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distinct: a novel approach to differential distribution analyses

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distinct: a novel approach to differential distribution analyses

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1 Abstract

2 We present *distinct*, a general method for dif-
3 ferential analysis of full distributions that is
4 well suited to applications on single-cell data,
5 such as single-cell RNA sequencing and high-
6 dimensional flow or mass cytometry data. High-
7 throughput single-cell data reveal an unprece-
8 dented view of cell identity and allow com-
9 plex variations between conditions to be discov-
10 ered; nonetheless, most methods for differential
11 expression target differences in the mean and
12 struggle to identify changes where the mean is
13 only marginally affected. *distinct* is based on
14 a hierarchical non-parametric permutation ap-
15 proach and, by comparing empirical cumulative
16 distribution functions, identifies both differen-
17 tial patterns involving changes in the mean, as
18 well as more subtle variations that do not in-
19 volve the mean. We performed extensive bench-
20 marks across both simulated and experimen-
21 tal datasets from single-cell RNA sequencing
22 and mass cytometry data, where *distinct* shows
23 favourable performance, identifies more differ-
24 ential patterns than competitors, and displays
25 good control of false positive and false discovery
26 rates. *distinct* is available as a Bioconductor R
27 package.

28 **keywords:** Differential distribution; Differential anal-
29 yses; Differential state; High-throughput single-cell
30 data; Single-cell RNA-seq; Single-cell flow and mass cy-
31 tometry; Permutation tests.

32 Background

33 Technology developments in the last decade have led to
34 an explosion of high-throughput single-cell data, such
35 as single-cell RNA sequencing (scRNA-seq) and high-
36 dimensional flow or mass cytometry data, allowing re-

37 searchers to investigate biological mechanisms at single-
38 cell resolution. Single-cell data have also extended the
39 canonical definition of differential expression by dis-
40 playing cell-type specific responses across conditions,
41 known as differential state (DS) [32], where genes or
42 proteins vary in specific sub-populations of cells (e.g.,
43 a cytokine response in myeloid cells but not in other
44 leukocytes [13]). Classical bulk differential expression
45 methods have been shown to perform well when used
46 on single-cell measurements [25, 26, 31] and on aggre-
47 gated data (i.e., averages or sums across cells), also re-
48 ferred to as pseudo-bulk (PB) [7, 32]. However, most
49 bulk and PB tools focus on shifts in the means, and
50 may conceal information about cell-to-cell heterogene-
51 ity. Indeed, single-cell data can show more complex
52 variations (Figure 1 and Supplementary Figure 1); such
53 patterns can arise due to increased stochasticity and
54 heterogeneity, for example owing to oscillatory and un-
55 synchronized gene expression between cells, or when
56 some cells respond differently to a treatment than oth-
57 ers [15, 31]. In addition to bulk and PB tools, other
58 methods were specifically proposed to perform differ-
59 ential analyses on single-cell data (notably: *scDD* [15],
60 *SCDE* [14], *MAST* [11], *BASiCS* [10, 29, 30] and mixed
61 models [27]). Nevertheless, they all present significant
62 limitations: *BASiCS* does not perform cell-type spe-
63 cific differential testing between conditions, *scDD* does
64 not directly handle covariates and biological replicates,
65 while *PB*, *SCDE*, *MAST* and mixed models performed
66 poorly in previous benchmarks when detecting differ-
67 ential patterns that do not involve the mean [7, 15].

68 Results

69 *distinct*'s full distribution approach

70 To overcome these challenges, we developed *distinct*, a
71 flexible and general statistical methodology to perform
72 differential analyses between groups of distributions.

73 *distinct* is particularly suitable to compare groups of
 74 samples (i.e., biological replicates) on single-cell data.

75 Our approach computes the empirical cumulative dis-
 76 tribution function (ECDF) from the individual (e.g.,
 77 single-cell) measurements of each sample, and compares
 78 ECDFs to identify changes between full distributions,
 79 even when the mean is unchanged or marginally in-
 80 volved (Figure 1 and Supplementary Figure 1). First,
 81 we compute the ECDF of each individual sample; then,
 82 we build a fine grid and, at each cut-off, we average the
 83 ECDFs within each group, and compute the absolute
 84 difference between such averages. A test statistic, s^{obs} ,
 85 is obtained by adding these absolute differences.

86 More formally, assume we are interested in compar-
 87 ing two groups, that we call A and B , for which N_A
 88 and N_B samples are available, respectively. The ECDF
 89 for the i -th sample in the j -th group, is denoted by
 90 $ecdf_i^{(j)}(\cdot)$, for $j \in \{A, B\}$ and $i = 1, \dots, N_j$. We
 91 then define K equally spaced cut-offs between the mini-
 92 mum, min , and maximum, max , values observed across
 93 all samples: b_1, \dots, b_K , where $b_k = min + k \times l$, for
 94 $k = 1, \dots, K$, with $l = (max - min)/(K + 1)$ being
 95 the distance between two consecutive cut-offs. We ex-
 96 clude min and max from the cut-offs because, trivially,
 97 $ecdf_i^{(j)}(min) = 0$ and $ecdf_i^{(j)}(max) = 1$, $\forall j, i$. At ev-
 98 ery cut-off, we compute the absolute difference between
 99 the mean ECDF in the two groups; our test statistic,
 100 s^{obs} , is obtained by adding these differences across all
 101 cut-offs:

$$s^{obs} = \sum_{k=1}^K \left| \frac{\sum_{i=1}^{N_A} ecdf_i^{(A)}(b_k)}{N_A} - \frac{\sum_{i=1}^{N_B} ecdf_i^{(B)}(b_k)}{N_B} \right|. \quad (1)$$

102 Note that in differential state analyses, these operations
 103 are repeated for every gene-cluster combination.

104 Intuitively, s^{obs} , which ranges in $[0, \infty)$, approximates
 105 the area between the average ECDFs, and represents
 106 a measure of distance between two groups of densities:
 107 the bigger s^{obs} , the greater the distance between groups.
 108 The number of cut-offs K , which can be defined by
 109 users, is set to 25 by default, because no detectable
 110 difference in performance was observed when further
 111 increasing it (data not shown). Note that, although at
 112 each cut-off we compute the average across each group's
 113 curves, ECDFs are computed separately for each indi-
 114 vidual sample, therefore our approach still accounts for
 115 the within-group variability; indeed, at a given thresh-
 116 old, the average of the sample-specific ECDFs differs
 117 from the group-level ECDF (i.e., the curve based on
 118 all individual measurements from the group). The null
 119 distribution of s^{obs} is then estimated via a hierarchical

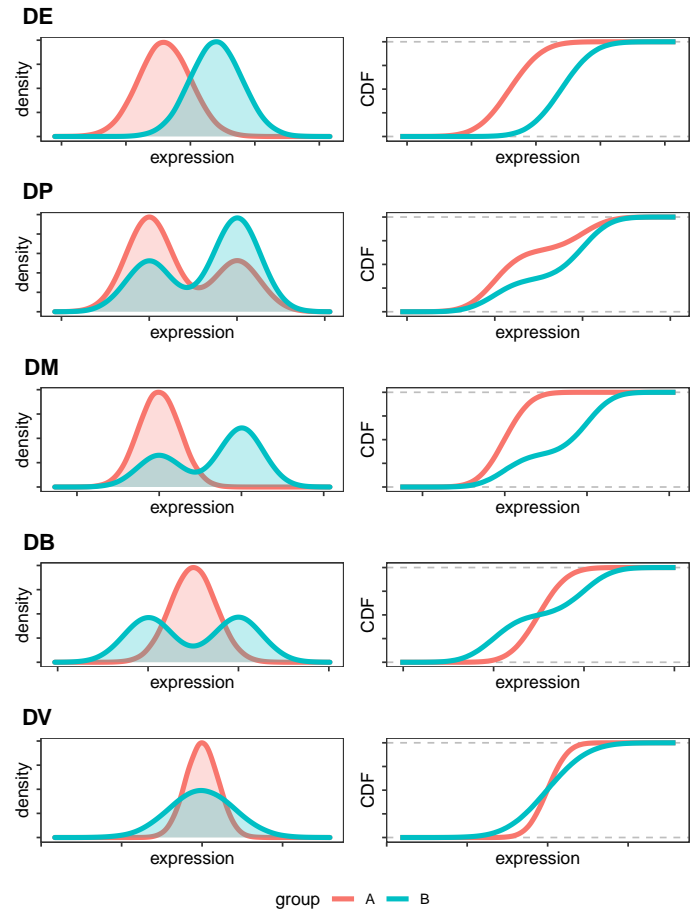


Figure 1: Cumulative distribution functions (CDFs) unravel differences between distributions. Density (left panels) and CDF (right panels) of five differential patterns: differential variability (DV), and the four proposed by Korthauer et. al. [15]: differential expression (DE), differential proportion (DP), differential modality (DM), and both differential modality and different component means (DB).

120 non-parametric permutation approach (see Methods).
 121 A major disadvantage of permutation tests, which of-
 122 ten restricts its usage on biological data, is that too
 123 few permutations are available from small samples. We
 124 overcome this by permuting cells, which is still pos-
 125 sible in small samples, because there are many more
 126 cells than samples. In principle, this may lead to an
 127 inflation of false positives due to lack of exchangeabil-
 128 ity (see Methods); nonetheless, in our analyses, *distinct*
 129 provides good control of both false positive and false
 130 discovery rates.

131 Importantly, *distinct* is general and flexible: it targets
 132 complex changes between groups, explicitly models bio-
 133 logical replicates within a hierarchical framework, does
 134 not rely on asymptotic theory, avoids parametric as-
 135 sumptions, and can be applied to arbitrary types of
 136 data. Additionally, *distinct* can also adjust for sample-
 137 level cell-cluster specific covariates (i.e., whose effect
 138 varies across cell clusters), such as batch effects. In par-
 139 ticular, *distinct* fits a linear mixed effects model with

140 the input data (e.g., normalized counts) as response
141 variable, nuisance covariates as fixed effects, and sam-
142 ples as random effects. The method then removes the
143 estimated impact of fixed effect covariates, and per-
144 forms differential testing on these normalized values
145 (see Methods).

146 Furthermore, to enhance the interpretability of differen-
147 tial results, *distinct* provides functionalities to compute
148 (log) fold changes between conditions, and to plot den-
149 sities and ECDFs, both for individual samples and at
150 the group-level.

151 Note that, although *distinct* and the Kolmogorov-
152 Smirnov [18] (KS) test share similarities (they both
153 compare distributions via non-parametric tests), the
154 two approaches present several conceptual differences.
155 Firstly, the KS considers the maximum distance be-
156 tween two ECDFs, while our approach estimates the
157 overall distance between ECDFs, which in our view is
158 a more appropriate way to measure the difference be-
159 tween distributions. Secondly, the KS test only com-
160 pares two individual densities, while our framework
161 compares groups of distributions. Thirdly, while the
162 KS statistic relies on asymptotic theory, our framework
163 uses a permutation test. Finally, a comparison between
164 *distinct* and *scDD* [15] based on the KS test (labelled
165 *scDD-KS*) shows that our method, compared to the KS
166 test, has greater statistical power to detect differential
167 effects and leads to fewer false discoveries (see Simula-
168 tion studies).

169 Simulation studies

170 We conducted an extensive benchmark, based on
171 scRNA-seq and mass cytometry simulated and experi-
172 mental datasets to investigate *distinct*'s ability to iden-
173 tify differential patterns in sub-populations of cells.

174 First, we simulated droplet scRNA-seq data via *mus-*
175 *cat* [7] (see Methods). We ran five simulation repli-
176 cates for each of the differential profiles in Figure 1,
177 with 10% of the genes being differential in each clus-
178 ter, where DE (differential expression) indicates a shift
179 in the entire distribution, DP (differential proportion)
180 implies two mixture distributions with different propor-
181 tions of the two components, DM (differential modal-
182 ity) assumes a unimodal and a bimodal distribution,
183 DB (both differential modality and different component
184 means) compares a unimodal and a bimodal distribu-
185 tion with the same overall mean, and DV (differential
186 variability) refers to two unimodal distributions with
187 the same mean but different variance (Figure 1 and
188 Supplementary Figure 1). Each individual simulation
189 consists of 4,000 genes, 3,600 cells, separated into 3 clus-

ters, and two groups of 3 samples each, corresponding
191 to an average of 200 cells per sample in each cluster.

192 We considered six different normalization approaches:
193 counts per million (CPMs), *scater*'s logcounts [19],
194 *linnorm* [34], *BASiCS* [10,29,30], *SCnorm* [3] and
195 residuals from variance stabilizing normalization from
196 *sctransform* (*vstresiduals*) [12]. We compared *dis-*
197 *tinct* to several PB approaches from *muscat*, based on
198 *edgeR* [24], *limma-voom* and *limma-trend* [23], which
199 emerged among the best performing methods for differ-
200 ential analyses from scRNA-seq data [7,26]. We further
201 considered three methods from *muscat* based on mixed
202 models (MM), namely *MM-dream2*, *MM-vstresiduals*
203 and *MM-nbinom* (see Methods). Finally, we included
204 *scDD* [15], which is conceptually similar to our ap-
205 proach: *scDD* implements a non-parametric method to
206 detect changes between individual distributions from
207 scRNA-seq, based on the Kolmogorov-Smirnov test,
208 *scDD-KS*, and on a permutation approach, *scDD-perm*.
209 For *scDD-perm* we used 100 permutations to reduce the
210 computational burden.

211 In all scenarios and on all six input data, *distinct* shows
212 favourable performance: it has good statistical power
213 while controlling for the false discovery rate (FDR)
214 (Figure 2). In particular, for DE, DP and DM, *distinct*
215 has similar performance to the best performing com-
216 petitors (*edgeR.linnorm* and *limma-trend.logcounts*),
217 while for DB and DV, it achieves significantly higher
218 true positive rate (TPR), especially when using *log-*
219 *counts*. PB methods in general perform well for differ-
220 ential patterns involving changes in the mean (DE, DP
221 and DM), but struggle to identify DB and DV patterns.
222 *scDD* provides good TPR across all patterns when us-
223 ing the KS test on *vstresiduals* (*scDD-KS.vstresiduals*),
224 while the TPR is significantly reduced when using
225 other inputs and with the permutation approach (*scDD-*
226 *perm*); however, *scDD* methods (in particular, *scDD-*
227 *KS.vstresiduals*) also show a significant inflation of the
228 FDR. In contrast, MM methods provide good control of
229 the FDR but have low statistical power in all differen-
230 tial scenarios. We also investigated how normalization
231 influences each method's results (Supplementary Fig-
232 ure 2): *distinct* appears to be the least affected method
233 and displays the smallest variation across normaliza-
234 tion inputs, possibly due to its non-parametric struc-
235 ture, which can more flexibly accommodate various in-
236 puts. Given the computational cost of *SCnorm*, which
237 is significantly higher than the other normalizations,
238 we only included this approach in the results from the
239 main simulations. Furthermore, among the 25 replicate
240 datasets in Figure 2, *SCnorm* ran in a few minutes on
241 10 simulations, while it failed to run within a week time

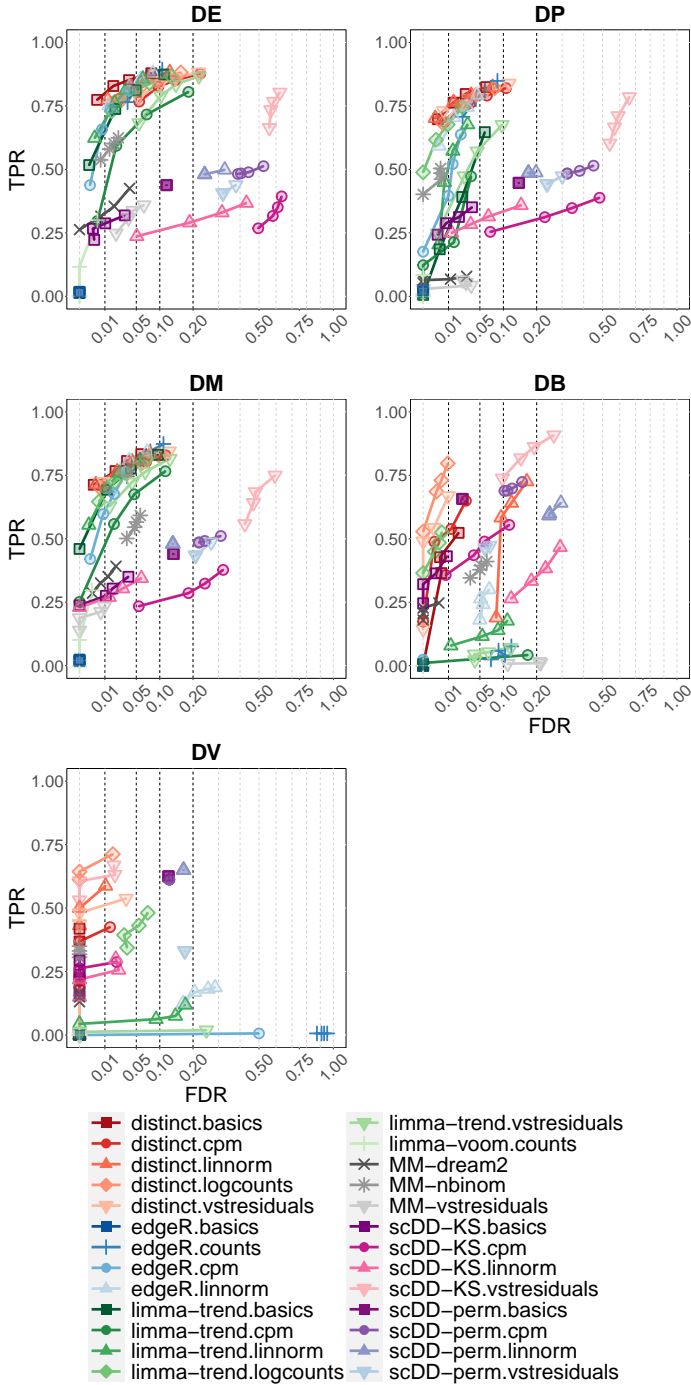


Figure 2: *distinct* identifies various differential patterns and controls for the FDR. TPR vs. FDR in *muscat* simulated data; DE, DP, DM, DB and DV refer to the differential profiles illustrated in Figure 1. Circles indicate observed FDR for 0.01, 0.05, 0.1 and 0.2 significance thresholds. Results are averages across the five simulation replicates. Each individual replicate consists of 4,000 genes, 3,600 cells, separated into 3 clusters, and two groups of 3 samples each, corresponding to an average of 200 cells per sample in each cluster.

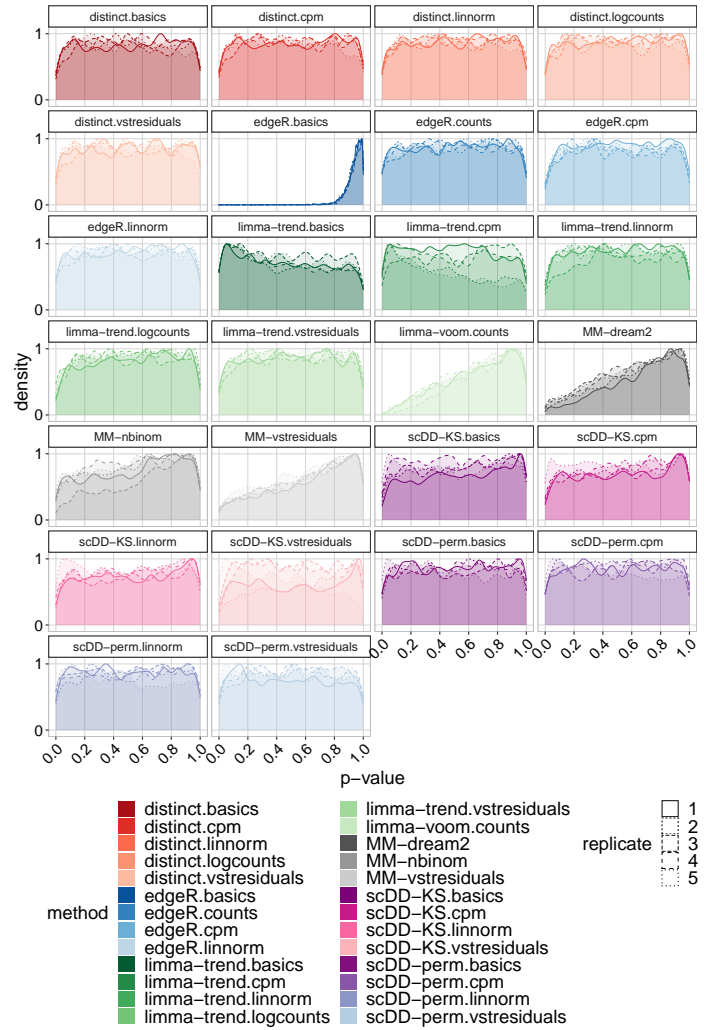


Figure 3: *distinct* has uniform null p-values. Density of raw p-values in *muscat* null simulated data; each replicate represents a different null simulation. Each individual replicate consists of 4,000 genes, 3,600 cells, separated into 3 clusters, and two groups of 3 samples each, corresponding to an average of 200 cells per sample in each cluster.

(on 10 cores) on the remaining 15 datasets. Therefore, we excluded *SCnorm* from Figure 2 and, in Supplementary Figures 3 and 4, we report a comparison of *SCnorm* to the remaining normalization methods, on the subset of 10 simulations where all normalizations successfully ran. For *distinct*, *edgeR* and *limma*, no noticeable differences are detected between *SCnorm* and

the remaining normalization methods, while for *scDD-KS SCnorm* leads to a higher inflation of the FDR.

We further simulated five null simulation replicates with no differential patterns; again with each simulation having 4,000 genes, 3,600 cells, 3 cell clusters and two groups of 3 samples each. In the null simulated data, only *limma-trend.basics* and *limma-trend.cpm* present a mild inflation of false positives, while MM and, particularly, *edgeR.basics* lead to overly conservative p-values; instead, *distinct* and *scDD* show approximately uniform p-values for all types of input data (Figure 3).

We also extended previous simulations to add a cell-type specific batch effect (i.e., a batch effect that affects differently each cell-type) [7,17]. In particular, we simulated 2 batches, that we call b_1 and b_2 , with one group of samples having two samples associated to b_1 and one to b_2 , and the other group of samples having two samples from batch b_2 and one from b_1 . Differential results

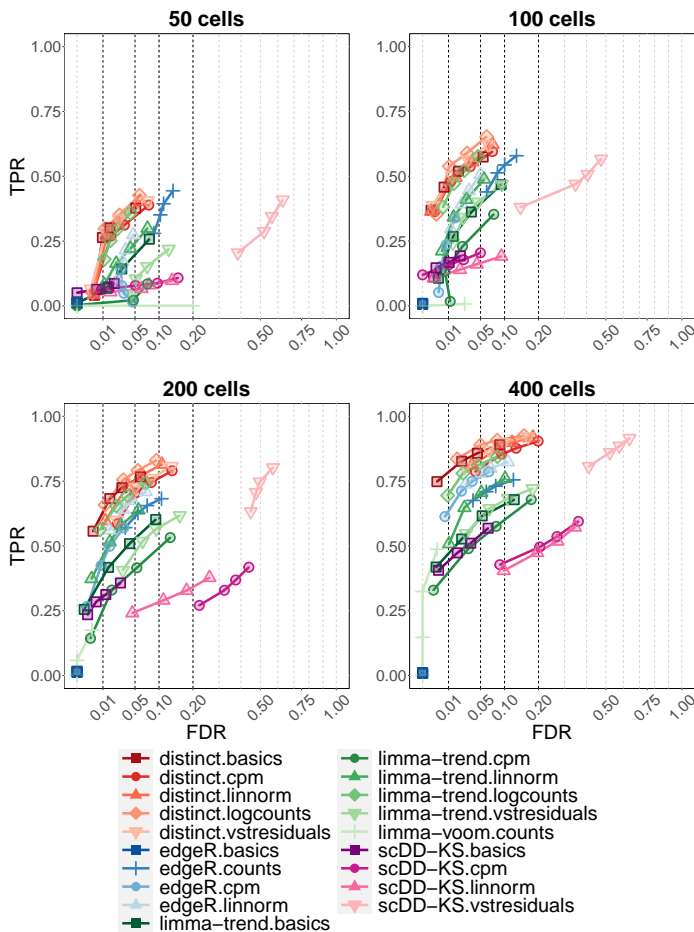


Figure 4: *distinct* achieves good performance when varying the number of available cells. TPR vs. FDR in *muscat* simulated data; with 50, 100, 200 and 400 cells per cluster-sample combination, corresponding to a total of 900, 1,800, 3,600 and 7,200 cells, respectively. Results are aggregated over the five replicate simulations of each differential type (DE, DP, DM, DB and DV), contributing in equal fraction. Each individual simulation replicate consists of 4,000 genes, 3 cell clusters and two groups of 3 samples each. Circles indicate observed FDR for 0.01, 0.05, 0.1 and 0.2 significance thresholds. Note that *scDD-perm* and MM were excluded from this analysis due to their computational cost.

are substantially unchanged (Supplementary Figure 5), which shows *distinct* can effectively remove nuisance confounders.

Furthermore, we performed various sensitivity analyses and investigated how results are affected when varying: i) the number of cells, ii) the library size, iii) the dispersion parameter, iv) the fraction of significant genes, and v) the sample sizes in each group. In particular, we simulated 50, 100, 200 (as in the original simulation) and 400 cells per sample in each cluster. We further modified the library size and dispersion parameters of the negative binomial model used by *muscat* to simulate scRNA-seq data, influencing the mean expression and cell-to-cell variability respectively, by considering values 1/5, 1/2, 2 and 5 times as big as those used in the original simulation. In addition, we varied the per-

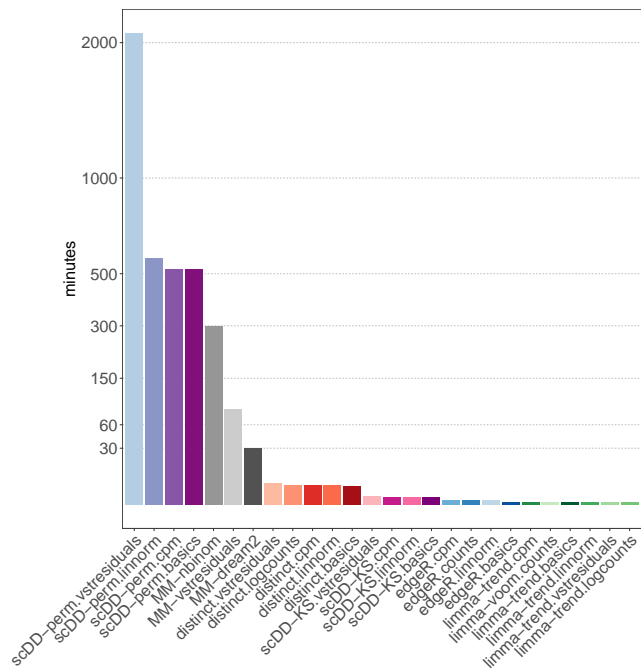


Figure 5: *distinct* requires more computational resources than PB and *scDD-KS* methods, but significantly less than MM and *scDD-perm* models. Average computing time, expressed in minutes, in *muscat* main simulations (Figures 2-3). For each method, times are averaged across simulation types (DE, DP, DM, DB, DV and null) and, for each type, across the five replicate simulations; in each replicate 3,600 cells are available (200, on average, per cluster-sample combination). *distinct*, MM and *scDD* models were run on 3 cores, while pseudo-bulk methods based on *edgeR* and *limma* used a single core because they do not allow for parallel computing. Note that *scDD-perm* requires much longer on vstresiduals than on the other normalized data, because *scDD* performs differential testing on non-zero values: vstresiduals, (unlike linnorm, cpm and basics normalized data) are not zero-inflated and, therefore, many more cells have to be used for differential testing.

percentage of simulated differential genes as 1, 5, 10 (as in the original simulation) and 20%, and considered various unbalanced designs by comparing two groups of different sample sizes: 3 vs. 2, 4 vs. 3, and 5 vs. 3. Overall, increasing the number of cells or the library size and decreasing the dispersion have a positive impact on the performance of all methods, by improving their ability to detect differential effects (i.e., true positive rate); nonetheless, none of these factors seem to affect the relative ranking of methods, which remains globally stable (Figure 4 and Supplementary Figures 6-7). In addition, changing the fraction of significant genes and considering unbalanced designs does not appear to introduce systematic changes in performance (Supplementary Figures 8-9). Note that, in these sensitivity analyses, we excluded MM models due to the high computational cost and low statistical power displayed in the previous analyses.

From a computational perspective, *distinct* required an average time of 3.2 to 4.5 minutes per simulation, which is higher than PB methods (0.1 to 0.2 minutes)

304 and *scDD-KS* (0.5 to 0.7 minutes), but significantly
 305 lower than MM approaches (29.4 to 297.3 minutes) and
 306 *scDD-perm* (544.7 to 2085.6 minutes) (Figure 5 and
 307 Supplementary Table 1). All methods were run on 3
 308 cores, except PB approaches, which used a single core,
 309 because they do not allow for parallel computing.

310 We also considered an alternative popular droplet
 311 scRNA-seq data simulator, *SplatPOP* [2], which rep-
 312 resents a generalization of *Splatter* [35], that allows
 313 multi-sample multi-group synthetic data to be gener-
 314 ated. In particular, we simulated 20,345 genes from
 315 a human genome with two groups of 4 samples each,
 316 and 100 cells per sample, belonging to the same clus-
 317 ter of cells, for a total of 800 cells across all samples.
 318 We ran 8 differential simulations, with 10% of genes
 319 truly differential between groups, by varying the loca-
 320 tion (*de.facLoc*) and scale (*de.facScale*) differential
 321 parameters, mainly affecting the mean and variance,
 322 respectively (see Methods). We considered the same
 323 normalization and differential methods as in the *mus-*
 324 *cat* simulation (except MM and *scDD-perm*, which were
 325 not considered due to the high computational cost and
 326 low statistical power displayed above). As expected, for
 327 all methods, differential patterns are easier to detect as
 328 the magnitude of the difference increases, with differ-
 329 ential location patterns having a higher true positive rate
 330 than differential scale patterns. While all methods con-
 331 trol the FDR, in all simulations, *distinct* achieves sub-
 332 stantially higher TPR than competitors (Figure 6). We
 333 also repeated the same simulations including a batch
 334 effect, with two batches, with the same scale and lo-
 335 cation differential parameters for the batch and group
 336 differences (i.e., increasing together from 0.2 to 1.5).
 337 Again, we excluded *scDD* from these analyses because
 338 it cannot handle covariates directly. Results agree with
 339 those from the *muscat* batch effect simulation study:
 340 FDR and TPRs are mostly unchanged when introduc-
 341 ing nuisance covariates, with only a minor decrease in
 342 the TPR in stronger batch effects, i.e., when *de.facLoc*
 343 and *de.facScale* are 1 and 1.5 (Supplementary Figure
 344 10), which again indicates that *distinct* can effectively
 345 control for nuisance covariates.

346 We further considered the semi-simulated mass cytom-
 347 etry data from Weber *et al.* [32] (labelled *diffcyt* sim-
 348 ulation), where spike-in signals were computationally
 349 introduced in experimental data [5], hence maintain-
 350 ing the properties of real biological data while also
 351 embedding a known ground truth signal. We evalu-
 352 ated *distinct* and two methods from *diffcyt*, based on
 353 *limma* [23] and linear mixed models (LMM), which out-
 354 performed competitors on these same data [32]. In
 355 particular, we considered three datasets from Weber

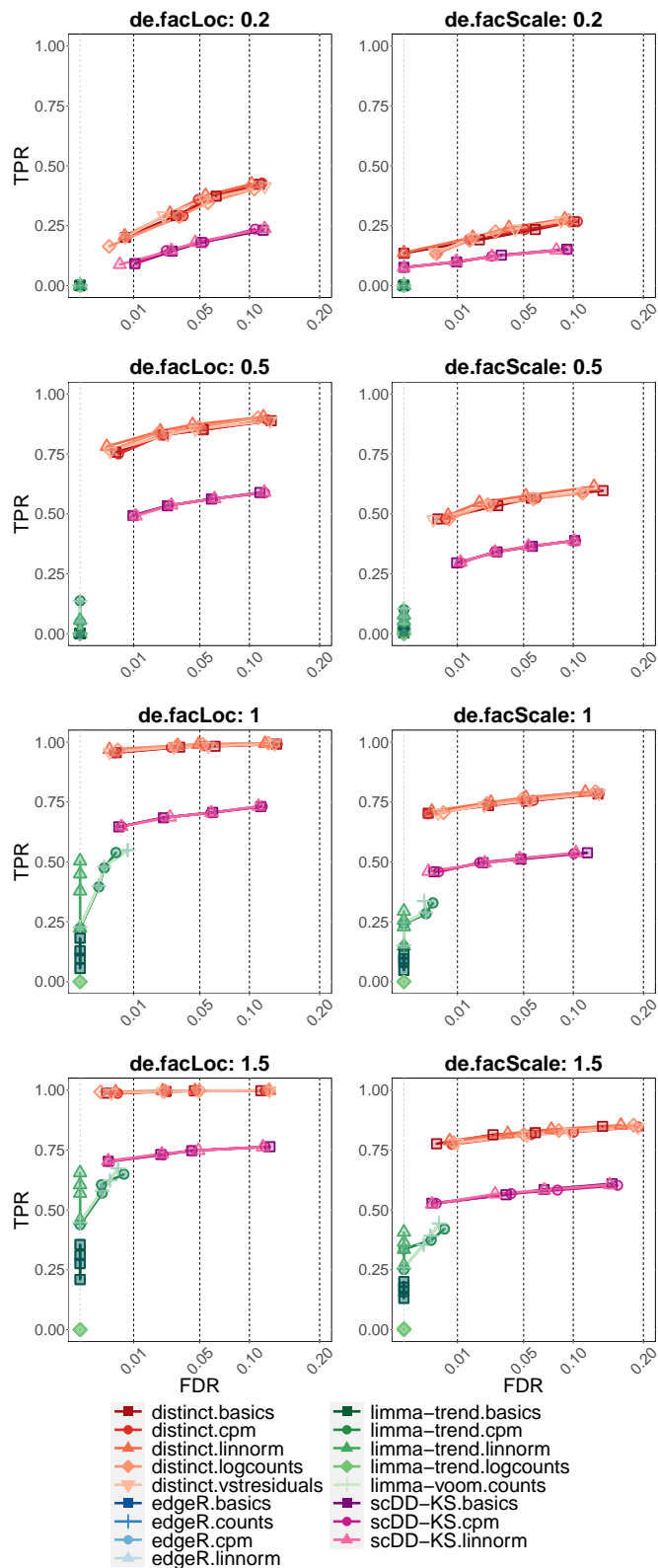


Figure 6: *distinct* displays higher TPR than competitors. TPR vs. FDR in *SplatPop* simulated data, with various degrees of differential location (left) and scale (right) parameters, primarily affecting the mean and variance, respectively. Circles indicate observed FDR for 0.01, 0.05, 0.1 and 0.2 significance thresholds. Each simulation consists of 20,345 genes, 800 cells (belonging to the same cluster), and two groups of 4 samples each, corresponding to an average of 100 cells per sample.

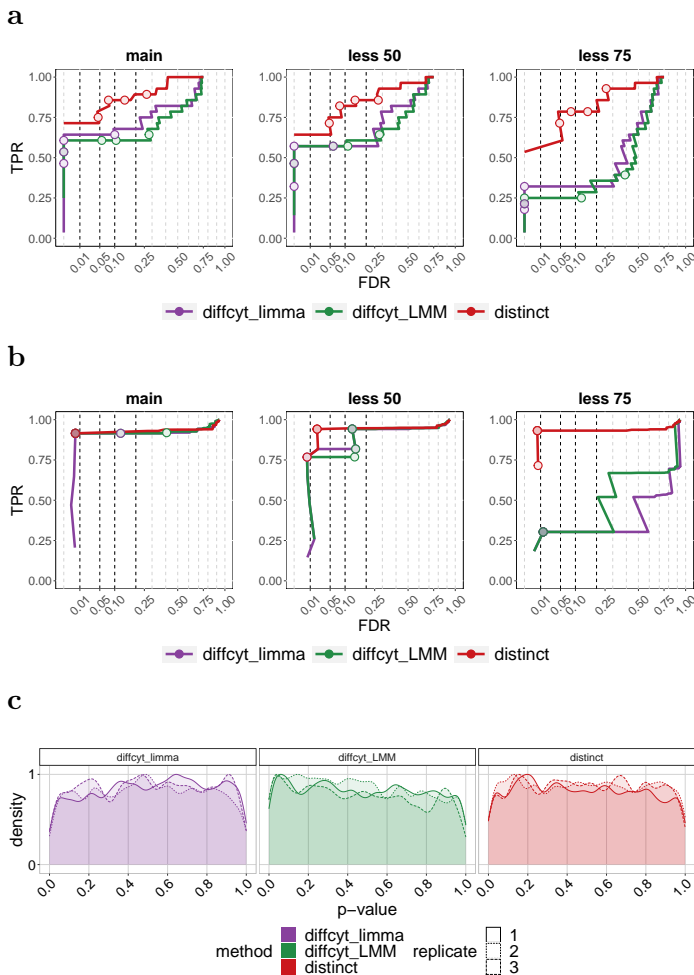


Figure 7: *distinct* shows high power while controlling for false positive and false discovery rates. (a-b) TPR vs. FDR in *diffcyt* semi-simulated data. ‘main’, ‘less 50’ and ‘less 75’ indicate the main simulation, and those where differential effects are diluted by 50 and 75%, respectively. Each simulation consists of 88,435 cells and two groups of 8 samples each. Circles indicate observed FDR for 0.01, 0.05, 0.1 and 0.2 significance thresholds. (a) As in the *muscat* simulation study, cells were clustered into 8 populations based on manually annotated cell types [32]. (b) As in Weber *et al.* [32], cells were grouped in 100 high-resolution clusters via unsupervised clustering. (c) Density of raw p-values in *diffcyt* null semi-simulated data; each replicate represents a different null simulation. Each replicate consists of 88,438 cells and two groups of 8 samples each. As in Weber *et al.* [32], cells were clustered in an unsupervised manner.

356 *et al.* [32]: the main DS dataset and two more where
 357 differential effects were diluted by 50 and 75%. Each
 358 dataset consists of 24 protein markers, 88,435 cells, and
 359 two groups (with and without spike-in signal) of 8 sam-
 360 ples each. Measurements were first transformed, and
 361 then cells were grouped into sub-populations with two
 362 separate approaches (see Methods): i) similarly to the
 363 *muscat* simulation study, cell labels were defined based
 364 on 8 manually annotated cell types [32] (Figure 7a),
 365 and ii) as in the original *diffcyt* study from Weber
 366 *et al.* [32], cells were grouped into 100 high-resolution clus-
 367 ters (based on 10 cell-type markers, see Methods) via
 368 unsupervised clustering (Figure 7b). In the main simu-

369 lation, *distinct* achieves higher TPR when considering
 370 cell-type labels (Figure 7a, ‘main’), while all methods
 371 exhibit substantially overlapping performance when using
 372 unsupervised clustering (Figure 7b, ‘main’). In both
 373 clustering approaches, as the magnitude of the differ-
 374 ential effect decreases, the distance between methods in-
 375 creases: *diffcyt* tools show a significant drop in the true
 376 positive rate whereas *distinct* maintains a higher TPR
 377 while effectively controlling for the false discovery rate
 378 (FDR) (Figures 7a-b and Supplementary Figure 11).
 379 This indicates that *distinct* has good statistical power
 380 to detect even small changes between conditions. We
 381 also considered the three replicate null datasets from
 382 Weber *et al.* [32] (i.e., with no differential effect), con-
 383 taining 24 protein markers and 88,438 cells across 8
 384 cell types, and found that all methods display approx-
 385 imately uniform p-values (Figure 7c).

386 Experimental data analyses

387 In order to investigate false positive rates (FPRs) in
 388 real data, we considered two experimental scRNA-seq
 389 datasets where no differential signals were expected, by
 390 comparing samples from the same experimental condi-
 391 tion. Given the high computational cost and low
 392 power of MM, and the high FDR of *scDD* models, for
 393 the real data analyses, we only included *distinct* and
 394 PB methods. We considered gene-cluster combinations
 395 with at least 20 non-zero cells across all samples. The
 396 first dataset (labelled *T-cells*) consists of a Smart-seq2
 397 scRNA-seq dataset of 19,875 genes and 11,138 T cells
 398 isolated from peripheral blood from 12 colorectal can-
 399 cer patients [36]. We automatically separated cells in
 400 11 clusters (via *igraph* [1, 8]), and generated replicate
 401 datasets, by randomly separating, three times, the 12
 402 patients to two groups of size 6. The second dataset
 403 (labelled *Kang*) contains 10x droplet-based scRNA-seq
 404 peripheral blood mononuclear cell data from 8 Lupus
 405 patients, before (controls) and after (stimulated) 6h-
 406 treatment with interferon- β (INF- β), a cytokine known
 407 to alter the transcriptional profile of immune cells [13].
 408 The full dataset contains 35,635 genes and 29,065 cells,
 409 which are separated (via manual annotation [13]) into 8
 410 cell types. One of the 8 patients was removed as it ap-
 411 pears to be a potential outlier (Supplementary Figures
 412 12-14). Here we only included singlet cells and cells
 413 assigned to a cell population, and considered control
 414 samples only, resulting in 11,854 cells and 10,891 genes.
 415 Again, we artificially created three replicate datasets
 416 by randomly assigning the 7 retained control samples
 417 in two groups of size 3 and 4. In both null analyses, we
 418 found that *limma-trend*, particularly when using CPMs,
 419 leads to an increase of FPRs, *distinct*’s p-values are only

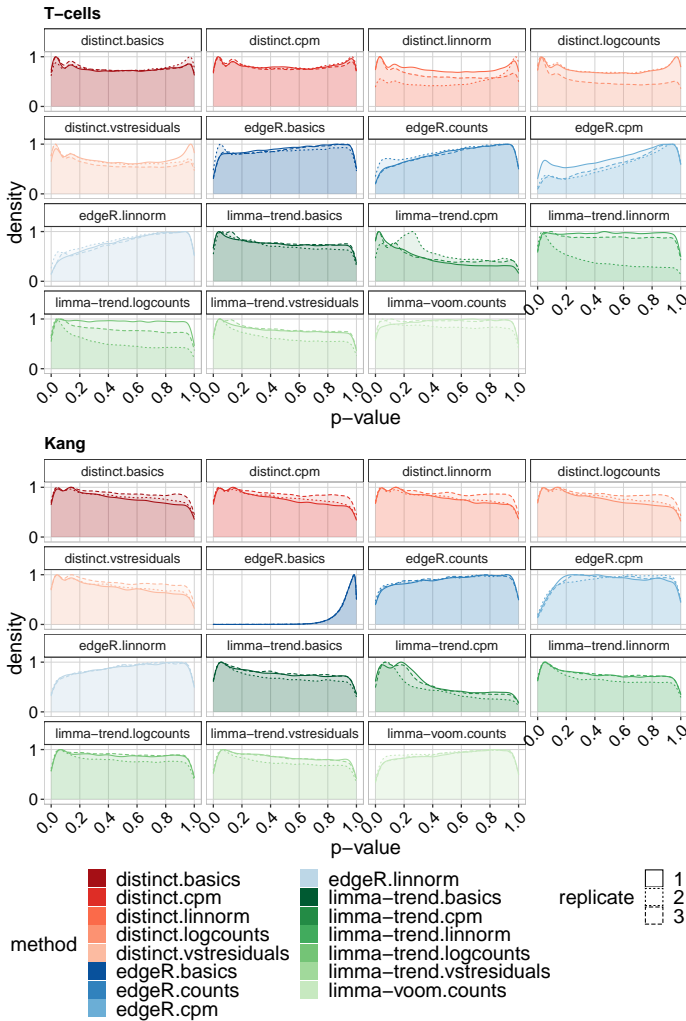


Figure 8: On experimental scRNA-seq data, *distinct* has almost-uniform null p-values. Density of raw p-values in the null *T-cells* (top) and *Kang* (bottom) experimental data. Each replicate represents a random partition of samples in two groups. The *T-cells* data consists of 12 samples and 11,138 cells across 11 clusters. For the *Kang* dataset, we retained 7 samples and 11,854 cells across 8 clusters.

marginally inflated towards 0, while *edgeR* and *limma-voom* are the most conservative methods and provide the best control of FPRs (Figure 8 and Supplementary Tables 2-3). Regarding normalization, *linnorm* and *BASiCS* lead to the most conservative p-values and smallest false positive rates.

We then considered again the *Kang* dataset, and performed a DS analysis between controls and stimulated samples. Again, we removed one potential outlier patient, and only considered singlet cells and cells assigned to a cell population; we further filtered gene-cluster combinations with less than 20 non-zero cells across all samples, resulting in 12,045 genes and 23,571 cells across 8 cell types and 14 samples. We found that *distinct* identifies more differential patterns than PB methods, with *edgeR* and *limma-voom* being the most conservative methods, and that its results are very coherent across different input data (Supplemen-

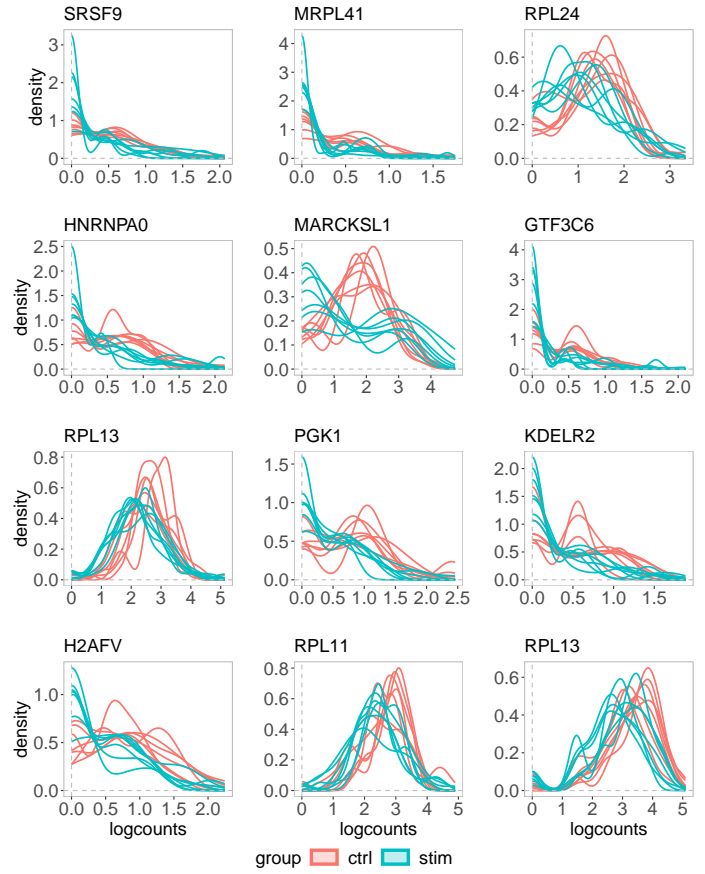


Figure 9: *distinct* discovers non-canonical differential patterns. Density of logcounts for nine examples of differential patterns identified by *distinct* on all input data (adjusted p-values < 0.05), and not by any PB tool (adjusted p-values > 0.05), on the *Kang* dataset when comparing controls and stimulated samples. Gene RPL13 was identified in FCGR3A+ Monocytes (third row) and in NK cells (fourth row), while all other genes were detected in Dendritic cells. Each line represents a sample.

tary Figure 15). When visually investigating the gene-cluster combinations detected by *distinct* (adjusted p-value < 0.1), on all five input data (CPMs, logcounts, linnorm, BASiCS and vstresiduals), and not detected by any of the ten PB approaches (adjusted p-value > 0.1), we found several interesting non-canonical differential patterns (Figure 9 and Supplementary Figures 16-27). In particular, gene MARCKSL1 displays a DB pattern, with stimulated samples having higher density on the tails and lower in the centre of the distribution, gene RPL13 mirrors classical DE, while the other genes seem to emulate DP profiles. Interestingly, ten out of eleven of these genes are known tumor prognostic markers: H2AZ2 for cervical and renal cancer, SRSF9 for liver cancer and melanoma, RPL24 for renal and thyroid cancer, HNRNPA0 for renal and pancreatic cancer, MARCKSL1 for liver and renal cancer, GTF3C6 for liver cancer, RPL13 for endometrial and renal cancer, PGK1 for breast, head and neck, cervical, liver, and pancreatic cancer, KDELR2 for renal, head

method	% of unique results
distinct.logcounts	0.3
distinct.basics	0.8
limma-trend.logcounts	0.9
distinct.cpm	1.0
distinct.vstresiduals	1.1
edgeR.linnorm	1.2
limma-trend.vstresiduals	1.5
limma-trend.basics	1.5
edgeR.counts	1.7
edgeR.basics	3.0
distinct.linnorm	3.6
limma-trend.linnorm	3.7
limma-voom.counts	5.6
edgeR.cpm	10.4
limma-trend.cpm	26.8

Table 1: Percentage of unique gene/cell-type identifications that are unique to each method. Since methods return significantly different number of significant results, for each method, we selected the most significant 1,000 results. For every method, we then compute the fraction of such results that are unique, i.e., not in common with the top 1,000 results returned by any other method.

and neck and glioma cancer, and RPL11 for renal and breast cancer [28]. This is an interesting association, considering that $\text{INF-}\beta$ stimulation is known to inhibit and interfere with tumor progression [9, 22]. Additionally, Supplementary Figures 16-27 show how *distinct* can identify differences between groups of distributions even when only a portion of the ECDF varies between conditions. Finally, we computed the fraction of detected genes that are unique by each method. Given that a ground truth is absent, we speculate that gene-cluster combinations detected by multiple methods are more likely to be truly differential, while those detected by a single method are more likely to be false positive detections. Since methods return widely different number of significant genes, for each method, we considered the top (i.e., smallest p-value) 1,000 genes per cell-type. We then computed the percentage of results that are unique to each method (Table 1), i.e., not in common with the top 1,000 results returned by any other method. Overall, *distinct* displays a lower fraction of unique results (1.4% on average across all input data) compared to *edgeR* (4%) and *limma* (6.7%). It is also interesting to note that *scater*'s logcounts normalization lead to the 2 smallest fractions of unique values (i.e., *distinct.logcounts* and *limma-trend.logcounts*).

Discussion

High-throughput single-cell data can display complex differential patterns; nonetheless, most methods for dif-

ferential expression fail to identify changes where the mean is not affected. To overcome the limitations of present differential tools, we have developed *distinct*, a novel method to identify differential patterns between groups of distributions, which is particularly well suited to perform differential analyses on high-throughput single-cell data. *distinct* is based on a flexible hierarchical multi-sample full-distribution non-parametric approach. In order to compare it to state-of-the-art differential methods, we ran extensive benchmarks on both simulated and experimental datasets from scRNA-seq and mass cytometry data, where our approach exhibits favourable performance, provides good control of the FNR and FDR, and is able to identify more patterns of differential expression compared to canonical tools, even when the overall mean is unchanged. In particular, our approach displays a higher statistical power (i.e., TPR) not only than PB methods, but also compared to other non-parametric frameworks from *scDD*, based on the Kolmogorov-Smirnov test statistic (*scDD-KS*) and on permutation tests (*scDD-perm*). *distinct* also allows for biological replicates, does not rely on asymptotic theory, which could be inaccurate in small sample sizes (typical of biological data), and avoids parametric assumptions, that may be challenging to meet in single-cell data. Additionally, *distinct* can also effectively adjust for sample-level cell-cluster specific covariates (i.e., whose effect varies across cell clusters), such as batch effects (Supplementary Figure 5). Importantly, *distinct* is a very general test that, due to its non-parametric nature, can be applied to various types of data, even beyond the single-cell applications shown here. Furthermore, thanks to its flexible form, we have shown in our simulations that *distinct* has the most consistent performance across normalization approaches (Supplementary Figure 2 and 4).

However, these advantages come at the expense of a higher computational burden, particularly when compared to PB methods or KS approaches (Figure 5). Nonetheless, by employing clever computational techniques (i.e., parallel computing and C++ coding within R), the method runs within minutes on a laptop, even for large datasets. Overall, we believe that *distinct* represents a valid alternative for differential detections from single-cell data, particularly when interest lies beyond canonical differences in means, as it allows to enhance statistical power at the cost of a reasonable increase in the computational time.

Finally, although we have focused here on comparing two groups of samples, several future extensions are possible to allow our framework to be applied to different scenarios. For instance, by suitably modifying

538 the test statistics in (1), one may ideally extend our ap-
539 proach to perform a joint differential test between three
540 of more groups of samples. Although, it is worth not-
541 ing that, in the presence of three or more experimental
542 conditions, at present, it is still possible to run pairwise
543 comparisons between pairs of conditions. While a joint
544 test across all groups may certainly be of interest in
545 some cases, from our experience, comparisons between
546 pairs of groups are usually more used among scientists.
547 In addition, as we were suggested by a user, *distinct*
548 could be employed to compare cell clusters instead of
549 experimental conditions, hence discovering differential
550 genes between cell clusters (e.g., cell types), even from
551 individual samples.

552 Availability

553 *distinct* is freely available as a Bioconductor R pack-
554 age at: <https://bioconductor.org/packages/distinct>.
555 The scripts used to run all analyses are avail-
556 able on GitHub ([https://github.com/SimoneTiberi/](https://github.com/SimoneTiberi/distinct_manuscript)
557 *distinct_manuscript*, version v3) and Zenodo (DOI:
558 10.5281/zenodo.6397114). The *diffcyt* simulated data
559 is available via FlowRepository (accession ID FR-FCM-
560 ZYL8 [32]) and *HDCytoData* R Bioconductor pack-
561 age [33]; the *Kang* dataset can be accessed via *musc-*
562 *Data* R Bioconductor package [6]; the *T-cells* dataset
563 is deposited on the European Genome-phenome (acces-
564 sion id EGAD00001003910 [36]).

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576 Author contributions

577 ST conceived the method, implemented it, performed
578 all analyses and wrote the manuscript. ST and MDR
579 designed the study. HLC and LMW contributed to
580 *muscat* and *diffcyt* simulation studies, respectively. PS
581 contributed to the computational development of *dis-*
582 *tinct* and to the revision process. All authors read,
583 contributed to, and approved the final article.

584 Competing interests

585 The authors declare no competing interests.

586 Methods

587 Permutation test

588 In order to test for differences between groups, we em-
589 ploy a hierarchical permutation approach: to estimate
590 the null distribution of s^{obs} , we permute the individual
591 observations (e.g., single-cell measurements) instead of
592 the samples. Note that this violates the exchangeability
593 assumption of permutation tests and, hence, p-values
594 are not guaranteed to be uniformly distributed under
595 the null hypothesis; nonetheless, in our simulated and
596 experimental analyses, we empirically show that *dis-*
597 *tinct* provides good control of both false positive and
598 false discovery rates. We randomly permute individual
599 observations P times across all samples and groups, by
600 retaining the original sample sizes. We denote by s_p
601 the test statistic computed at the p -th permutation,
602 $p = 1, \dots, P$. A p-value, \tilde{p} , is obtained as [21]:

$$\tilde{p} = \frac{\sum_{p=1}^P \mathbf{1}(s_p \geq s^{obs}) + 1}{P + 1}, \quad (2)$$

603 where $\mathbf{1}(cond)$ is 1 if *cond* is true, and 0 otherwise. In
604 order to accurately infer small p-values, when \tilde{p} is below
605 some pre-defined thresholds, the number of permuta-
606 tions are automatically increased and \tilde{p} is re-computed.
607 By default, *distinct* initially computes 100 permuta-
608 tions; when $\tilde{p} \leq 0.1$ these are increased to 500; when
609 the new $\tilde{p} \leq 0.01$ we use 2,000 permutations, which
610 are further increased to 10,000 if $\tilde{p} \leq 0.001$. Note that
611 the number of permutations (i.e., 100, 500, 2,000 and
612 10,000) can be specified by the user.

613 Covariates

Assume we observe Z nuisance covariates, and that N
samples are available across all groups, where for the
 i -th sample we observe C_i values (e.g., single-cell mea-
surements). We fit the following linear mixed effects
model:

$$y_c^{(i)} = \beta_0 + \sum_{z=1}^Z \beta_z X_z^{(i)} + \alpha_i + \epsilon_c^{(i)}, \text{ for } i = 1, \dots, N, \\ \text{and } c = 1, \dots, C_i, \quad (3)$$

614 where $y_c^{(i)}$ represents the c -th observation for the i -th
615 sample, β_0 is the intercept of the model, $X_z^{(i)}$ indicates
616 the z -th covariate in the i -th sample, β_z denotes the

617 fixed effect coefficient for the z -th covariate, α_i rep- 666
618 resents the random effect term for the i -th sample, 667
619 and $\epsilon_c^{(i)}$ is the (zero-mean) residual for the c -th obser- 668
620 vation in the i -th sample. We assume that random 669
621 terms are normally distributed as $\alpha_i \sim \mathcal{N}(0, \sigma_i^2)$, where 670
622 $\mathcal{N}(a, b)$ denotes the normal distribution with mean a 671
623 and variance b . Note that, due to the random effect 672
624 terms, observations from the same sample are posi- 673
625 tively correlated while, observations between different 674
626 samples are independent. We infer model parameters 675
627 via maximum likelihood, with the estimated values for 676
628 the fixed effect terms denoted by $\hat{\beta}_0, \dots, \hat{\beta}_Z$. We then 677
629 remove the estimated effect of nuisance covariates as 678
630 $y_c^{(i)} - \sum_{z=1}^Z \hat{\beta}_z X_z^{(i)}$; differential testing is performed, as 679
631 described above, on these normalized values. In DS 680
632 analyses, model (3) is fit, separately, for every gene- 681
633 cluster combination, hence accommodating for cell-type 682
634 specific effects of covariates. 683

635 Normalization

636 In scRNA-seq datasets, CPMs and logcounts were com- 687
637 puted via *scater* Bioconductor R package [19], *vstresiduals* 688
638 were calculated via *sctransform* R package [12] 689
639 (except for the *T-cells* data, where, due to a fail- 690
640 ure of *sctransform*'s variance stabilizing normalization, 691
641 we used *DESeq2*'s *vst* transformation [16]), while *lin-* 692
642 *norm*, *BASiCS* and *SCnorm*, normalized data were 693
643 calculated with the respective Bioconductor R pack- 694
644 ages [3, 10, 29, 30, 34]. For *SCnorm*, following the 695
645 author's suggestions, we normalized each cell cluster (3 in 696
646 total) separately, using samples as *Conditions* param- 697
647 eter. 698

648 In mass cytometry datasets, measurements were trans- 699
649 formed via *diffcyt*'s *transformData* function, which ap-
650 plies an *arcsinh* transformation.

651 *muscat* simulation and *Kang* data

652 In all *muscat* simulations, we used the control samples
653 of the *Kang* dataset as a anchor data; as in the real
654 data analyses, we excluded one sample as it emerged as
655 a potential outlier (Supplementary Figures 12-14), and
656 only considered singlet cells and cells assigned to a cell
657 population. In *muscat*'s simulation studies, we con-
658 sidered gene-cluster combinations with simulated ex-
659 pression mean greater than 0.2; for DB patterns, we
660 increased this threshold to 1 because with low expres-
661 sion values differences are not visible by eye. In the
662 simulation when varying the library size (Supplemen-
663 tary Figure 6), we filtered gene-clusters combinations
664 with at least 50 non-zero cells. For every simulations,
665 five replicates were simulated, and results were aver-

aged across replicates. In the main simulation (Figure
2) and the batch effect simulation (Supplementary Fig-
ure 5), we simulated from a paired design 2 groups of
3 samples each, with 4,000 genes, and 3,600 cells dis-
tributed in 3 clusters (corresponding to an average of
200 cells per sample in each cluster). For the simu-
lation study when varying the number of cells (Fig-
ure 4), the total numbers of available cells were 900,
1,800, 3,600 and 7,200, corresponding to an average of
50, 100, 200 and 400 cells per sample in every clus-
ter. For the differential simulations, we used log2-FC
values of 1 for DE, 1.5 for DP and DM, and 3 for DB
and DV. For the batch effect simulation study we used a
modified version of *muscat*, developed by Almut Luetge
at the Robinson lab (available at: https://github.com/SimoneTiberi/distinct_manuscript), which allows sim-
ulating cluster-specific batch effects [7, 17]. All *mus-*
cat simulation studies, as well as the *Kang* non-null
data analysis, were performed by editing the original
snakemake workflow from Crowell *et al.* [7]. PB meth-
ods were applied on aggregated data by summing cell-
level measurements; for differential testing, we used
muscat's *pbDS* function [7]. Mixed model methods
were implemented, via *muscat*'s *mmDS* function, us-
ing the same approaches as in Crowell *et al.* [7]: in
MM-dream2 and *MM-vstresiduals* linear mixed models
were applied to log-normalized data with observational
weights and variance-stabilized data, respectively, while
in *MM-nbinom* generalized linear mixed models were
fitted directly to raw counts. In the *muscat* simulations
and in the *Kang* non-null data analysis, we accounted
for the paired design by modelling the patient id as a
covariate in all methods that allow for covariates (i.e.,
distinct, PB and MM).

700 *splatPop* simulation

701 In *SplatPOP* simulated data, we used a hu-
702 man genome, version 19, downloaded from
703 https://www.encodegenes.org/human/release_19.html.
704 We ran a total of 16 simulations: 8 with and 8 without
705 batch effects as nuisance covariate. In each case, we
706 ran 4 differential location ("de.facLoc" parameter)
707 and 4 differential scale ("de.facScale" parameter)
708 simulations, with differential parameters equals to
709 0.2, 0.5, 1 and 1.5. In every simulation, 10% of
710 genes were differential between groups, and a total
711 of 20,345 genes and 800 cells were simulated (100
712 per sample). In the simulation with batch effects,
713 the 8 samples were randomly assigned to 2 batches,
714 and the differential location and scale parameters
715 between batches ("batch.facLoc" and "batch.facScale",
716 respectively) matched those between groups of samples

717 (“de.facLoc” and “de.facScale”). For more details on
718 how *SplatPOP*’s data is simulated, please refer to the
719 original manuscript [2] and vignettes.

720 *diffcyt* simulation

721 The *diffcyt* semi-simulated data originates from a real
722 mass cytometry dataset of healthy peripheral blood
723 mononuclear cells from two paired groups of 8 samples
724 each [5]; one group contains unstimulated cells, while
725 the other was stimulated with B cell receptor/Fc recep-
726 tor cross-linker. The original dataset contains a total
727 of 172,791 cells and 24 protein markers: 10 of these
728 are cell-type markers used for cell clustering, while 14
729 are cell state markers used for differential state anal-
730 yses; the distinction between cell state and cell-type
731 markers is based on prior biological knowledge [32].
732 In Weber *et al.* [32], semi-simulated data were gener-
733 ated by separating the cells of each unstimulated sam-
734 ple in two artificial samples; a differential signal was
735 then computationally introduced by replacing, in one
736 group, unstimulated B cells with B cells from stimu-
737 lated samples. Measurements were transformed and
738 cells clustered via *diffcyt*’s *transformData* (which ap-
739 plies an *arcsinh* transformation) and *generateClusters*
740 functions, respectively. For the DS simulation in Fig-
741 ure 7b, as in Weber *et al.* [32], we evaluated methods’
742 performance in terms of detecting DS for phosphory-
743 lated ribosomal protein S6 (pS6) in B cells, which is
744 the strongest differential signal across the cell types in
745 this dataset [20, 32]. For the DS simulation in Figure
746 7a, we considered previously manually annotated cell
747 types [32] and included all 14 cell state markers. *dif-*
748 *fcyt*’s *limma* and LMM methods were applied via *dif-*
749 *fcyt*’s *testDS_limma* and *testDS_LMM* functions, re-
750 spectively [32]. We accounted for the paired design by
751 modelling the patient id as a covariate.

752 P-values adjustment

753 All p-values were adjusted via Benjamini-Hochberg cor-
754 rection [4]. In *diffcyt* simulations we used globally ad-
755 justed p-values for all methods, i.e., p-values from all
756 clusters are jointly adjusted once. However, since PB
757 methods were found to be over-conservative when glob-
758 ally adjusting p-values [7], in *muscat* simulations and
759 *Kang* discovery analyses, we used locally adjusted p-
760 values for all methods.

761 Software versions

762 All analyses were performed via R software version
763 4.0.0, with Bioconductor packages from release 3.11.

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