSA, A730, Sigma-Aldrich) for 1 h under gentle orbital shaking. Slides were removed from hybridization cassettes and washed three times in 10 ml TBST for 15 min per wash (45 min total), also under gentle orbital shaking. Slides were then similarly probed and washed with fluorescently labeled secondary antibodies in TBST supplemented with 2% BSA. Slides were washed a final time in 10 ml DI water for 5 min and dried under a nitrogen stream. Imaging was conducted using a GenePix 4300A microarray scanner with PMT gains of 400–550 and laser powers of 30-100%, optimized for maximum dynamic range without saturation of target band fluorescence values. Filter sets were employed for three-channel detection using Alexa Fluor 488-, 555- and 647-labeled secondary antibodies using 488-, 532- and 635-nm lasers, respectively. 12.5-mm-diameter emission filters for the 488- and 532-nm spectral channels were from Omega Optical (XF3405 and XF3403, respectively); the 635-nm channel employed a built-in far-red emission filter.

Spectral bleed-through was below noise thresholds of on-target fluorescence line profiles, except for coprobing of ERK or β -tubulin (Alexa Fluor 555–labeled secondary antibody) with EGFP (Alexa Fluor 488–labeled secondary antibody) in **Figures 3** and **4**, respectively. Ratio metrics in **Figure 3d,e** for which ERK bands were affected by EGFP bleed-through above technical noise were discarded from analysis. Ratio metrics in **Figure 4f** derived from β -tubulin bands similarly affected by EGFP bleed-through were also discarded. No fluorescence micrographs or derived data sets were fluorescence compensated for spectral bleed-through.

Stripping of slides was performed via 3-h incubations in a stripping buffer heated to 50 °C consisting of 2.5% SDS and 1% $\beta\text{-mercaptoethanol}$ (M3148, Sigma-Aldrich) in 62.5 mM Tris titrated to pH 6.8 with HCl. Following stripping, slides were washed three times in 10-ml aliquots of TBST for 5 min per wash and stored in TBST at 4 °C until reprobing. For longer-term archiving, stripped, air-dried slides could be successfully reprobed after extended (>1 month) storage at 4 °C.

scWestern data analysis. Cell-per-microwell scoring was conducted manually or via custom software designed in-house that employed scripts to mate thresholding and particle analysis on the basis of cell size and circularity in ImageJ (http://rsbweb.nih.gov/ij/) to downstream gating to identify microwells containing

single cells in R (http://www.r-project.org/) (Supplementary Software).

To quantify the performance of automated cell-per-microwell scoring, we calculated precision = tp/(tp + fp) and sensitivity = tp/(tp + fn), where tp is the number of microwells scored as containing single cells that actually contained single cells, fp is the number of microwells scored as containing single cells that did not contain single cells and fn is the number of microwells scored as not containing single cells that actually contained single cells⁴⁵. Precision = 1 means that all microwells scored as containing single cells actually contained single cells. Sensitivity = 1 means that all microwells actually containing single cells were scored as containing single cells. Precision and sensitivity metrics were 0.90 ± 0.09 and 0.68 ± 0.17 , respectively ($\pm s.d.$, n = 56 blocks of 420 microwells on 8 separate slides), reflecting stringent selection of single-cell microwells at the expense of the total number of microwells included in downstream analysis.

Fluorescence images from the GenePix scanner were registered using landmark correspondences in Fiji (http://fiji.sc/Fiji). A custom script extracted line profiles from grids of regions of interest (ROIs) from each fluorescence image. Line profiles were background subtracted using linear interpolation between points set to the approximate boundaries of peaks of interest. Data quality control was performed by manually reviewing separation ROIs flagged owing to outlying line profiles. Separations that were clearly affected by the presence of, for example, autofluorescent particulates were discarded from data sets, as were zero-cells-per-microwell separations incorrectly scored as single-cell separations that did not contain β -tubulin loading control signals above technical noise.

Total areas under peaks (AUCs) of interest (or metrics derived from them, such as AUC ratios and calibrated AUCs) were transformed, where applicable, using the function $AUC_t =$ arcsinh(AUC/F), where AUC_t is the arcsinh-transformed value and *F* is a cofactor prescribing the transition from linear to loglike behavior. The value of *F* was optimized by setting it according to $F = \mu_{\text{ones,below}} + 3\sigma_{\text{ones,below}}$ where $\mu_{\text{ones,below}}$ and $\sigma_{\text{ones,below}}$ are the mean and s.d. of the set of single-cell-per-microwell separations with fluorescence AUCs (or metrics) below a technical noise threshold. The technical noise threshold *T* was set at $T = \mu_{zeros}$ + $3\sigma_{zeros}$, where μ_{zeros} and σ_{zeros} are the mean and s.d. of the AUCs or metric values from zero-cells-per-microwell separations in a given experiment. Where applicable, separations with AUCs in the numerator of ratio metrics falling below T were flagged to display as such when plotted. Separations with AUCs below T in the denominator were discarded from data sets.

For analysis of β TUB distribution in **Figure 2b**, histogram data were fit to the gamma distribution $f(x) = (x^{a-1}e^{-x/b})/(\Gamma(a)b^a)$ using an implementation of the least-squares Marquardt-Levenberg algorithm in gnuplot (http://www.gnuplot.info); x = total probed band fluorescence, $a = \mu_p^2/\sigma_p^2 = 14.8$, $b = \sigma_p^2/\mu_p = 1.6 \times 10^5$ AFU, $\mu_p =$ mean band fluorescence, $\sigma_p^2 =$ variance in band fluorescence and Γ is the gamma function.

Statistical analysis. Nonparametric comparison of scWestern data (single comparisons only) was performed using the Mann-Whitney *U*-test in conjunction with Shapiro-Wilk and Levine tests for normality and equality of variance, respectively, in SPSS v.21 software (IBM).

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scWestern calibration. A conceptual overview and schematics of 'direct' and 'indirect' calibration assays are provided in **Supplementary Note 6** and **Supplementary Figure 11**. For direct calibration of EGFP, an eight-aliquot dilution series (40 μ l per aliquot) of EGFP in denaturing RIPA buffer supplemented with $4\,\mu\text{M}$ BSA (approximating total protein levels in single-cell separations) was added to distinct microwells of scWestern slides in the ArrayIt hybridization cassette (Supplementary Fig. 12). Slides were sandwiched and assayed as for purified protein assays (see "Purified protein scWesterns") with one additional step. A subset of microwells in each block were imaged for EGFP fluorescence (10× magnification, 200-ms exposure time, 1×1 -pixel binning) immediately before the electrophoresis step using a preset position list to guide the mechanical stage on the IX71 fluorescence microscope. Partition coefficients across the concentration range were determined from these images according to $K = ([EGFP]_{gel} -$ [EGFP]_{gel,bg})/([EGFP]_{microwell} - [EGFP]_{microwell,bg}), where [EGFP]_{gel} and [EGFP]_{microwell} are in-gel and in-microwell concentrations of EGFP at equilibrium determined by fluorescence calibration in a separate microfluidic channel of 30-µm depth (Supplementary Fig. 4). Custom straight-channel microfluidic chips were fabricated in soda lime glass using standard wet-etching processes (PerkinElmer). [EGFP]_{gel,bg} and [EGFP]_{microwell,bg} correct for the background fluorescence of the scWestern slide before incubation with the EGFP solutions. The number of molecules of EGFP in each microwell voxel was also estimated from these data, assuming cylindrical microwells of nominal dimensions: 20-µm diameter, 30-µm depth (9.4-pl volume).

Indirect calibration was performed by capturing to the scWestern gel and probing a dilution series of a given purified protein in denaturing RIPA supplemented with 4 µM BSA in the absence of an electrophoresis step (Supplementary Figs. 11 and 12). Spot UV exposures were applied to the underside of the slide within each microwell block via the 10× objective for 45 s each on the Olympus IX71 fluorescence microscope through a custom UV-longpass filter set (excitation 300-380 nm, emission >410 nm; XF1001, XF3097; Omega Optical) with a UV power at the slide surface of ~40 mW/cm² (320- to 400-nm UV meter; C6080-365, Hamamatsu). The in-gel concentrations of purified proteins captured in this manner were determined from separate partition coefficient measurements using Alexa Fluor 568-labeled aliquots of each protein (Supplementary Fig. 4). Indirect calibration of EGFP reports molecule number using the inferred in-gel concentrations for a voxel size equivalent to that of a typical probed EGFP band from an scWestern experiment (45 μ m \times 45 μm in area, 30 μm in depth). Probe AFU and SNR values in indirect calibration data were corrected for fluorescence background caused by nonspecific probing of UV-exposed gel spots in the absence of calibration standard.

Integration of scWestern analysis with FACS. Live EGFP-expressing NSCs were sorted using an Influx v7 Sorter (BD Biosciences). BD FACS Sortware 1.0.0.650 was used to establish a 4×16 grid over the surface of a dried scWestern slide for deposition of sorted cells. 10- μ m polystyrene fluorescent microspheres (Flow-Check Fluorospheres, 6605359, Beckman Coulter) were test sorted for fine positioning adjustment. Cells were gated for EGFP expression, and sorting was calibrated so that each droplet exiting the nozzle contained a single EGFP-positive cell or no cells.

The sorting purity was ~96%. After FACS, gel slides were rehydrated by immersion in PBS and analyzed by scWestern. For propidium iodide (PI) cell staining, PI (1 mg/ml, P3566, Life Technologies) was added to cell suspensions at 1:100 dilution. Dead cells were imaged after drying of FACS droplets on scWestern slides by fluorescence microscopy.

Determination of bulk buffer velocity during in-microwell lysis. Bulk maximum flow speeds during lysis (ignoring vector information) were estimated by wide-field fluorescence microscopy (4× objective, EGFP filter set) during pouring of a 15-µm fluorescent microsphere–spiked RIPA buffer over a scWestern slide (10⁵ microspheres/ml) at an exposure time of 10 ms (**Supplementary Fig. 3**). Velocities were extracted from fluorescence streaks caused by movement of microspheres in the horizontal plane over the exposure period, with the objective focused ~1 mm above the center of the scWestern slide plane to observe bulk fluid behavior.

COMSOL fluid modeling. Fluid flow in scWestern microwells was modeled in COMSOL Multiphysics 4.2a (Supplementary Note 1 and Supplementary Fig. 3). Bulk flow above microwells was simulated as steady-state laminar flow of water in a square channel of cross-section $100\,\mu\text{m}\times100\,\mu\text{m}$. The top and side walls of the channel were set to a slip boundary condition. The bottom wall of the channel and the microwell walls were set to no-slip. Inlet velocity was set to 0.0087 m/s to achieve a maximum bulk flow velocity of 0.013 m/s. Outlet pressure was set to 0. Microwell recirculation flow was visualized by the particle tracing module in COMSOL.

Flow cytometry. For flow cytometry for EGFP expression; EGFP-expressing NSCs and uninfected NSCs were detached with Accutase, fixed by suspension in 4% paraformaldehyde (P6148, Sigma-Aldrich) for 15 min and then blocked and permeabilized with flow staining buffer (5% donkey serum with 1 mg/ml saponin; D9663 and 47036, Sigma-Aldrich; in PBS) for 15 min. Cells were incubated with goat anti-GFP (1:100; see "scWestern probing, imaging, and stripping" for product details) in flow staining buffer for 1 h; followed by incubation with Alexa Fluor 555–labeled donkey antigoat IgG (1:100) in flow staining buffer for 1 h. Cells were washed twice for 5 min each with staining buffer between application of primary and secondary antibodies, and finally for 5 min with staining buffer and twice for 5 min each with PBS immediately before performing flow analysis. Flow cytometry was performed using an EMD Millipore EasyCyte 6HT-2L.

Conventional western blotting. For the signaling study in **Figure 3c**, EGFP-expressing NSCs were seeded at 2.5×10^5 cells per well in a six-well culture plate. Cells were FGF-2 starved for 16 h, incubated with 20 ng/ml FGF-2 for the desired stimulation time and lysed in RIPA buffer (50 mM Tris, 150 mM NaCl, 1% NP-40, 0.5% sodium deoxycholate, 0.1% SDS, pH 8) containing protease and phosphatase inhibitor cocktails (87786 and 78420, ThermoFisher Scientific) and 10 mg/ml PMSF (78830, Sigma-Aldrich). For the differentiation assay in **Figure 4e**, EGFP-expressing NSCs were seeded at 5×10^5 cells per dish in 6-cm dishes. Day 0 differentiated cells were lysed the following day; day 6 differentiated cells were cultured in differentiation medium (DMEM-F12-N2, 0.5 ng/ml

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FGF-2, 1 µM RA, 1% FBS) for 6 d and then lysed. Cell lysates of equal total protein concentrations determined by a bicinchoninic acid assay (23227, ThermoFisher Scientific) were mixed with 5× Laemmli buffer (final 50 mM Tris, 2% SDS, 0.1% bromophenol blue, 10% glycerol), 2-mercaptoethanol was added to 10% v/v and samples were boiled at 95 °C for 5 min. Samples were electrophoretically separated on SDS-PAGE gels of between 6 and 10%T and transferred onto nitrocellulose membranes using standard methods. Blots were blocked for 1 h in TBST and 3% BSA for phosphoprotein antibodies or 5% nonfat powdered milk (6250, EMD Millipore) for all other antibodies. Blots were probed overnight with primary antibodies in the same blocking buffer: rabbit anti-pERK1/2 (1:2,000; see "scWestern probing, imaging, and stripping" for product details), rabbit anti-ERK1/2 (1:1,000), mouse anti-ERK1/2 ("ERK #2," 1:1,000), mouse anti-MASH1 (1:1,000), mouse anti-SRC (1:1,000), goat anti-EphB4 (1:1,000), goat anti-GFP (1:1,000), rabbit anti-pMEK1/2 (1:1,000), rabbit anti-MEK1/2 (1:1,000), goat anti-SOX2 (1:500), mouse anti-nestin (1:1,000), mouse anti-nestin ("NEST #2," clone: rat-401, 1:2,000), goat anti-GFAP (1:1,000), mouse anti-βIII-tubulin (1:2,000), rabbit anti- β -tubulin (1:500); followed by 1 h incubation with appropriate horseradish peroxidase-conjugated secondary antibodies: mouse anti-goat HRP (1:5,000, 31400), goat anti-mouse HRP (1:10,000, 32430) and goat anti-rabbit HRP (1:10,000, 32460), all from ThermoFisher Scientific. Protein bands were detected using SuperSignal West Dura Chemiluminescent Substrate (34076, ThermoFisher Scientific), and blots were digitally imaged on a ChemiDoc XRS+ Imaging System (Bio-Rad). Blots were stripped in a solution of 3% acetic acid, 0.5 M NaCl, pH 2.5, for 10 min, neutralized with 0.5 M NaOH for 1 min, and then reprobed as needed. Blot densitometry was performed in ImageJ by measuring background-subtracted ROI intensities.

For purified protein samples (**Supplementary Fig. 6**), 1 μ g of Alexa Fluor 488–labeled OVA and/or 1 μ g Alexa Fluor 488–labeled BSA was incubated in denaturing or standard RIPA buffer for 30 min at room temperature, protected from light. Samples were then mixed with 5× Laemmli buffer. For reducing conditions, 2-mercaptoethanol was added to samples to 10% v/v. For boiling conditions, samples were boiled at 95 °C for 5 min; nonboiled samples were incubated at room temperature for 5 min. All samples were electrophoretically separated on a 10% SDS-PAGE gel. Fluorescent protein bands were directly imaged in-gel via the ChemiDoc instrument.

Immunocytochemistry. For the signaling study in Figure 3f, EGFP-expressing NSCs were seeded at 5 × 10³ cells per well in a 96-well plate. Cells were FGF-2 starved and stimulated as described for conventional western blotting. For the differentiation assay in standard cell culture conditions (Fig. 4a), EGFP-expressing NSCs were seeded at 4 × 10⁴ cells per well in a 24-well plate and differentiated. For scWestern microwells, EGFP-expressing NSCs were differentiated in culture plates, suspended on the appropriate day, settled into scWestern slides and processed within ArrayIt hybridization cassettes. Cell cultures and settled cells were fixed with 4% paraformaldehyde for 15 min and then blocked and permeabilized with staining buffer (5% donkey serum with 0.3% Triton X-100 in PBS) for 30 min. Cultures and cells were incubated 24–48 h with combinations of primary antibodies in staining buffer: rabbit anti-pERK1/2 (1:200; see "scWestern probing, imaging and

stripping" for product details), mouse anti-ERK1/2 (1:50, 4696, Cell Signaling), rabbit anti-pMEK1/2 (1:200), mouse anti-MEK1/2 (1:25, 4694, Cell Signaling), goat anti-SOX2 (1:100), mouse antinestin (1:200), goat anti-GFAP (1:500), mouse anti-BIII-tubulin (1:500); followed by 2 h incubations with appropriate Cy3–, Alexa Fluor 555–, and Alexa Fluor 647–labeled donkey anti-mouse, rabbit or goat IgG secondary antibodies (1:250, Life Technologies; 15-165-150, 715-605-150, 711-605-152, 705-605-147, Jackson ImmunoResearch), with 4,6-diamidino-2-phenylindole (DAPI) as a nuclear counterstain (5 μ g/ml, D1306, Life Technologies). Cell cultures were imaged using a Nikon Eclipse Ti inverted fluorescence microscope (Nikon Instruments) or an ImageXpress Micro XL Widefield High Content Screening System (Molecular Devices). In-microwell cells were imaged using the Olympus IX71 microscope (see "scWestern").

Confocal images were obtained on a BX51W1 microscope (Olympus) with swept-field confocal optics (Prairie Technologies) and analyzed with Icy bioinformatics software (http://icy.bioimageanalysis.org). For confocal imaging of differentiated cells in scWestern microwells in **Figure 4c**, rabbit anti-GFAP (1:500, ab7260, Abcam) was used; all other antibody reagents were identical to those listed.

Immunocytochemistry data analysis. For the signaling study in Figure 3f, cells were identified via custom ImageJ scripts using thresholding and particle analysis to locate DAPI-stained nuclei. Single cells for analysis were isolated and selected by gating for distance to nearest neighbor cells and uniformity of background signal in R. Fluorescence was quantified by summing pixel intensities of a background-subtracted 75×75 -pixel ROI around each single cell. Approximately 50% of pixels in each ROI consisted of background signal, which was Gaussian in distribution. The intensity value with highest pixel count was taken to be the mean background intensity and used for background subtraction for individual ROIs. A noise threshold was set to $T = 3\sigma_{bg}$, where σ_{bg} is the maximum s.d. of background signal intensity in the fluorescence micrographs at each experimental condition. Measurements with fluorescence below T in the numerator were identified as such in plotted data. Measurements with fluorescence below T in the denominator were discarded from data sets.

Fluorescence micrographs from ICC experiments in culture plates and scWestern microwells for the differentiation experiment in **Figure 4a**,**b** were manually scored for marker expression according to arbitrarily determined fluorescence thresholds in ImageJ. Different, blinded researchers conducted ICC counting and scWestern marker expression analyses.

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