SYSTEMATIC EVALUATION OF CELL-TYPE DECONVOLUTION PIPELINES FOR SEQUENCING-BASED BULK DNA METHYLOMES

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ABSTRACT

DNA methylation sequencing is becoming increasingly popular, yielding genome-wide methylome data at single-base pair resolution through the novel cost- and labor-optimized protocols. It has tremendous potential for cell-type heterogeneity analysis, particularly in tumors, due to intrinsic read-level information. Although diverse deconvolution methods were developed to infer cell-type composition based on bulk sequencing-based methylomes, their systematic evaluation has not been performed so far. Here, we thoroughly review and evaluate five previously published deconvolution methods: Bayesian epiallele detection (BED), PRISM, csmFinder + coMethy, ClubCpG and MethylPurify, together with two array-based methods, MeDeCom and Houseman as a comparison group. Sequencing-based deconvolution methods consist of two main steps, informative region selection and cell-type composition estimation. Accordingly, we individually assessed the performance of each step and demonstrated the impact of the former step upon the performance of the following one. In conclusion, we demonstrate the best method showing the highest accuracy in different samples, and infer factors affecting cell-type deconvolution performance according to the number of cell types in the mixture. We found that cell-type deconvolution performance is influenced by different factors according to the number of components in the mixture. Whereas selecting similar genomic regions to DMRs generally contributed to increasing the performance in bi-component mixtures, the uniformity of cell-type distribution showed a high correlation with the performance in five cell-type bulk analyses.

Keywords DNA methylomes, deconvolution, heterogeneity, sequencing, computational epigenetics

1 Introduction

Although single-cell analyses have been spearheading the progress in biomedicine lately, profiling of bulk samples is still in high demand for the analyses which cannot be readily accomplished using single-cell methods due to technical or cost-related reasons. For instance, single-cell methods are still laborious and costly [1] for profiling of long-time biobanked samples or large tumor patient cohorts [2][3][4]. The bulk analysis of epigenetic modifications can demonstrate epigenetic variations in large cohorts and relate those with genomic and phenotypic characteristics [5][6].

DNA methylation, particularly occurring at cytosines in CpG context in mammals, carries highly distinguishable cell-type-specific signals [7][6]. The inheritability over cell divisions and the chemical stability make it significantly more accessible for profiling. These features of DNA methylation motivated research in disease-associated cell-type-specific signals in DNA methylomes in the form of disease-specific differentially methylated regions (DMRs). To this end,
it was shown that tumor sub-types can be successfully classified based on bulk DNA methylation patterns of patient biopsies \[8\]–\[10\]. Nevertheless, cell-type-specific signals of DNA methylation can suffer from confounding factors, because DNA methylation is also associated with gender, age, various environmental influences etc. \[11\]–\[12\]. These confounding factors make cell subpopulation analysis difficult by increasing the complexity of overall methylation patterns and obscuring cell-type-specific signals.

Even though there are in vitro techniques to purify individual cell types from bulk samples, such as cell sorting or cell enrichment, other spurious variation can be introduced into the samples during the experimental process and eventually further confound cell-type-specific methylation signals. As an alternative, cell-type deconvolution, a computational approach to infer the cell-type composition of bulk samples, is applicable for cell subpopulation analysis post-experimentally. Cell-type deconvolution has been extensively utilized to dissect array-based DNA methylation bulk data, e.g. Infinium 450K/EPIC microarrays \[13\]–\[14\], that generates matrices of average methylation levels at specific loci in multiple samples (we refer to them as "array-shaped data"). Reference-based deconvolution method infers cell-type proportions based on reference methylationomes of purified cell populations. Various statistical methods and algorithms were employed to establish and fit reference-based deconvolution models including different forms of regressions \[15\]–\[16\]–\[17\]. Expectation-Maximisation (EM) algorithm and deep neural networks \[18\]–\[19\]. In contrast, reference-free deconvolution methods do not require reference data to infer the proportions of underlying cell types from methylationomes. Numerous reference-free methods have been proposed by us and others \[20\]–\[21\]–\[22\]–\[23\]–\[24\], also broadly reviewed \[25\] and compared elsewhere \[26\]. Overall, reference-based and reference-free deconvolution of bulk DNA methylationomes are well established and have been used routinely, in particular for the analysis of cellular composition in complex tumors \[27\]–\[28\]–\[29\]–\[30\]–\[16\].

More recently, DNA methylation sequencing approaches such as reduced representation bisulfite sequencing (RRBS) \[31\] or whole genome bisulfite sequencing (WGBS) \[32\] have become increasingly popular owing to dropping sequencing costs, providing much broader genome coverage \[33\]–\[34\]–\[35\] and single-molecule resolution in the form of read-level methylation calls. Furthermore, single-cell bisulfite sequencing (scBS-seq) is being used increasingly \[36\]–\[37\] exceptionally in conjunction with multi-omics analyses \[38\]–\[39\]. However, few issues still have to be carefully managed in sequencing data. Since it has high genome coverage, the multiple methylation states from reads covering respective sites has to be refined and well-summarized. In addition, sequencing data consists of read-level methylomes that each CpG site includes multiple methylation state values from all reads covering the site. This abundance of information makes sequencing data analysis more intricate by concealing some cell-type-specific signals. Consequently, dealing with sequencing data highly depends on which genomic regions are analysed and how to interpret read-level information altogether.

Read-level DNA methylationomes from sequencing data, in theory, should be more suitable and more information-rich for cell composition inference in bulk samples, compared to array-shaped data \[28\]. Diverse methods have been developed for cell-type deconvolution of sequencing-based methylome data, making use of the aforementioned advantages, foremost read-level resolution \[40\]–\[41\]–\[42\]–\[43\]–\[44\]. However, the methods published so far were tested using different data sets and variable standards. Additionally, each method requires different preprocessing procedures and generates different forms of output which might confuse less experienced users. Nevertheless, previous studies mainly assessed array-based cell-type deconvolution methods \[26\]–\[25\]. With these distinctions and rapidly growing demand of sequencing data analysis, a comprehensive, standardized and unbiased assessment of deconvolution methods targeting sequencing data has become more necessary.

In order to bridge this gap, we thoroughly compare and evaluate five previously published sequencing-based deconvolution algorithms: Bayesian epiallele detection (BED) \[40\], PRISM \[42\], MethylPurify \[43\], csmFinder + coMethy \[41\] and ClubCpG \[44\]. We evaluate their performance with respect to informative feature selection results and deconvolution accuracy under various experimental scenarios, using in silico mixtures of single-nucleus methylomes as well as realistic mixtures of tumor and normal WGBS data. Based on our analysis results, we finally propose efficient and trustworthy pipelines to deconvolve complex cell-mixture samples in different scenarios. To our knowledge, this benchmarking study is the first attempt to systematize and comprehensively evaluate the methodological developments in sequencing-based methylome deconvolution.

2 Results

2.1 Comparison of methodological design

For our benchmarking study, we chose five published sequencing-based cell-type deconvolution methods (BED, PRISM, MethylPurify, csmFinder + coMethy and ClubCpG). As a comparison group, we added two cell-type deconvolution methods for array-shaped data (reference-based constrained projection method (Houseman) and our own reference-
Figure 1: Characteristic overview of our cell-type deconvolution benchmarking. A Overall scheme of cell-type deconvolution benchmarking. We synthesized in silico cell mixtures called pseudo-bulks by mixing up reads randomly sampled from mouse neuronal scBS-seq dataset and tumor WGBS dataset respectively. Sequencing-based cell-type deconvolution methods directly take sequencing data and select informative CpG sites (informative regions) supposedly related to cell-type heterogeneity (upper row pipeline). Then, cell-type proportions are estimated from selected CpGs. For array-based cell-type deconvolution methods, we converted sequencing data into array shape with pre-identified CpGs and the methods estimate cell-type compositions from the array (bottom row pipeline). B Example of a informative region showing a cell-type-specific methylation pattern (chr1:75244319-75244379). Methylation pattern of reads (each row) overlapping with a specific region was extracted from three different samples, two pure cell-type samples (mL6-2 and mPv) and bulk_1 comprised of these cell types. Some missing methylation patterns in the figure occurred owing to the CpG sites not covered by each read. In this region, majority of reads from mL6-2 are fully methylated whereas most reads from mPv show a fully unmethylated pattern.

free method (MeDeCom)). To cope with the completely different prerequisite of input data, sequencing-based and array-based cell-type deconvolution procedures follow different pipelines in our project (Figure 1A).

All compared sequencing-based methods consist of two common steps, informative region selection and cell-type composition estimation. In the informative region selection step, the sequencing-based cell-type deconvolution methods filter out CpGs where the methylation patterns do not clearly demonstrate cell-type heterogeneity. Since sequencing data has significantly broader genome-wide coverage than array-shaped data, using all available CpG sites for deconvolution is not the most efficient strategy for cell-type composition estimation as well as escalating computational complexity. In many genomic regions, methylation patterns are identical across all reads covering the CpG sites regardless of the cell type and thus non-informative. On the other hand, some of patterns are over-complicated because of other confounding factors. Even, at such loci where read coverage is low, local cell-type population (distribution of reads in terms of cell type at the loci) usually does not correspond to global cell-type composition (cell-type composition in the entire bulk) due to the lack of reads. Informative region selection step alleviates these problems by clearing out these confounding methylation signals.

First of all this process, CpGs which are proximate to each other or overlap with a specific genomic region are grouped together. Here, we call each such group a region. After that, regions only satisfying particular conditions are selected for the next step and these selected regions are referred to as informative regions here. There are three common conditions...
**Method** | **Main data type** | **Informative region selection** | **Cell-type composition estimation**
--- | --- | --- | ---
BED | RRBS | # CpG Region Size(bp) Coverage | Class # Components Estimation Scope
ClubCpG | WGBS | >4 NA*** >20 | ref-based 2 local
csmFinder + coMethy | WGBS | >4 NA*** >10 | ref-free 2 or more global
MethylPurify | WGBS | >10** 300/200** >20 | ref-free 2 local
Prism | RRBS | >4 NA*** >20 | ref-free 2 or more local
Houseman | Methylation microarrays | CpGs overlapping with DMRs | ref-based 2 or more global
MeDeCom | Methylation microarrays | CpGs showing high methylation variance over covering reads | ref-free 2 or more global

* The original setup in [44] is 2 for minimum number of CpG and 10 for minimum read coverage.
** The original setup in [43] is 300 bp for region size and 10 for minimum read coverage, but we chose 10 for CpG number threshold. In tumor-normal pseudo-bulk analysis, we changed region size parameter value to 200 bp.
*** BED, csmFinder and Prism do not require specific region size.

Table 1: Comparison of benchmarked deconvolution methods. Sequencing-based methods have three common criteria in the informative region selection: number of CpG, region size and read coverage. Some criteria were altered to be more suitable for dataset we used in our analyses. For array-based methods, we specifically designated CpG sites based on methylation variance or DMRs. Furthermore, we described differences in cell-type composition estimation part based on three criteria again: reference-requirements, number of detectable subpopulations and estimation scope.

considered in all our benchmarked methods: number of CpGs, region size and read coverage. The respective method retains only regions with an abundant number of overlapping CpGs, sufficient region length and high read coverage. The detailed informative region selection parameter values of each method are described in Table 1.

For the two array-based methods, however, informative region selection is not a necessarily required step, since DNA methylation array-shaped data is already designed as a matrix of CpG sites by samples. Therefore, we converted our sequencing data to an array shape with pre-designated CpG sites. We described CpG site selection procedure for each array-based method in the Methods section.

With the selected informative regions (sequencing-based methods) or pre-designated CpG sites (array-based methods), cell-type deconvolution methods predict cell-type proportions within given input cell-mixture samples. This step can be characterized with three main specifications: reference requirement, number of detectable cell types/subpopulations and estimation scope (Table 1).

**Requirement of reference methylomes.** Among the benchmarked methods in this project, BED, ClubCpG and Houseman are categorized as reference-based methods. BED and Houseman require methylome profiles of pure cell type. ClubCpG fits a regression model to training cell-mixture bulk data given together with known cell-type composition.

**Number of detectable components.** We also categorized the benchmarked methods with respect to whether the method specifically targets tumor samples or not. The difference between tumor targeting and broadly applicable methods is their assumption about cell-type composition within given samples, and the methods targeting tumor is generally referred as tumor purity estimation methods. These assume that only two sub-populations comprise given cell mixtures, tumor and healthy stroma, whereas normal cell-type deconvolution method does not limit the number of subpopulations. Among the methods we considered, BED and MethylPurify were developed particularly for the analysis of tumor samples. So, we additionally tested our benchmarked methods with realistically simulated tumor-normal bulk samples, since those might be better suited to deal with abnormal methylation patterns from tumor cell types.

**Estimation scope.** Exploring all the methods, we have found a prevalent computational approach which summarizes methylation patterns across all of selected informative regions and compute the final estimates. BED, MethylPurify and PRISM locally calculate statistics in respective informative regions and predict compositions from the peak of
region-wise estimated cell-type composition distribution. On the other hand, ClubCpG, coMethy, MeDeCom and Houseman *globally* calculate final cell-type proportions directly over all selected informative regions.

<table>
<thead>
<tr>
<th>Resource</th>
<th># bulks</th>
<th>Cell types</th>
<th>Biological sample</th>
<th>Sequencing protocol</th>
</tr>
</thead>
<tbody>
<tr>
<td>2 cell-type mouse neuron</td>
<td>20</td>
<td>mL6-2, mPv</td>
<td>Mouse neuron</td>
<td>scBS-seq (Illumina HiSeq 4000)</td>
</tr>
<tr>
<td>pseudo-bulk</td>
<td></td>
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<tr>
<td>5 cell-type mouse neuron</td>
<td>20</td>
<td>mL6-2, mPv, mDL-2, mL2-3, mL5-1</td>
<td>Mouse neuron</td>
<td>scBS-seq (Illumina HiSeq 4000)</td>
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<tr>
<td>pseudo-bulk</td>
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<tr>
<td>Tumor-normal</td>
<td>20</td>
<td>Normal B cell, B cell lymphoma</td>
<td>B cell</td>
<td>WGBS (Illumina NovaSeq 6000)</td>
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<td>pseudo-bulk</td>
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Table 2: Details of pseudo-bulk datasets. We created three datasets of pseudo-bulk samples for evaluating our benchmarked methods. Two datasets are comprised of two and five cell-types of mouse neuronal single-cell BS-seq data each, and one dataset was created with tumor and normal cell types. We generated 20 pseudo-bulk samples in each dataset.

2.2 Benchmarking performance analysis study design

To estimate the influence of each step upon the final result, we comprehensively assess all methods not only with the final cell-type composition estimation results, but also with the interim results of selected informative regions for deconvolution.

Our benchmarking analysis was performed with three different datasets to test the capability of methods in varied biological scenarios, two and five cell-type mouse neuronal pseudo-bulks and tumor pseudo-bulk (Table 2). Firstly, we used single-nucleus bisulfite sequencing data of mouse neuron population from Luo et al [46] to generate *in silico* pseudo-bulk read/cell mixtures for two and five cell types. Secondly, to test methods in a realistic scenario of tumor bulk deconvolution, we created *in silico* mixtures of WGBS methylomes from normal non-cancer B-cell and B-cell lymphoma samples from Do et al [47].

In the analysis of five cell-type pseudo-bulk group, we excluded BED and MethylPurify, because tumor purity estimation methods are not eligible to detect more than two cell types. Although Prism is not a tumor purity estimation method, it failed in detecting five cell types so we excluded it from five cell-type pseudo-bulk analysis.

2.3 Evaluation of the informative region selection methods using cell type DMRs

We hypothesized that ideal informative region selection results should overlap with cell-type DMRs, i.e. genomic regions showing significant methylation pattern difference between cell types (Figure 2B). Conversely, informative region selection results with low similarity to DMRs are unlikely to supply enough cell-type-specific signals. Hence, we assessed informative region selection results primarily by comparing them with cell-type DMRs. The DMRs were generate by comparing one pure cell-type bulk to all others (details in Methods).

Three methods notably selected many more informative regions overlapping with DMRs compared to other methods in all three groups of pseudo-bulks: ClubCpG and csmFinder (Figure 2A, B and Supplementary Figure 2). For two and five cell-type mouse neuron pseudo-bulks, ClubCpG detected the largest number of overlaps, but csmFinder detected more than ClubCpG for tumor pseudo-bulks. Nonetheless Prism detected the most number of bases overlapping with cell-type DMRs in tumor pseudo-bulk analysis.

Since the size of the selected informative region set is different among methods, the number of overlaps may not exactly correspond to how similar each selected informative region set is to DMRs. We also evaluated the informative region selection results based on the proximity to DMRs in terms of genomic base. Favorov et al. designed a statistic called *genomic correlation* which measures the distributions of distances between genomic regions in two given sets [48] (details in the Methods). This statistic is [0,1] interval-bound, with lower value indicating larger distance between genomic regions from the two compared sets.

Although informative regions of ClubCpG overlapped the largest number of cell-type DMRs, their genomic correlation with each cell-type DMR set was lower than 0.2 in all analyses (Figure 2C, D and E). This implies that DMRs comprised
Figure 2: Plots for informative region selection. **A, B** Overlaps between cell-type DMRs for (A) two cell-type mouse neuronal pseudo-bulks and (B) tumor pseudo-bulks. The coloured box plots at the top of each graph shows the number of overlaps between a pair of methods connected at the middle across all pseudo-bulks, and the grey box plots at the right side mean the number of informative regions detected by each method or the number of regions in DMRs. Size of overlaps were calculated with respect to the number of bases. In each plot, different cell-type DMRs are distinguished by different colors. **C-E** Genomic correlation between DMRs and selected informative regions in (C) two cell-type mouse neuronal pseudo-bulks, (D) tumor pseudo-bulks and (E) five cell-type mouse neuronal pseudo-bulks. Higher genomic correlation means higher similarity between DMRs and selected informative regions with respect to the number of overlaps and the proximity.

a very small fraction of the informative regions selected by ClubCpG. In contrast, csmFinder and MethylPurify showed high genomic correlation with cell-type DMRs, meaning that the majority of selected informative regions by csmFinder and MethylPurify overlapped with DMRs.
2.4 Mouse neuronal single-cell pseudo-bulk cell-type deconvolution

As explained, we evaluated all methods with two groups of mouse neuronal single-cell pseudo-bulk samples. Our results showed that reference-based methods, with the exception of BED, perform better than reference-free methods in both groups (Figure 3A and B). Furthermore, we analysed predicted proportion of each cell type within each pseudo-bulk sample (Figure 3C and D). We realized that ClubCpG can produce an cell-type proportion estimate lower than 0 or higher than 1, when ground-truth proportion is relatively low or high. This is because ClubCpG does not restrict its prediction value between 0 and 1. Yet, other methods successfully made all predictions in the range of 0 and 1.

Figure 3: Cell-type composition estimation for mouse neuronal single-cell pseudo-bulk scenario. A, B Absolute error between ground-truth and estimated cell-type proportion, calculated in each sample and each cell type, for (A) two cell-type and (B) five cell-type mouse neuronal pseudo-bulks. Black line at the middle refers to the median and the both ends of bar are the first and third quantiles. Numbers above box plots indicate the median value. C, D Estimated cell-type proportions and ground-truth values (black line) ordered with respect to the ground-truth proportion, in (C) two cell-type and (D) five cell-type mouse neuronal pseudo-bulks. For two cell-type pseudo-bulks, only results of mPv cell type are shown to avoid duplicates of cell-type proportions in bi-component mixtures.

In two cell-type pseudo-bulk analysis, the best accuracy was achieved by Houseman’s method requiring pure cell-type methylome profiles. Among reference-free methods, coMethy performed best with a median absolute error value of $4.4 \times 10^{-2}$, even though MethylPurify and Prism inferred more accurate values for bulks with high proportion of mPv.

The pseudo-bulk with five cell types showed the same results as with two cell types: the lowest median absolute error was recorded by Houseman and coMethy among reference-based and reference-free methods each. In bulk-wise performances separated by cell types (Figure 3D), mL2-3 was the the most difficult cell type to estimate for coMethy, but ClubCpG and MeDeCom recorded the lowest accuracy in mL2.

Whereas Houseman exhibited lower median absolute error with five cell-type pseudo-bulks than with two cell-type pseudo-bulks, all other methods showed better performance with two cell-type pseudo-bulks. This may indicate that the Houseman’s method works better on samples with more cell types by introducing more pure cell-type methylome profiles as reference.
2.4.1 Tumor-normal cell mixture deconvolution

As described in Table 1, BED and MethylPurify are particularly designed for tumor purity estimation of tumor-normal cell mixture. For this reason, we additionally assessed the benchmarked methods using pseudo-bulk samples comprised of B cell lymphoma and normal non-cancer B cell (Figure 4).

In the reference-based setting, Houseman’s method again showed the best performance in tumor cell-type deconvolution. Compared to mouse neuronal pseudo-bulk deconvolution, Houseman’s method accomplished the lowest median absolute error of $1.7 \times 10^{-5}$ in tumor cell-type deconvolution. Among reference-free methods, MeDeCom estimated tumor pseudo-bulk cell-type compositions better than coMethy. Sample-wise performance for one cell type also showed that Houseman estimated the most accurate proportions for all samples and ClubCpG exceeded the range of 0 and 1 for extreme proportions similarly to mouse neuronal pseudo-bulk analysis results.

![Figure 4: Cell-type composition estimation results of tumor pseudo-bulk scenario. Absolute error between estimated and ground-truth cell-type proportions (left) and estimated proportions by each method and ground-truth values in order (right). All details of two plots are same as Figure 3. Again, we present the results only for normal B cell due to the symmetric cell-type proportions in bi-component mixtures.](image)

2.5 Identifying factors that influence cell-type deconvolution performance

Based on the results so far, we investigated whether informative region selection results have detectable influence upon cell-type composition estimation. We plotted the absolute error together with genomic correlation separately in pseudo-bulk samples comprised of two components (Figure 5A). Both absolute error and genomic correlation results from a method were averaged across all samples in each dataset. For two cell-type mouse neuronal pseudo-bulks, results from both cell types are included, not like cell-type composition estimation results, since these cell types generated different DMRs. Consequently, the accuracy of cell-type composition estimation (complemented of the mean absolute error value) is proportional to genomic correlation between selected informative regions and DMRs. From this result, even though all methods are designed with different algorithms, we claim that detecting CpGs overlapping with DMRs significantly contributes to the performance of cell-type deconvolution generally in the cell mixture of bi-components.

We further discovered that all four methods tested with five cell-type pseudo-bulks could infer more accurate cell-type proportions in the mixtures with more balanced cell-type composition (Figure 5B). This was determined using the entropy value which measures the uniformity of given proportions or given distribution (details in Method). The entropy of cell-type distribution was negatively correlated with the mean absolute error showing the p-value lower than 0.05 in all five cell-type deconvolution results. We presume that, in cell-type mixtures with low entropy values where cell population is extremely distributed, minor cell types may not provide enough cell-type specific signals. Also, extreme value of proportions in high entropy of cell-type distribution, very close to 0 or 1, could deteriorate the performance. Nevertheless, in both two cell-type deconvolution scenarios, not all methods performed better with higher entropy of samples. For instance, Houseman in the two cell-type pseudo-bulk analysis and coMethy, Prism in tumor pseudo-bulk analysis rather showed positive correlation between two values (Supplementary Figure 3). This might be caused by some bulks with high entropy, where cell types are more uniformly distributed, resulting in more complicated methylation patterns as similar amount of cell-type specific signals from each cell type.
Figure 5: Influential factors in cell-type deconvolution performance. A Mean absolute error vs mean genomic correlation between selected informative regions and DMRs. The points indicate results of each cell type from each deconvolution method. B Mean absolute error vs entropy of cell-type proportions in each bulk sample. In both plots, we fitted dots in a linear function (the line with grey background).

3 Discussion

Here we extensively reviewed and assessed five sequencing-based cell-type deconvolution methods with standardized measurements for unbiased evaluation. Two more array-based deconvolution methods, MeDeCom and Houseman, were also added to the analyses as a comparison group, to evaluate the effectiveness of the benchmarked methods to leverage the unique properties of sequencing data.

In order to reflect various biological scenarios, our analyses were done with two different datasets, mouse neuronal scBS-seq dataset and B cell tumor WGBS dataset. We generated pseudo-bulk samples by merging randomly sampled reads from pure cell-type samples in each dataset. For mouse neuronal pseudo-bulk samples, we generate mixtures of two and five cell types respectively to study scenario of different number of subpopulations.

All sequencing-based methods we evaluated include two essential steps. First, each method selects genomic regions which are considered to show clear cell-type-specific methylation patterns (informative region selection). Then, in the second step, final cell-type composition is inferred in the selected regions (cell-type composition estimation). Thus, we evaluated the performance separately in respective steps and finally examined the influential factors in cell-type deconvolution performance.

While evaluating informative region selection, we regarded cell-type DMRs as the gold-standard cell-type-specific loci and compared them with informative regions selected by each method. Although, in mouse neuronal pseudo-bulk analyses, ClubCpG detected the largest number of overlaps with DMRs, csmFinder showed the best genomic correlation. This is because that large number of overlaps can be caused by the large number of total selected informative regions, not by the capability of detecting cell-type-specific regions (Supplementary Figure 1).

To assess cell-type composition estimation results, we calculated the absolute error between the estimated and ground-truth proportion of each cell type in each sample. Since introducing prior knowledge from reference data ordinarily improves estimation performance, for the evaluation we grouped methods according to whether reference data is required or not. Among reference-based methods, Houseman’s method strongly outperformed other methods. In the comparison of reference-free methods, coMethy inferred the most accurate cell-type compositions in both two and five cell-type pseudo-bulks.

Cancer-associated DNA methylations are particularly aberrant and the subpopulation in cancerous tissue often brings more complicated composition with the mixture of healthy normal and tumor cells [49]. Considering the unusual situation of methylation patterns in tumor, we separately evaluated cell-type deconvolution results in tumor pseudo-bulk samples generated as described above. In informative region selection analysis, csmFinder not only showed the large number of overlaps with the DMRs, but also reached the highest genomic correlation. For cell-type composition
estimation, Houseman and MeDeCom outperformed all others in reference-based methods and in reference-free methods each.

Lastly, we have noticed the significance of selected informative regions for the overall cell-type deconvolution performance. As shown in Figure 5, mean absolute error and genomic correlation of evaluated methods have a negative correlation in bi-component pseudo-bulks. This result highlights that the more similar informative regions to cell-type DMRs a method can detect, the better performance the method would achieve in cell-type composition estimation for cell mixtures comprised of two distinct subpopulations. Yet, in five cell-type mouse neuronal pseudo-bulks, entropy of cell-type distribution showed a negative correlation with the absolute error (Figure 5). This implies that distribution of subpopulations can be more influential in cell-type deconvolution of complex bulk compositions.

Although our benchmarked methods carried out rational cell-type deconvolution results in most analyses, there are still some limitations which have to be resolved in sequencing-based cell-type deconvolution. Firstly, based on our results, sequencing-based methods did not outperform array-based methods in cell-type composition estimation. Not only Houseman’s method performed best of all reference-based methods, but also MeDeCom achieved the lowest median absolute error among reference-free methods in the tumor pseudo-bulk analysis. One of the factors making sequencing-based approach intricate is the high complexity of sequencing data itself. Sequencing-based method should be capable of eliminating redundant methylation patterns and statistically well-modeling subpopulation distribution with remaining informative methylation patterns, because sequencing data involves methylation state of each CpG site in every read unlike array data summarizing multiple methylation states at the same CpG site in one methylation beta value.

With the leverage of single-cell profiling, it was convincingly demonstrated that the variable composition of numerous normal cell types and tumor cell subclones underlies intra-tumor heterogeneity [50, 51, 52]. However, tumor purity estimation methods in our analyses could only consider tumor samples as binary mixtures of tumor and normal cell subpopulations. These results are not able to explain the actual complexity of tumor microenvironment which has been widely investigated, particularly in relation to success of various therapeutic strategies, e.g. immunotherapy [53]. Therefore, to support the upcoming tumor studies with the analysis of cell-type composition variation, more advanced tumor-targeting cell-type deconvolution methods, addressing multi-component intra-tumor heterogeneity needs to be implemented.

Last but not least, the software availability, sustainability and deployment should be considered important when cell-type deconvolution tools are implemented and released. To this end, we found that some of our benchmarked methods have significant methodological and technical limitations. For example, PRISM and BED do not ensure its feasibility to other types of data than RRBS. Furthermore, most of benchmarked methods are available only for samples aligned with Bismark [54]. The limited availability, complexity of deployment and lack of input standardization eventually hinder efficient utilization of evaluated methods for real-life analyses. In addition, software maintenance is another crucial issue to develop a sustainable cell-type deconvolution tool. Importantly, to be able to execute the tools, we had to implement multiple bugfixes. Given that bisulfite sequencing technology has been rapidly innovated recently [53, 54], sequencing-based cell-type deconvolution tools should be well maintained and updated to keep the usefulness correspondingly.

Taken together, our analysis results suggest a clear paradigm of how to conduct cell-type deconvolution for sequencing data. It will pave the way towards more accurate cell-type composition inference and more precise analysis of cell-type-specific methylation patterns to allow further method development and improvement. Because of the intrinsic benefit of read-level information, which provides detailed methylation states at each CpG, it should be possible to accomplish better accuracy in the inference of cell-type composition from sequencing-based DNA methylation data compared to summarized (array-shaped) data, something which currently available tools cannot achieve yet. For future works, it has to be more thoroughly pondered how to extract cell-type-specific signals and adjust for confounding factors affecting sequencing data. Precise cell-type deconvolution of sequencing data can broaden the range of available analyses and support a strong cell population examination tool in diversified clinical and biomedical applications.

4 Methods

4.1 Benchmarking datasets

In order to evaluate the sequencing-based deconvolution methods, we generated synthetic cell-mixture samples called pseudo-bulk. For the pseudo-bulk generation, we have chosen two different datasets, mouse brain single-cell methylCseq data [46] and B cell tumor WGBS data [37].
4.1.1 Single-cell methylome data processing

3377 single-nucleus methylomes derived from 8-week old mouse cortex tissue were downloaded from the Gene Expression Omnibus (GEO) with the accession number GSE97179. This dataset was created through high-throughput single-nucleus methylome sequencing (smnC-seq). We firstly trimmed reads twice to remove sequencing adaptors, random primer index sequence and C/T tail attached, using Adaptase with Cutadapt 2.6 [55]. Trimmed reads were aligned to the mm10 reference genome using Bismark 0.22.3 [54]. After the alignment, we sorted the result BAM files using samtools 1.9 [56] and removed duplicated reads using picard MarkDuplicates 1.141. Finally, the reads whose mapping quality is lower than 30 were filtered out with samtools 1.9 again. Parameters used in each step and details about these processes are clarified in Supplementary Table 1.

4.1.2 Tumor WGBS data processing

To generate realistic tumor-normal cell mixtures, we downloaded diffuse large B cell lymphoma and normal non-cancer B cell WGBS data from one subject each (GEO accession number GSE137880) [57]. Trimming was conducted using Trim Galore 0.6.6, then reads were aligned by Bismark 0.22.3 [54] in paired-end mode. Only reads not aligned in paired-end mode were re-aligned in single-end mode, and all reads were merged and sorted through samtools 1.9 [56]. At the end, duplicated reads were removed by picard Mark Duplicates 1.141. Detailed pipeline is clarified in Supplementary Table 1.

4.1.3 Pseudo-bulk generation

We created pseudo-bulks by merging reads randomly sampled from the aligned mouse brain single-nucleus methylome data and B cell WGBS data. The cell-type proportion for bulks were decided based on Dirichlet distribution which can mimic diverse biological scenarios. For mouse neuron pseudo-bulk, five cell types including both excitatory (mDL-2, mL2-3, mL5-1 and mL6-2) and inhibitory (mPv) neuron classes were chosen. These cell types are distinctly clustered in the two-dimensional t-SNE visualization calculated by Luo et. al. [46] and secure sufficient single-cell samples. For tumor pseudo-bulk, diffuse large B cell lymphoma and normal B cell cell-types were mixed. We used generateExample function from R package MeDeCom (https://rdrr.io/github/lutsik/MeDeCom/src/R/utilities.R) to generate the cell-type proportions for pseudo-bulk samples. We followed all default parameter setup with the exception of 10 for proportion.var.factor and 1 million for number of genomic features.

4.2 Differentially methylated regions

As gold-standard to be compared with informative region selection results showing cell-type-specific features, we extracted differentially methylated regions (DMRs). DMRs refer to genomic regions whose methylation states are different across given multiple samples. It is also known to be associated with the cell development and the cell reprogramming stages [58].

For mouse neuronal cell types, we compared each of 11 pure cell-type bulks (mL4, mL6-1, mL6-2, mDL-2, mL5-1, mL5-2, mL2-3, mSst-1, mNdnf-2, mPv and mVip) with all other bulks and identified DMRs respectively. For tumor analysis, we called DMRs between normal B cell non-cancer and diffuse large B cell lymphoma. All DMRs were called using DSS package 2.34.0 [59] with the following parameters: 0.2 for delta, 0.05 for threshold of p-values, 4 for minimum number of CpG sites and default values for minimum length and the distance to merge, which are 50 bps each.

4.3 CpG selection scheme for array-based deconvolution methods

Array-shaped methylome data comes in the matrix form, CpG sites by samples. To apply array-based deconvolution methods, Houseman and MeDeCom, as a comparison group, we transformed our sequencing data to array shape using methrix [60]. When converting the sequencing data, we specified a set of CpG sites to comprise the array shape. During data conversion for MeDeCom, we chose top 20000 CpGs with the highest beta-value variance. CpGs overlapping cell-type-specific DMRs were taken to generate array-shaped data for Houseman’s method to adhere to the reference-based scenario.

4.4 Parameter values and procedures for each deconvolution method

We describe the detailed algorithm and parameter setting of each deconvolution method in this section. The pipelines are also available at https://github.com/CompEpigen/SeqDeconv_Pipeline.git.
4.4.1 ClubCpG

ClubCpG \cite{44} clusters reads fully covering informative regions satisfying given informative region selection conditions, using density-based spatial clustering of applications with noise (DBSCAN). For our experiments, we applied informative region selection conditions as given in Table 1 which suited our dataset than the default values suggested by authors. Even though ClubCpG package itself does not have cell-type composition estimation function, we followed the cell-type deconvolution strategy proposed in \cite{44}. Firstly, we created ClubCpG clustering results from another 100 pseudo-bulk samples as training data and extracted 20 principal components (PCs) using PCA algorithm from the result. Then, a multivariate linear regression model for cell-type proportion was fitted on the extracted 20 PCs. Finally, the cell-type composition was estimated using the trained multivariate linear regression model.

4.4.2 PRISM

PRISM \cite{42} infers the composition of epigenetically distinct subpopulations in tumor bulk samples based on methylation patterns. It mainly improves the accuracy by correcting erroneous methylation pattern using Hidden Markov Model. After the correction, the method remains only loci comprised of fully methylated and unmethylated patterns, then cell-type proportions are estimated on the loci using EM algorithm. We applied PRISM with default setup as given in Table 1 and yielded cell-type proportions in respective samples by calculating the ratio of inferred subclones.

4.4.3 MethylPurify

MethylPurify \cite{43} adopts EM algorithm to estimate tumor purity in bisulfite sequencing data. The EM algorithm in MethylPurify not only estimates methylation level of subpopulations, but also decides which subpopulation each read would be assigned to over iterations. We used the same parameter values in both mouse neuronal and tumor pseudo-bulks, 10 for read coverage and 50 for sampling time. However, bin length parameter was set to 300 bp for mouse neuronal pseudo-bulks and to 200 bp for tumor pseudo-bulks.

4.4.4 csmFinder + coMethy

Yin et al. developed two distinct computational tools called csmFinder and coMethy \cite{41}. csmFinder determines genomic regions showing cell-type specific signals denoted as putative cell-type-specific methylated (pCSM) loci. coMethy can decompose methylation level matrix of pCSM loci by samples, using NMF-approach similar to MeDeCom. For the specific input file format of csmFinder, we extracted methylation call results from our pseudo-bulk samples via \texttt{bismark\_methylation\_extractor} with \texttt{comprehensive}, \texttt{gzip} and \texttt{cytosine\_report} options. CpG site coordinates of hg19 and mm10 genomes were used for corresponding samples. Running csmFinder, we followed the default parameters, minimum methylation difference of hypo- and hyper-methylation patterns 0.3 and p-value of the difference 0.05. Since coMethy works on a matrix which consists methylation patterns at same CpG sites from multiple samples, we collected pCSM loci detected from all samples in each experiment and filtered out only loci involving missing methylation values in any other sample than the sample that the locus was detected.

4.4.5 BED

BED algorithm \cite{40} is based on a Bayesian model and recognizes the distribution of epialleles which indicates all possible methylation patterns at Cpg sites in a specific genomic range. For the preprocessing step to deal with contiguous and missing methylation patterns, we used pipelines elucidated in bed-beta Github page (\url{https://github.com/james-e-barrett/bed-beta}). This pipeline includes all epiallele estimation processes. Barret et al. demonstrated that the proportion of reads which are not attributed to normal tissue at each loci $i$ can be calculated as follow:

$$
\zeta_i = \frac{1}{2} \sum_q |\phi_q - n_q|
$$

This equation is calculated with respect to all epialleles $q$. $\phi$ and $n$ mean the distribution of epialleles at the locus and normal tissue samples each. We considered the rightmost maximum value in $\zeta_i$ distribution graph to be the estimated tumor purity as this is where the local epiallele distributions and normal tissue epiallele distributions showed the biggest discrepancy.

4.4.6 MeDeCom

MeDeCom \cite{20} estimates cell-type proportions in array-shaped DNA methylation data based on non-negative matrix factorization (NMF). When converting our sequencing-based data to array shape, we used \texttt{methrix} which is capable of
loading bedGraph file format including methylation call and creates a CpG sites by samples matrix of methylation levels [60]. A bedGraph file for each pseudo-bulk sample was created through a computational tool called MethylDackel (github.com/dpryan79/MethylDackel). For reference genome to read bedGraph files in, mm10 and hg19 genomes were used for mouse neuronal and B cell tumor pseudo-bulks each. As an input matrix of MeDeCom, we selected 20,000 CpG sites with the largest methylation level variance over covering reads. We ran MeDeCom with a regularization parameter $\lambda$ from $10^{-5}$ to $10^{-2}$, 10 cross-validation folds, maximum iteration number 500 and random initialization number 30.

4.4.7 Houseman’s method

Houseman’s method [15] (or Houseman short) was proposed to infer cell-type distribution from DNA methylation array-shaped data based on regression calibration. We used array-shaped data converted from our sequencing pseudo-bulk samples processed in the same way as for MeDeCom above. For selecting informative CpG sites, we mimicked the marker-CpG selection step of the original algorithm and chose CpG sites overlapping with DMRs which can provide cell-type specific signals. We increased the number of CpG sites to 1000 in the original code considering the larger number of total CpG sites in our pseudo-bulk dataset compared to microarray data they used. For the rest, we followed up the pipeline given in the supplementary material of the publication [15].

4.5 Performance measurement

4.5.1 Genomic correlation

We used genomic correlation to calculate overall genomic base-wise proximity between informative region selection results and DMRs. This statistic was suggested by Favorov et al. for determining the distribution of distances between two sets of genomic regions [48].

Relative distance of a selected informative region $q_i$ with respect to given DMRs is defined as below:

$$\delta_i = \min(|q_i - r_k|, |r_{k+1} - q_i|); k = \arg \min_{q_i > r_k} (q_i - r_k)$$

(2)

$r_k$ and $r_{k+1}$ are two nearest DMRs from given informative region $q_i$. It is calculated by dividing the distance between selected informative region $q_i$ and the closest DMR by the distance between two nearest DMRs.

If a set of selected informative regions and DMRs are independent, $\delta_i$ will have a uniform distribution. Hence, they test if the distribution of calculated relative distances makes a uniform distribution.

$$GenomicCorr = \frac{\int_0^{1/2} |ECDF(\delta) - ECDF_{\text{ideal}}(\delta)| d\delta}{\int_0^{1/2} ECDF_{\text{ideal}}(\delta) d\delta}$$

(3)

$ECDF$ refers to empirical distribution cumulative function and $ECDF(\delta)$ creates the distribution of observed $\delta_i$. $ECDF_{\text{ideal}}$ is the uniform distribution. So, Equation 3 yields the ratio of difference between area under $ECDF(\delta)$ and under $ECDF_{\text{ideal}}$. When given selected informative regions are independent of DMRs, $ECDF(\delta)$ will create a uniform distribution and $GenomicCorr$ will reach 0. Conversely, if selected regions are identical to DMRs, $GenomicCorr$ is 1.

4.5.2 Absolute error between estimate and ground-truth

In cell-type composition estimation analysis, we assessed the performance by calculating absolute error between estimated proportion and ground-truth value of each bulk sample. For coMethy and MeDeCom, we had to match cell types to estimated components by choosing the one with the lowest mean absolute error out of all possible combinations of estimate and ground-truth pair.

4.5.3 Entropy of cell-type distribution

In information theory, entropy is a concept to describe the level of uncertainty or information in a set. We used this measurement to determine how equally cell types are distributed in the cell mixture, because cell-type composition
estimation can be more intractable with extremely low or high proportion of subpopulations. Applying entropy to our cell mixture, entropy of cell-type proportion in the cell mixture $C$ comprised of $n$ cell types, $c_1, c_2, ..., c_n$ is defined as:

$$H(C) = -\sum_{i=1}^{n} P(c_i) \log P(c_i) \tag{4}$$

$P(c_i)$ refers to the cell-type proportion of cell type $c_i$ here. Thus, the entropy value is higher when all cell-type proportions are more uniformly distributed.

For example, let us assume we have two cell mixtures $A$ and $B$ comprised of bi-components with different proportions. The dominant cell type takes apart of 90% in cell mixture $A$ and 60% in cell mixture $B$. According to Equation 4, the entropy of cell mixture $A$ is smaller than cell mixture $B$.

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References


