Cross-Tissue Identification of Somatic Stem and Progenitor Cells Using a Single-Cell RNA-Sequencing Derived Gene Signature

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ABSTRACT

A long-standing question in biology is whether multipotent somatic stem and progenitor cells (SSPCs) feature molecular properties that could guide their system-independent identification. Population-based transcriptomic studies have so far not been able to provide a definite answer, given the rarity and heterogeneous nature of these cells. Here, we exploited the resolving power of single-cell RNA-sequencing to develop a computational model that is able to accurately distinguish SSPCs from differentiated cells across tissues. The resulting classifier is based on the combined expression of 23 genes including known players in multipotency, proliferation, and tumorigenesis, as well as novel ones, such as Lcp1 and Vgll4 that we functionally validate in intestinal organoids. We show how this approach enables the identification of stem-like cells in still ambiguous systems such as the pancreas and the epidermis as well as the exploration of lineage commitment hierarchies, thus facilitating the study of biological processes such as cellular differentiation, tissue regeneration, and cancer.

SIGNIFICANCE STATEMENT

This novel transcriptomics-based approach exploits the increased molecular resolution provided by single-cell RNA-sequencing to accurately identify somatic stem and progenitor cells (SSPCs) across a wide range of tissues. The developed classifier uniquely combines expression information of only few genes, suggesting that SSPCs share specific molecular properties across tissues. The method provides a valuable resource to identify SSPCs and order them across differentiation stages in yet poorly characterized systems. Among its downstream applications we envision, besides the detection of novel stem-like cells, the estimation of the fraction of SSPCs in a heterogeneous sample or, in an expanded version, the detection of highly malignant cells in tumors.

INTRODUCTION

The identification and molecular characterization of multipotent somatic stem and progenitor cells (SSPCs) is of fundamental interest for understanding development, homeostasis, and regeneration of complex multicellular organs. In addition, the study of cells with multipotent capacity provides novel regenerative therapy opportunities [1] as well as new insights into disease mechanisms or treatments [2]. While hematopoietic, neural, epidermal, and gastrointestinal stem cells have already been phenotypically well-characterized, it has proven very challenging to identify SSPCs and establish their hierarchy in a broad range of other systems, including the lung, kidney, mesenchyme, heart, liver, and pancreas [3–5]. This is largely because such SSPCs tend to constitute small and heterogeneous populations that reside in tissues of complex composition and lack universal markers, which leads to technological or experimental limitations such as population-level molecular measurements and tedious in vivo functional assays [6].

Recent advances in single-cell genomics have revolutionized our ability to reveal the composition of individual tissues or specific developmental patterns [7–10]. To determine which single-cell RNA-sequencing (scRNA-seq)-assessed cells are stem-like, studies tend to cluster or arrange them along a differentiation time-line (termed “pseudo-time”) and attribute labels to groups based on marker gene expression [8–11]. This is an iterative and highly variable process, as it is sensitive to particular algorithmic (clustering, graph walking etc.) as well as biological (markers, cell stages) choices. The recently developed StemID alleviates these issues by streamlining the procedure, but it is only applicable to cells...
across multiple differentiation stages and not to homogeneous single-cell populations [12]. For such cases, one is obliged to leverage on the knowledge accumulated in other systems to assign functionality. To formalize such an approach, we developed a cross-tissue SSPC identification method.

A wide range of efforts have already been directed toward delineating a stem cell (SC) molecular signature (referred to as “stemness” signature) [13–19]. For example, over 100 stemness-related resources, including curated gene sets, computationally derived signatures, and transcription factor (TF) targets have been collected and made centrally available through the online platform StemChecker [20]. Importantly, while there is substantial overlap between the individual resources, there is by no means a consensus on the molecular program that underlies the core properties of SSPCs [20] or a universally accepted marker gene set that facilitates their identification [6]. This may be because SSPCs may only be captured through a combination of system-specific features rather than single marker genes [4, 16, 21]. At the same time, while scRNA-seq studies revealed cell hierarchies and novel cell types within systems of interest [7–10, 22–24], cross-study analyses have so far only focused on cell cycle stage identification [25]. In this study, we aimed to bridge this gap by exploring how well multipotent SSPCs could be identified and distinguished from differentiated cells based on scRNA-seq data. We demonstrate here that a lasso logistic regression-based model is able to accurately identify various SSPCs by using a unique combination of genes that together reach good cross-system performance.

**Materials and Methods**

In our approach, we carefully considered (a) the quality/diversity of the training data, (b) the genes included as features in the training, (c) the algorithm used, and (d) the normalization applied prior to training, exploring several options in order to obtain a robust model.

**Datasets**

We used the following data, also listed in Supporting Information Table S1: (1) as positive SSPCs, (a) as train and test data, we included quiescent and activated neural stem cells assessed after injury (qNSCs and aNSCs, d1) [10], macrophage dendritic cell progenitors (MDPs, d2) [22], hematopoietic stem cells (HSCs, d3, and d4) [26, 27] and (b) as independent test data quiescent and activated NSCs (qNSCs and aNSCs, d9) [28], mesenchymal stem cells (MSCs, d10) [29], short- and long-term HSCs (stHSCs and lHSCs, d11–12) [30], intestinal stem cells (ISCs, d13) [9]; (2) as negative data (non-SSPCs), (a) as train and test data, we included astrocytes (STR and CTX, d5) [10], T cells (d6) [31], AT2 lung epithelial cells (AT2 cells, d7) [8], and adult non-stem neuronal cells (neuro, d8) [23] and (b) as independent test data non-stem intestinal cells (REG4+ and LGR5+, d14–15) [9], lymphoblastoid cell lines (LCLs, d16) [32], adult non-stem neuronal cells (neuro, d17) [33] as well as adult hepatocytes and endothelial cells isolated from human liver (d33) [34]. To test predictions on cells with a gradually decreasing level of stemness, we used: (a) oligodendrocytes (Oligo, d18), neuroblasts (NB, d19), neural transit amplifying cells (TAPs, d20), activated and quiescent NSCs (aNSCs and qNSCs, d21–22); (b) megakaryocyte progenitor cells (CD150- MkPCs, d23); (c) dendritic cell precursors (PreDCs, d24) and common dendritic cell progenitors (CDPs, d25) [22]; (d) an unsorted mixture of cells progressively losing progenitor status from embryonic day 14 over 16 to 18 (E14, E16, and E18, d26–27) [8]. To keep the fraction of positive and negative cells balanced, we selected only a (random) subset of cells among those that were part of large datasets: d7, d8, and d11–12 [23, 30, 31]. Finally, we used dissociated mouse epidermis cells [35] as well as human pancreas cells [12, 36, 37] for generating novel predictions.

**Data Normalization**

We used the expression estimates provided by the individual studies, which were publicly available through Gene Expression Omnibus [38], irrespective of the underlying experimental and computational methods that they were based on (Supporting Information Table S1). For developing the stem and progenitor cell identification model, due to the highly heterogeneous nature of scRNA-seq, exacerbated by both experimental and data processing methodological differences among the studies as well as variability across distinct biological sources, we did not use these raw expression estimates. Rather, we applied rank-based normalization and reduced the dynamic range of the data by using quantiles instead of ranks or expression estimates.

We ran the lasso logistic regression model described below on a range of ranked data (for non-zero data points: 10, 20, 30, 50, 100, 500, 1,000, and 1,500 quantiles, with zero as an additional lowest rank in each set; plus the full ranked matrix, Supporting Information Fig. S2A), finding that differences in the binomial deviances were minimal. We further used ventiles (20-quantiles), which showed minimal average values, for all reported results.

To generate the t-SNE maps [39] with the R library Rtsne, we used log-normalized expression estimates of all genes having maximal log expression across all cells in the datasets ≥ c, where c was calculated for each dataset to equal the third quartile of log expression estimates of a reference gene set (housekeeping genes and ERCC spike-ins when available) expressed in ≤ 50% of all cells. Cells in which a high fraction (> third quartile + 2*standard deviation across all cells) of reference genes were not measured (=0) were removed from the analysis in a first step.

**Model Training and Testing**

We trained lasso logistic regression (cv.glmnet ≪ nfolds = 10 type.measure = “deviance” lower.limit = 0 family = “binomial” alpha = 1≫), elastic-net logistic regression (cv.glmnet ≪ nfolds = 10 type.measure = “deviance” lower.limit = 0 family = “binomial” alpha = 0.5≫), and random forest (randomForest ≪ importance = T proximity = T ntree = 1000 ≫) models on two of these d1–8 and tested them on the remaining one third as well as on the full d10–28 (Supporting Information Fig. S2B, S2C; Table S1) datasets, using the R libraries glmnet [40] and randomForest [41]. We used a set of 4,528 genes commonly annotated (but not necessarily expressed) in all train and test data (d1–d28) (All.g; Supporting Information Table S3) as starting features. We define commonly annotated genes as human-mouse orthologous genes with shared gene symbol across species and for which expression information (even if equal to 0) is present across all used datasets. For
comparison, we also trained: (a) lasso logistic regression models (cvglmnet $\leftarrow$ nfolds = 10 type.measure = “deviance” lower.limit = 0 family = “binomial” alpha = 1 $\Rightarrow$) using as features only genes part of previous population-derived stemness signatures—PluriNet [42] and Wong et al. [43] (Lasso.LogReg, l1.Pruni-
Net.g, and l1.Wong.g); (b) logistic regression models (glm $\leftarrow$ family = “binomial”) using as features only genes part of previous population-derived stemness signatures—PluriNet [42] and Wong et al. [43] or retained from the full set of 4,528 genes after l1 (lasso) regularization (LogReg, PluriNet.g, Wong.g, and SSPCI.g); (3) a logistic regression model (glm $\leftarrow$ family = “binomial”$\Rightarrow$) using as features the predicted probabilities for a cell to be in G1 or G2/M phase of the cell cycle [25] (Fig. 2A, 2C, Supporting Information Fig. S2B, S2C and Table S3). We also calculated the average correlation between a cell’s gene expression and the average expression of known positive SSPCs (using log-normalized expression estimates of the 4,528 shared genes and Spearman’s rank correlation coefficient), exploring how well this measure can be used to classify cells (Fig. 2A, 2C). ROCR curves were generated to visualize the performance of all distinct trained models on both the test and the independent test datasets using the ROC R package (performance $\leftarrow$ “tpr”, “fpr” $\Rightarrow$) [44].

The final reported model (referred to as SSPCI), was trained using lasso logistic regression and cross-validation, as implemented in the R library glmnet [40] (cvglmnet $\leftarrow$ nfolds = 10 type.measure = “deviance” lower.limit = 0 family = “binomial” alpha = 1 $\Rightarrow$). We generated predictions on new data using the function predict with the parameters $\prec$ pred.type = response $\succ$ for obtaining fitted probabilities of class 1, referred to as “stemness probabilities.” We used $\prec$ pred.type = class $\succ$ s = lambda.1se $\Rightarrow$ for binary predictions (1 = SSPC, 0 = non-
SSPC), corresponding to the largest value of lambda for which the error is within one standard error of the minimum. Stemness probabilities were displayed using the function beanplot [45] (what = c(1,1,1,0) overallline = “median” log = “bw” “nrdo” $\Rightarrow$).

The model was trained on two thirds of the data by 10-fold-cross-
validation and then tested on the remaining one third of the data (not used for parameterization) as well as the completely inde-
pendent test data. Feature selection was automatically per-
formed by the lasso regularization, which sets the coefficients of a large number of genes to zero. The intersection of features retained >10% of the time (>10 across 100 repetitions) was fur-
ther used as final feature set, corresponding to 23 genes in the final model. By constraining the model with $\prec$ lower.limit = 0 $\succ$, we only obtained positive coefficients, more easily biologically interpretable as genes associated with SSPCs. We refer to the probability of being classified as positive as the “probability (p) of stemness” and to the cells predicted as positives (i.e., $p \geq 0.5$) as “somatic stem and progenitor cells” or “SSPCs” throughout the manuscript.

Gene Sets

One constraint of the initial training gene set is that it should ideally contain universally expressed orthologous genes across mouse and human, which are likely to be present in a new prediction dataset. The intersection of all our train and test data (d1-d28, irrespective of their expression level) resulted in 4,528 genes (All.g, Supporting Information Table S3), the majority of which we assume is likely to be universally measurable and thus present in future datasets. We assessed the robustness of the obtained gene signature to variations in the initial starting gene set by altering the test data (removing one or two datasets, and thus starting with 4,841, 6,803, and 7,317 genes, respectively), or by subsampling (10 repetitions) from the largest (7,317 genes) initial gene set. The vast majority (19 of 23) of signature genes were also retained in $\geq 80\%$ of these alternative starting sets (Supporting Information Fig. S5C, Table S4). Moreover, among the top 10 signature genes ordered by logistic regression coefficient (and thus relative importance), 9 were retained in $>80\%$ of all tested models. Together, these results strongly sug-
gest that the gene signature used by the classifier is robust.

Furthermore, we used genes part of previous population-
derived stemness signatures—PluriNet [42] and Wong et al. [43] (PluriNet.g and Wong.g) to test how informative they are in separ-
ating scRNA-seq assessed SSPCs from non-SSPCs, as well as gene sets previously specifically associated with ISC function (1: pre-
vously used ISC marker genes, as listed on Wikipedia on 03.12.2015, ISC.g1 and 2: Munoz et al. ISC signature genes, ISC.g2 [46]), genes annotated with the “cell cycle” GO Term (GO:0007049) (Cyc.g), genes annotated with various differentiation and stemness-related terms, including “differentiation,” “development,” “stem cell main-
tenance,” “stem cell development,” “stem cell division,” “stem cell commitment” (“GO:00030154,” “GO:0048468,” “GO:0019827,” “GO:0048864,” “GO:0017145,” “GO:0072089,” “GO:0048865,” “GO:0048863”) (Diff.g), and genes annotated with the “Metabolism” GO Term (GO:0008152) (Metab.g) to validate the biological relevance of the reported stemness probability (Supporting Information Table S3). We calculated the Spearman’s rank correlation coefficient (rho) of the stemness probability with the expression level of genes in these sets. We found that the correlations obtained for the two ISC-specific gene sets were signific-
antly higher than those obtained for the background or for all other gene sets ($p \leq 0.005$ for any of the comparisons, Wilcoxon rank sum test), suggesting that our approach indeed uncovered genes that are functionally relevant for stemness.

To identify stemness-specific genes in the intestinal organ-
oids [9] (Supporting Information Table S3, ISC.g3), in the human pancreas [12, 36] (Supporting Information Table S2), and in the mouse epidermis [35] (Supporting Information Table S2) based on stemness probability, we took a two-step approach: (a) we only included genes expressed (>0 read counts) in $\geq 60\%$ of the predicted positives and $\leq 20\%$ of the predicted negatives and (b) we only included genes with a log expression estimate highly correlating (Pearson’s r, FDR 0.05) with the stemness p. Finally, we analyzed the properties of genes retained by SSPCI (SSPCI.g) using StemChecker as well as gene ontology (Supporting Information Table S4).

We also used a list of “housekeeping” genes, expected to be universally expressed at similar levels across diverse scRNA-seq data based on their presence (>50% of cells) and high (FDR 0.05) Gaussian rank correlation with ERCC spike-ins in $\geq 2$ datasets (Supporting Information Table S3).

Other Methods

Comparisons of stemness probabilities and expression values were performed using one-sided Wilcoxon rank sum tests and the significance of overlaps was assessed using one-sided Fisher’s exact tests. The gene ontology enrichment was performed using the topGO library, the $\ll$elimCount$\gg$ method and a $p$ value cut-off of .001. Overlaps with population-derived signature gene sets were assessed using the webservice StemChecker [20]. Cell cycle
stages were predicted using Cyclone with default parameters and based on known markers [25]. All analyses were performed using R version 3.1.1 and Bioconductor version 3.0.

For the intestinal organoid analyses, statistical computations were performed using GraphPad Prism6. Experimental data are presented as mean ± standard deviation. Statistical significance was assessed by two-tailed unpaired Student’s t test; the data were considered not significant (ns) for p > .05.

**Mouse Model**

Animal experiments were performed in accordance with protocols approved by the “Service de la Consommation et des Affaires Vétérinaires” of Canton Vaud, Switzerland. APC<sup>m<sub>−min</sub></sup> mice were described previously [47]. This strain was backcrossed onto C57BL/6 for at least 10 generations.

**Isolation of Intestinal Tissue and Organoid Culture**

Organoid cultures were established from total intestinal crypt preparations of APC<sup>m<sub>−min</sub></sup> mice as described previously [48]. Freshly isolated small intestines were incised along their length and villi were removed by scraping. Then, the adenomas were isolated and after several washes, the tissue was incubated in PBS/EDTA (Sigma-Aldrich, St. Louis, MO, US) (2 mM), pH8 for 5 minutes at 4°C. Gentle shaking removed remaining villi, and intestinal tissue was subsequently incubated in phosphate-buffered saline (PBS)/EDTA (2 mM), pH8 for 30 minutes at 4°C. Upon dissociation, samples were passed through a 70 μm filter and washed four times in cold Advanced Dulbecco’s Modified Eagle’s Medium (DMEM)/F12, (Gibco, NY, US) media. Afterward crypt cells were embedded in Matrigel (BD Biosciences, San Jose, CA, US) and plated in 400 μl of organoid media (Advanced DMEM/F12 with B27, N2, and N-acetylcysteine; containing growth factors EGF 50 ng/ml (Invitrogen, NY), R-Spondin 1 μg/ml, Noggin 100 μg/ml) into 24-well plates at a concentration of 500 crypts per well. Growth factors were added every other day and the entire medium was changed every 4 days. Secondary organoid culture was achieved by removing the organoids from Matrigel followed by mechanical dissociation with a glass pipette, gentle centrifugation (800 rpm), and then transferred to fresh Matrigel. Serial passaging (passage 1, passage 3, and passage 4) of shRNA-expressing organoids was compared to control organoids.

**Plasmids, shRNAs, and Protein Production**

Lcp1 (5' TTGAAAGATGTCGGTGGTGGG 3') and Vglil4 (5' TCACTGCTTCTTCTTACGGG 3') were suppressed by shRNAs with the indicated antisense oligos in a MIR30 backbone expressed from a lentiviral vector (pSM2, Openbiosystems). The expression plasmid for R-spondin1 was a kind gift from Calvin Kuo (Stanford University, U.S.) and this and Noggin proteins were produced as published [49] using proteinG or Ni-NTA purification, respectively.

**Cell Culture and Selection of shRNA-TurboRFP Gene Suppression Cells by Flow Cytometry**

The mouse colon carcinoma-derived cells CT26 were cultured in DMEM 10% fetal bovine serum (FBS, Invitrogen, NY, US). The shLcp1-TurboRFP, shVglil4-TurboRFP, and their respective controls were generated by lentiviral transduction. To this end, we generated a lentiviral vector driving TurboRFP expression that enables identification of positive cells by fluorescence-activated cell sorting (FACS). TurboRFP<sup>+</sup> cells were sorted from a population of transduced cells by flow cytometry (FACSAria II; Beckton Dickinson (BD) Biosciences, Franklin Lakes, NJ, USA, or MoFlo; Dako, Hamburg, Germany).

**Lentiviral Production and Transduction**

Lentiviruses were generated in 293T cells using third generation lentivirus packaging vectors, and virus particles were concentrated by ultracentrifugation for 2 hours at 22,000 rpm. Lentiviral titration was performed in 293T cells. Cell lines were infected with lentiviruses overnight at 37°C. Colon carcinoma cell lines (CT26) and murine intestinal organoids were infected by lentiviral transduction as published [50]. Lentiviral infection was performed after 5 minutes Trypsin-EDTA (37°C) organoid dissociation. After several washes with 5%FBS-PBS, the cells were spun with the lentiviral particles at 1,300 rpm for 20 minutes and incubated with organoid media for 2 hours at 37°C. The pellet of single cells was then embedded in Matrigel and fresh organoid medium was added. Single cells gave rise to organoids after 10 days of culture. Images and qRT-PCR analyses were performed between day 13 and 17. Serial passaging was then performed at the following time points: P1 day 18, P2 day 28, P3 day 35, and P4 day 40.

**Real-Time-qPCR**

RNA was prepared using the mini or micro RNA kit (Qiagen, Germany) from organoids. cDNAs was synthesized using Superscript-II reverse transcriptase (Invitrogen, NY, US) and oligoD<sub>7</sub> priming. qPCR was performed in a StepOnePlus thermocycler (Applied Biosystems Thermo Fisher, NY, US) using the Power SYBR green PCR Master Mix (Applied Biosystems, Thermo Fisher, NY, US) and the specific primers are listed below. Relative gene expression was determined by the comparative Ct method.

<table>
<thead>
<tr>
<th>Murine gene</th>
<th>Forward primer</th>
<th>Reverse primer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anpep</td>
<td>CTTGTAGCAAAGATGTTGAGATT</td>
<td>GGATGAGATGGCAAAAGAGAAA</td>
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<tr>
<td>Lgr5</td>
<td>CTCCACAGCTTCGAGATACAG</td>
<td>AACCAAGGCTAATGGCAGCGAAT</td>
</tr>
<tr>
<td>MkI67</td>
<td>GATGGAAGATTGGAGAACA</td>
<td>CTGCTCTTTCACAGATTTCAAG</td>
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<tr>
<td>Tff3</td>
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</table>

**RESULTS**

**Accurate Identification of SSPCs Across Biological Systems**

We collected diverse publicly available scRNA-seq data across biological systems and germ layers (Fig. 1, Supporting Information Fig. S1; Table S1, and Methods), using only two-thirds of the cells from each dataset (138 positives consisting of mostly stem cells but also early progenitors, including neural and hematopoietic stem cells as well as MDPs, and 170 negatives consisting of differentiated cells such as neuronal cells, T cells, and lung epithelial cells) for 10-fold-cross validation.
based feature (gene) selection and parameterization. We separately retained a test set (one-third, 69 positives and 85 negatives) completely untouched during training and also used completely independent (467 positives and 554 negatives, including distinct studies and germ layers) test data (Fig. 1, Supporting Information Table S1 and Methods). We note that for simplicity, we refer to the probability of being classified as positive as the “probability (p) of stemness” and to the cells predicted as positives (i.e., p > .5) as “somatic stem and progenitor cells” or “SSPCs” throughout the article.

We evaluated a broad range of models on both test and independent test data and found that de novo lasso logistic regression (referred to as somatic stem and progenitor cell identifier, SSPCI, hereafter) showed best overall performance, followed by de novo elastic-net logistic regression and de novo randomForest (Fig. 2A–2D, Supporting Information Fig. S2B, S2C and Methods). While average correlation-based and logistic-regression models trained on population-derived stemness gene sets (both with or without regularization) performed only marginally inferior to SSPCI on the test samples (Fig. 2A), they showed worse performance on the independent test samples (Fig. 2C). Remarkably, SSPCI identified 89% of all SSPCs and 97% of all non-SSPCs as such (Fig. 2B, 2D, Supporting Information Table S1).

Importantly, SSPCI was also able to identify SSPCs in completely independent biological systems such as the mouse intestine [9], corresponding to a germ layer not originally covered in the training data (Fig. 2D, 2E, Supporting Information Fig. S2D). In particular, LGR5+ ISCs were attributed significantly higher stemness probabilities than the LGR5- cells (Fig. 2E, p = 10^{-32}, Wilcoxon rank sum test) and only 17 (of a total of 229) LGR5-cells were classified as positives. All of the latter were transcriptionally similar to the LGR5+ population, as visualized in a 2D t-SNE projection [39] (Fig. 2E, Supporting Information Fig. S2D). These cells likely represent highly proliferative and multipotent transit amplifying cells and are also partially characterized by high expression of Hopx and Gmi3, other commonly used stem cell markers (Fig. 2E, Supporting Information Fig. S2D) [46, 51, 52]. It is thus possible that some of these misclassification events do not represent false positive predictions, as these multipotent cells may genuinely be very close to ISCs and may thus also be categorized as SSPCs [5]. As an additional control, we tested the classifier on human liver data and confirmed that neither adult hepatocytes nor endothelial cells were classified as positives (Supporting Information Fig. S2E, S2F).

We next asked whether the cell cycle stage of the cells influences the performance of the classifier, given that proliferation and the ability to self-renew by cell division are intrinsically linked to stemness. To do so, we used a recently developed scRNA-seq-based method to attribute each cell to one specific cell cycle stage (G1, G2/M, S or if the assignment is ambiguous, na) [25]. This analysis revealed that across all our datasets, SSPCs did not show an association with a specific cell cycle stage (Fig. 2E, Supporting Information Fig. S2G, S2H). Furthermore, we found that a linear regression model trained using only the probabilities of cells being in one of the three cell cycle stages as features performs very poorly in identifying SSPCs (Supporting Information Fig. S2B, S2C). Thus, we conclude that SSPCs cannot be identified solely based on their cell cycle stage.

**De Novo Identification of SSPCs in scRNA-Seq Dissected Tissues**

We subsequently tested the power of our approach to resolve still ambiguous systems that were recently dissected by scRNA-seq: the human pancreas [12, 36, 37] and the mouse epidermis [35]. In the pancreas, we found that virtually all...
tested endocrine cells (alpha, beta, gamma, and delta) had very low stemness probabilities and were classified as negatives (Fig. 3A–3E). This result is in line with the notion that proliferation of differentiated cells is the main mechanism active in the islets [53, 54]. In contrast to endocrine cells, a substantial (10%–60%) percentage of ductal, acinar, and most notably stellate cells showed high stemness probabilities and were predicted positive (Fig. 3A–3E). Importantly, these results were highly consistent across three independent studies (Fig. 3A–3E) [12, 36, 37], and individuals [36] (Supporting Information Fig. S3A, S3B), demonstrating the robustness of our approach to laboratory-induced variability. Our findings are compatible with the increased clonogenic capacity and phenotypic plasticity of these three cell types compared to endocrine cells [55]. They also support the notion that stellate cells are major contributors to pancreatic maintenance and repair [56, 57], as our prediction suggests that the stellate cell fraction contains most SSPCs in this system. Interestingly, both disease (type I and II diabetes, respectively), and age seems to alter the SSPC proportion in the pancreas, with a higher fraction of stem-like cells in affected adults as well as healthy children compared to healthy adults (Fig. 3E). This is consistent with the observation of increased de-differentiation of alpha and beta cells in diabetes patients compared to adult controls [37].

A second system that we examined is the mouse epidermis. Using scRNA-seq of thousands of epidermal cells, our predictor revealed almost twice as many SSPCs among basal cells compared to all other cells (Fig. 3F, left), and interestingly, no difference in the fraction of predicted SSPCs between basal cells expressing at least one of the epidermal stem cell markers Cd34, Lgr5, Lgr6, Gli1, Lrig1, Krt14, and those not expressing any (Fig. 3F, right). Surprisingly, when examining which subclass of basal cells is predicted to contain most SSPCs, we found that it is the interfollicular epidermis (IFE), with 7%–13% of stem-like IFE basal cells compared to only ~4% of bulge cells (Fig. 3G). We thus focused on the IFE area and examined where high scoring cells are positioned on the previously estimated differentiation trajectory [35]. We found that these cells are indeed largely located at the root of the trajectory (Fig. 3H). Consistently, our predicted stemness probability was significantly negatively correlated to previously estimated differentiation pseudotime values (Spearman’s rho = −0.417, p < 10⁻³). In sum, our analysis supports the notion that there is no specific subcluster of SSPCs in this system.
interspersed, preponderantly basal cells with stem-like expression patterns. We found that many of the latter cells are located in the IFE and the upper hair follicle region, as also suggested recently [35].

Finally, for all predicted SSPCs in both systems, we highlighted their associated genes (Supporting Information Table S2; Fig. S3C, S3D), which can guide follow-up experiments aimed at their isolation and characterization or validate previously reported associations. For example, it has been recently suggested that the TF STMN1 may act as a marker for progenitor-like acinar cells [59]. Our data supports this association given that its expression is highly correlated with acinar SSPCs in both pancreas datasets (Supporting Information Table S2; Fig. S3E, S3F).

**Toward a Quantitative Measure of Multipotency**

Prediction results on the intestinal, pancreas, and epidermal data suggested that intermediate stemness probabilities may reveal cells in transition stages between multipotent SSPCs and fully differentiated cells (Fig. 2E, 3A–3E, Supporting Information Fig. S2D). To test the capacity of our classifier to distinguish among stem-like and differentiated cells, we analyzed...
four distinct systems featuring cells with increasing level of commitment: (a) lung development [8], (b) dendritic cell development [22], (c) megakaryocyte differentiation [26] and neurogenesis [10] (Fig. 4A, Supporting Information Table S1). In all of these datasets, we observed a progressive decrease in stemness probabilities across time/differentiation (Fig. 4A, Supporting Information Fig. S4A; Table S1). For instance, while the vast majority of multipotent MDPs and CDPs were predicted as positives, the stemness probabilities assigned to CDPs were significantly lower ($p < 10^{-9}$, Wilcoxon rank sum test) and further, the majority (68%) of dendritic cell precursors (PreDCs) were identified as negatives (Fig. 4B, Supporting Information Fig. S4A). These results suggest that the stemness probability provided by our classifier can be used as a quantitative indicator of a cell’s progression along a specific differentiation path, independently of the cell’s particular cell cycle stage at the time of measurement (Supporting Information Fig. S4B, S4C).

Having established that our derived stemness probability is a robust indicator of a cell’s differentiation status, we next asked whether it could be used to functionally associate genes with stemness. The large number of stem and non-stem cells present in the intestinal organoid data [9] enabled us to explore this question in detail. Among genes that showed significant expression correlation with stemness probability and that were largely confined to cells predicted as positives (Methods), we identified the bona fide stemness marker LGR5 as well as other surface-associated stem cell markers such as GNL3 and CD44 (Fig. 5A). A broad range of TFs were also among these genes as well as several metabolic factors such as PHGDH, which is part of the known intestinal SC (ISC) signature [46] (Fig. 5A). In total, we found 98 genes that were ISC-associated (ISC.g3, Supporting Information Table S3), representing known (26% overlapped with an ISC stemness signature) and putatively novel players in ISC function. Gene set enrichment analysis also revealed terms previously associated with stem cell function, including retinoic acid signaling, DNA replication, cell cycle, and proliferation (Supporting Information Fig. S5A).

**Signature Genes Are Required for a Stem-Like Phenotype**

Given the robust performance of our classifier across biological systems, we reasoned that some of the genes that were retained in our model (Methods and Supporting Information Table S4) are likely to be more generally required for SSPC function. Using the resource collection “StemChecker” [20], we thus asked which of these 23 genes (Fig. 5B) had previously been implicated in stem cell biology. We found that 70% of them...
were part of at least one population-derived stemness gene set, most often characteristic of neural, hematopoietic, or intestinal SCs (Bonferroni adjusted \( p \leq 0.03 \)) (Fig. 5B, 5C, Supporting Information Table S4). Thus, while the majority of our classifier genes are individually part of population-derived stemness gene sets, they have not been previously used collectively to discriminate SSPCs from differentiated cells.

Interestingly, 12 out of the 23 classifier genes showed significantly higher expression in ISCs versus all other cells (Supporting Information Fig. S5B; Table S4). This is despite the fact that no ISCs, or any other endodermal SSPCs, have been used for training the model. Among genes with higher expression in ISCs were those that have already been implicated in ISC biology such as \( \text{Myc} \) and \( \text{Ccnd2} \) [60, 61] (Fig. 5D, Supporting Information Fig. S5B). Others have only been partially linked to cell differentiation, and have so far not been implicated in ISC nor SSPC function in general. While a role in ISC function is not an absolute requirement for classifier genes with higher expression in ISCs, it is clearly intuitive. Therefore, we focused on two such genes, \( \text{Vgll4} \) and \( \text{Lcp1} \), and examined the impact of their shRNA-mediated gene suppression on intestinal organoid growth. We used intestinal organoids from APC\(^{\text{min}} \) mouse precancerous adenomas (Fig. 6A), since the upregulation of the Wnt signaling pathway induces activation and expansion of LGR5\(^+\) ISC-like cells [9], depicting expression of the TFs \( \text{Myc} \) and \( \text{Etv6} \) (gray to orange gradient, orange highest expression). Abbreviations: HSC, hematopoietic stem cell; ISC, intestinal stem cell; NSC, neural stem cell; TF, transcription factor.
We first demonstrated successful shRNA-mediated gene suppression of the two genes in the murine CT26 colon carcinoma cell line and intestinal organoids (Fig. S6A). While shRNA-mediated gene suppression in CT26 cells did not affect cell phenotype or proliferation per se (data not shown), we observed a striking reduction in the number and size of intestinal organoids in the gene suppression samples (Fig. 6B, 6C, Supporting Information Fig. S6B–S6D). Strikingly, organoid growth after Vgll4 shRNA suppression was completely impaired, such that organoids could not be passaged and RNA could not be collected for further experiments. For Lcp1, the gene suppression effects were observed both at initial and subsequent passages (Fig. 6C, Supporting Information Fig. S6B–S6D). Importantly, we also found that canonical stemness marker genes such as Lgr5, Vdr, and the potential oncogene Wwp1 [64], were significantly downregulated at passage 0 when Lcp1 was suppressed, while Ki67, a proliferation marker, showed no significant difference (Fig. 6D, 6E). Further passaging of these organoids (passage 3 and 4), showed that these shRNA-mediated gene suppressions increased differentiation markers such as Anpep (absorptive lineage) and Tff3 (secretory lineage).

Together, these results demonstrate for the first time the requirement of two classifier genes, Lcp1 and Vgll4, in APC<sup>min</sup> intestinal organoid growth and propagation. However, follow-up studies will be required to consolidate the role of these genes in stem cell function in general throughout many tissues.

**DISCUSSION**

Despite substantial progress in understanding SSPC function and their essential role in maintaining and repairing mammalian tissues, it is currently unknown how many kinds of distinct SSPCs exist, which tissues harbor them and what their molecular and functional similarities and differences are [4, 5]. Recent technological developments in the field of scRNA-seq now facilitate analyses of unprecedented molecular resolution, which enabled us to develop a universally applicable method for the de novo identification of self-renewing and multipotent cells. The timeliness of our study is demonstrated by the fact that numerous SSPC and non-SSPC scRNA-seq datasets have only very recently become available, enabling for the first time cross-system analyses. Nevertheless, we acknowledge that we were still constrained in the choice of training and test sets, leading to overrepresentation of certain lineages (hematopoietic) and germ layers (mesoderm) in our analyses as well as the lack of distinction between true stem cells and multipotent progenitor cells. Ideally, all germ layers and SSPC types (e.g., quiescent vs. active) should be equally well represented in both training and test sets to derive the best classifier. Therefore, when more datasets will become available, one can envisage updating and further scaling up the binary classification approach that we used here to a multi-class system. The underlying aim would be to increase the discriminative power of our approach by distinguishing activated and quiescent somatic SCs, multipotent and unipotent progenitors, for instance. This will in turn allow the exploration of specific transcriptional properties for each of the individual stages as well as detailed delineation of commonalities and differences between SSPCs.

A further limitation of our approach is the relative simplicity of the used data normalization (quantiles), which dampens the large dynamic expression range that is a powerful feature of RNA-seq. We took this route given the large technical and biological noise that is currently characterizing scRNA-seq.
data, likely exacerbated by working across experimental and data analysis protocols [65]. We anticipate that, with the maturation of experimental and computational scRNA-seq techniques, the actual distribution of expression estimates will be taken into account, such that the full dynamic range of this powerful method can be exploited.

Despite these limitations, we showed here that our SSPC identification method can be applied to previously completely unseen scRNA-seq datasets, across different technologies (e.g., CEL-seq and SMART-seq) [66, 67] and experimental systems, including highly proliferating (e.g., ISCs) and quiescent (long-term HSCs) SSPCs. Consequently, we envision several downstream applications for our classifier. First, it may allow the detection of stem-like cells in underexplored, heterogeneous tissues, with the aim of identifying specifically expressed marker genes to enable downstream phenotypic analyses, as shown here for the mouse epidermis and the human pancreas. Indeed, while the markers that we propose in these two systems will require further validation, we also demonstrated the feasibility of marker detection given the de novo identification of the established markers LGR5 and GNL3 in intestinal epithelial organoids [62, 68]. Second, such a classifier may support estimating the fraction of stem-like cells in a heterogeneous mix of cells. This may provide novel insights into tissue organization, as we also illustrated here for the epidermis and pancreas, or into a tissue's or cell fraction's regenerative capacity, for example over the course of aging [30], or to inform cell transplantation experiments [69]. A third application is the detection of highly malignant cells in tumors. That scRNA-seq data can be used in the context of tumor heterogeneity to reveal stemness-exhibiting cells has in fact recently been demonstrated in glioma samples [70]. While these latter applications go beyond the scope of the current study, we envisage that they will constitute very fruitful research areas in the future.

CONCLUSION

We developed a sc RNA-seq-based classifier that accurately detects SSPCs across tissues, aiding in the resolution of still poorly characterized systems. In addition, we demonstrate that this classifier can provide a quantitative measure of a cell’s progression along a differentiation path. The method’s good cross-system performance supports the notion that SSPCs can be recognized based on shared, but not necessarily identical molecular properties. Specifically, genes with high discriminative power have putative roles in SSPC biology, of which many are known such as those of Myc and Ccnd2, but others are novel such as Vgll4 and Lep1’s function in the intestinal system.

AVAILABILITY

We provide the trained model as well as the two best-performing alternative models (elastic-net logistic regression and random forest), training and test data as well as an R script to produce the predictions discussed in this manuscript and use on novel data. Available at https://github.com/DeplanckeLab/SSPCI. Predictions can also be obtained and visualized through the web-based platform ASAP at https://asap.epfl.ch/[71].

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AUTHOR CONTRIBUTIONS

P.C.S.: conception and design, data analysis and interpretation, manuscript writing, conducted all scRNA-seq related analyses; P.O.M.: performed all experiments, analyzed all data on intestinal organoids; J.H.: supervised the intestinal organoids experiments, data evaluation and financial support; B.D.: conception and design, data analysis and interpretation, manuscript writing, financial support; P.C.S., P.O.M., J.H., and B.D.: read and approved the final manuscript.

DISCLOSURE OF POTENTIAL CONFLICTS OF INTEREST

The authors indicated no potential conflicts of interest.

REFERENCES


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