Hybrid Bayesian-rank integration approach improves the predictive power of genomic dataset aggregation

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ABSTRACT

Motivation: Modern molecular technologies allow the collection of large amounts of high-throughput data on the functional attributes of genes. Often multiple technologies and study designs are used to address the same biological question such as which genes are over-expressed in a specific disease state. Consequently, there is considerable interest in methods that can integrate across datasets to present a unified set of predictions.

Results: An important aspect of data integration is being able to account for the fact that datasets may differ in how accurately they capture the biological signal of interest. While many methods to address this problem exist, they always rely on external standards or internal statistics, which may not always be available. We present a new rank aggregation method for data integration that requires neither external standards nor internal statistics but relies on Bayesian reasoning to assess dataset relevance. We demonstrate that our method outperforms established techniques and significantly improves the predictive power of rank-based aggregations. We show that our method, which does not require an external gold standard, provides reliable estimates of dataset relevance and allows the same set of data to be integrated differently depending on the specific signal of interest.

Availability: The method is implemented in R and is freely available at http://www.pitt.edu/~mchikina/BIRRA/

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Supplementary information: Supplementary data are available at Bioinformatics online.

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1 INTRODUCTION

The availability of high-throughput experimental technologies has greatly increased the rate of data generation. However, the reliability of individual high-throughput datasets is often uncertain, and the development of techniques to integrate the results across different studies to improve the validity and reproducibility of findings has been an important area of research.

In samples studying disease-related expression changes, for example, sample size is a common limitation, and data integration by statistical meta-analysis has been used repeatedly to boost statistical power and find reliable signals. Formal meta-analyses rely on the statistical properties of each independent dataset, and common strategies involve combining effect sizes and $P$-values [see (Tseng et al., 2012) for a comprehensive review]. However, the fundamental limitation of this approach is that statistical properties may not correlate with how well a particular biological question is addressed. A single study may show large effect sizes and small $P$-values but can be biologically invalid owing to design flaws. For example, a study purporting to find large expression differences in ethnic groups (Spielman et al., 2007) was later shown to be confounded by batch effect (Akley et al., 2007).

An alternative data integration approach that is robust to discordance between statistical power and biological validity is Bayesian integration with an external gold standard. This approach does not use any internal dataset statistics and has the additional advantage that it can be used to perform integration tasks that cannot even in principle be addressed with statistical meta analysis, such as integrating data from completely unrelated experimental platforms. In the Bayesian integration approach, the accuracy of each dataset is measured in terms of how well it is able to reproduce known information in the gold standard. For example, in integrating genomic datasets for functional network predictions, each dataset is evaluated against a gold standard of functionally interacting proteins (Huttenhower et al., 2009; Lee et al., 2004). In this case, datasets only contribute a set of final predictions (which may be categorical or real valued) whose accuracy is judged independently of the underlying data structure.

The limitation of the Bayesian approach is that it requires the existence of an acceptable gold standard against which individual predictions are evaluated. For some integration tasks, such as disease gene expression studies, such a standard is typically not available. Moreover, even well-established standards of knowledge, such as the Gene Ontology or KEGG, are only limited to capturing accuracy relative to what is already known. Thus, if a dataset captures correct but completely novel biological information, its value is not recognized by the comparison with a gold standard lacking any sampling of this signal, and its contribution will be discarded.

A practical alternative to formal statistical meta-analysis or Bayesian analysis is combining datasets directly without any treatment of the underlying statistics or any reference to the external standard. Such methods can be applied in a variety of contexts where statistics and external standards may be unavailable or prone to producing misleading quality assessments. In this integration approach, each dataset contributes equally to the consensus signature. Without loss of generality, the approach...
can be viewed as an instance of rank aggregation where each dataset supplies a prediction list (that is not necessarily complete) in order of differential expression, correlation or some other metric of relevance. Unfortunately treating all datasets equally can make these methods unacceptably sensitive to noise and outliers.

To address these limitations, we propose a new data integration method that combines the rank aggregation framework with the power of Bayesian reasoning. Our method assumes that there is a ground truth to be discovered and this is represented with varying fidelity in the datasets to be integrated. Starting with an equally weighted consensus generated by rank averaging, we iteratively fit Bayes Factors to each dataset and recompute the integration result. Thus, this approach combines the advantages of being agnostic to dataset-specific statistics and external standards while using the robustness of the Bayesian framework to handle outliers and inconsistencies. We demonstrate that this approach has favorable theoretical properties on an established simulation pipe-line and improves over other rank aggregation methods in real data integration tasks. The method is computationally simple, yet significantly increases the predictive power of rank-based aggregations. Additionally it provides a reliable estimate of dataset relevance and can be used to produce context-sensitive integrations that focus in on specific areas of interest. Because it requires only ranked lists, it is readily applicable to many integration problems.

2 APPROACH

Our method is inspired by Latent Class Analysis (LCA), a statistical technique widely used in social science and medical diagnostics to analyze a series of related measurements by assuming that they capture an underlying latent trait. For example, in the absence of definitive knowledge regarding disease status, LCA can be used to infer the sensitivities and specificities of a series of diagnostic tests. In essence, the technique works by consensus, if multiple diagnostic tests report a positive result the individual is more likely to have the disease. The entire combination of diagnostic test-specific false-positive and false-negative rates and the disease status probability of individual patients can thus be estimated by maximum likelihood.

While the underlying problem appears to be similar to genomic data integration and a related approach has been successful in integrating binary interaction networks (Weile et al., 2012), the methodology is not directly transferable to arbitrary genomic data. LCA operates on categorical data, while genomic data are usually continuous valued. Moreover, the assumption of the existence of a few discrete latent classes is often incorrect and genomic datasets can be extremely discordant, resulting in many solutions with similar likelihoods. For example, in a case of large disagreements, solutions dictating that one dataset is right and all others are wrong are all equally valid. We address these limitations with several heuristic techniques, which give good results in practice. Our methods works by iteratively computing an aggregated ranking from dataset-specific Bayes factors and using the ranking to update the Bayes factor calculation. We refer to our method as Bayesian Iterative Robust Rank Aggregation (or BIRRA).

While the method can be applied to any data integration task, for the purpose of illustration we can assume that the input to our method is a set of datasets that rank genes in order of disease relevance, for example, a set of differential expression studies. If we had a representative set of genes that were known to be differentially expressed in the disease, we could infer the probability of disease relevance (DR) for the rest of the genes given the set of evidence (gene’s ranks within a dataset) \( E_{1,n} \) via the standard Bayesian approach

$$P(DR|E_1, E_2, \ldots, E_n) = C \cdot \prod_{i=1}^{n} P(E_i|DR)$$

Where \( C \) is a normalization constant and \( P(E_i|DR) \) is calculated from the set of known positive examples. This approach naturally captures the fact that individual datasets vary in how well they are able to discriminate disease-relevant genes and their relative contribution to the final ranking should be adjusted accordingly. Our method uses the same principle without relying on a set of positive examples to calculate \( P(E_i|DR) \) but rather estimating it from the structure of the data.

**Require:** input data to be integrated for \( n \) genes, rank ordered by some measure of relevance (such as differential expression \( P \)-values or correlation). Missing values in each dataset are assigned the lowest rank \( P \) a priori probability for the ‘positive class’ \( B \) number of bins.

1: Compute initial aggregate ranking by mean ranks
2: Compile a working standard by assigning the top \( np \) of aggregated predictions to be the positive class and the rest of the predictions to the negative class
3: Discretize the data into \( B \) bins
4: repeat
5: Compute bin-wise cumulative Bayes factors for each dataset by comparing with the working standard
6: Aggregate the data by naive Bayesian integration according to Equation 1
7: Update working standard with new rankings obtained in step 6
8: until rankings unchanged or max iterations exceeded
9: return Final ranking

In principle, for complex genomic datasets, this kind of procedure can be unstable because in cases of large disagreements or low signal, multiple solutions are equally valid. We implement several heuristic techniques to make this approach more robust. In particular, we compute Bayes factors cumulatively starting from the top bin and enforce a monotone decrease. Thus, the minimum Bayes factor is limited by the prior probability, and each piece of data can only contribute positive evidence. Secondly, we use a winzorization step whereby for each bin we decrease the maximum Bayes factor to that of the second maximum. This ensures that at least two datasets are always combined and prevents the appearance of singleton solutions whereby only one datasets is predicted to have any signal.
While we find that our experiments with real data were robust to variations in bin size, we intentionally leave this as free parameters because the optimal choice is likely to be data dependent. Because the algorithm proceeds entirely by bootstrapping any noise in Bayes factor, estimation may be propagated and thus choice of bin size must strike a balance between using all the available signal while minimizing the influence of random fluctuations. We find that for integrations of data from mammalian genomic datasets, a choice of 50 bins was appropriate. Our implementation includes a plotting functionality that tracks the progress of the Bayes factor estimates (as in Fig. 2) and can be used to investigate the influence of different binning. We also find that in all our analyses, using real data convergence was achieved in ≤10 steps (which is the default number of maximum iterations).

3 RESULTS

3.1 Simulations

While using a Bayesian framework, our approach is still most accurately classified as rank aggregation because it requires only ranked lists and the conditional probabilities are bootstrapped from the data itself. Rank aggregation methods are attractive data integration tools because they require no prior knowledge and are thus widely applicable and easy to implement. However, they are fundamentally limited by their ability to handle noise and outliers because there is no mechanism to prevent a large number of noisy datasets from overwhelming the results. We expect that the Bayesian framework should perform well in this regard and we evaluate the theoretical properties of our method with a simulation strategy similar to that outlined in Kolde et al. (2012).

We simulate a differential expression meta-analysis with 1000 genes where 5% of the genes are truly differentially expressed and the differential expression is assayed in independent experiments. To create a single dataset, we draw values form a $N(1,1)$ distribution for the differentially expressed genes, and for the remaining 95% of background genes, we drawn values from a $N(0,1)$ distribution. The final values are subsequently rank transformed. Using these distribution parameters produces individual datasets that rank the true differentially expressed genes with an average Area Under the Curve (AUC) of 0.75 and we refer to these as ‘signal’ datasets. We also introduce ‘noise’ datasets where the genes are ranked randomly.

We use the complete set of signal and noise datasets to evaluate different rank aggregation approaches [mean ranks, Robust Rank Aggregation (RRA) (Kolde et al., 2012), Stuart (Stuart et al., 2003), and our method BIRRA]. Aggregating 10 signal and 30 noise datasets with several different methods we find that even on a dataset with 75% uninformative data all methods tested produce results that outperform individual signal datasets (Fig. 1) and the computationally sophisticated RRA and Stuart methods surpass mean ranks at low sensitivity. However, RRA outputs a P-value of 1 for almost 80% of the genes and thus has no discriminative power at higher recall, producing a total AUC that is lower than mean ranks. Overall, the more sophisticated methods improve only slightly if at all over the simple mean ranks, demonstrating that these methods provide little additional robustness. The BIRRA method, on the other hand, is specifically designed to address robustness to noise, as it explicitly estimates the information content of individual datasets down-weighting the uninformative datasets directly.

Figure 2 shows a sample run of the BIRRA method that highlights this key advantage of our approach. We plot the estimated log Bayes factors against bin number for each iteration of our algorithm and we observe that the pure noise datasets (gray lines) and signal datasets (red lines) separate quickly, with the noise datasets converging to lower Bayes factors. Of course, if we had a representative gold standard to evaluate the conditional probabilities, we would find that the noise datasets contain no information and the corresponding log Bayes factors should be 0. While in the absence of an independent standard the noise datasets are not discounted completely, they are still significantly down-weighted, producing a result whose performance falls between various single-step aggregation methods and optimal naive Bayes (Fig. 1).

Varying the number of non-informative datasets we find that BIRRA maintains an advantage across a large range of added noise and produces the most improvement for integration tasks that contain between 50 and 85% uninformative datasets (Fig. 3). BIRRA also performs comparably with other methods for values outside this range. When little or no noise is present,
We have compiled a set of 14 related gene expression datasets. In these cases, it is difficult to evaluate performance, as typically no universally accepted standards of disease-related expression changes exist. However, one important source of validation can be reproducibility, and we can use this criterion to evaluate our method. We have compiled a set of 14 related gene expression datasets that compare tissue from Parkinson’s disease (PD) patients with controls. Given these datasets we can ask how well the results of a single one of them can be predicted by integrating the remaining 13. We evaluate our method alongside mean ranks and the more sophisticated RRA and Stuart rank aggregation methods.

Figure 4 shows that for a number of datasets their results could not be predicted by any of the rank aggregation methods, demonstrating that they capture entirely different signals. However, for those datasets for which mean rank aggregation successfully captures significant signal, BIRRA was able to improve on that result in 9 of 11 cases, producing the best overall performance of the four methods tested. Importantly, the two datasets that were better recapitulated with mean ranks than with BIRRA were difficult to predict with all methods, and the performance difference was small; on the other hand, the datasets for which integration was a good predictor (>0.7 AUC in mean ranks), our method produced a substantial improvement. We also investigated how well the relative ranking within the top 500 genes was predicted by different aggregation approaches and find that BIRRA showed an improvement on this metric as well (Supplementary Fig. S1).

An additional advantage of our BIRRA approach is that it not only outputs aggregated results but also the computed Bayes factors, which can serve as an alternative measure of dataset quality because these values capture how well each dataset reproduces the consensus signal. When datasets are combined by BIRRA, these Bayes factors determine the relative contribution of each. On the other hand, in formal statistical meta-analyses, the relative contribution is determined by the statistical properties of the underlying data. It is natural to ask how the statistical quantities that are intrinsic to each dataset and the BIRRA...
quality estimates that are computed from the consensus are related. While there is no single metric that can describe dataset quality in a statistical sense, we focus on two parameters that would be of relevance in various meta-analysis approaches. Common meta-analysis strategies include directly merging raw data and combining effect sizes or P-values (Tseng et al., 2012). Thus, one relevant parameter is the sample size of the study, which is an a priori measure of the expected statistical power. We also choose the maximum P-value for the top 50 differentially expressed genes to represent the actual observed magnitude of the overall disease effect. Plotting the sum of bin-wise Bayes factors against these statistical quantities we find that there is no relationship between these measures of intrinsic dataset quality and quality as judged by replication (Fig. 5).

For example, the dataset with the seemingly largest effect (GSE22491) had a poor BIRRA quality score, revealing that it produced differential expression rankings that were not supported by other datasets, while for the dataset that BIRRA judged to be most informative (GSE20291) the P-value for the first 50 DE genes was greater than would be expected from a uniform distribution. This analysis demonstrates the great discordance between intrinsic statistical properties and the ability of datasets to capture reproducible biological signal and underscores the potential pitfalls of relying exclusively on dataset statistics when performing meta-analysis.

### 3.3 Predicting transcription factor-target networks

To evaluate how well our method performs in a real data integration task for which an external source of validation is available, we follow the approach described in Kolde et al. (2012). The goal of this integration task is to predict transcription factor (TF) targets from correlation between the TFs and other genes in several gene expression datasets.

We perform the analysis for 14 TFs involved in embryonic stem-cell (ES) renewal and assayed by ChIPseq in Chen et al. (2008) and 12 ES related microarray datasets. We compute correlations between the TF gene and all other genes in each dataset, and the resulting ranked lists serve as the input to various rank aggregation methods. The individual datasets and results of four aggregation approaches (mean ranks, BIRRA, RRA, and Stuart) are evaluated for correctness against the ChIPseq dataset that directly measures physical binding (Fig. 6). We find that on this integration task, both the Stuart and RRA methods typically outperformed mean ranks but not the best individual dataset. On the other hand, our method BIRRA outperformed the best dataset in 10 of the 14 cases, a substantial improvement over alternative rank aggregation techniques. Importantly, BIRRA showed the largest improvement in cases where the individual datasets varied most with respect to how well they were able to capture the relevant signal, for example, Myc and Smad1, demonstrating that BIRRA correctly identifies the more relevant datasets without relying on an external gold standard.

Using the independent gold standard of ChIPseq interactions we can not only evaluate the predictive power of our final rank aggregation but also the estimates of dataset quality that are a byproduct of our approach. We compare the BIRRA-estimated Bayes factors with the values obtained by directly plugging in the true-positive and false-positive counts obtained from the ChIPseq gold standard (as in standard naive Bayes analysis).

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**Fig. 5.** Evaluating BIRRA estimated dataset relevance against dataset statistical properties. We plot the sum of log Bayes factors against maximum P-value of top 50 DE genes (A) and study sample size (B). We find that the BIRRA-estimated quality is unrelated to dataset intrinsic statistical properties.

**Fig. 6.** TF targets predicted from expression correlation. We aggregate the results of 10 different ES expression datasets to predict TF target interaction and evaluate the result using a ChIPseq dataset. We find that for most TFs tested BIRRA aggregation produced outperformed other methods and the best individual dataset.
that the BIRRA method outperforms established approaches on individual datasets, weighing them accordingly. We have shown aggregation approaches in that it directly estimates the reliability of Bayesian reasoning. Our method differs from previous rank aggregation methods that can only output a single result, BIRRA works depending on the TFs of interest. Thus, unlike other rank complete correlation network, our method produces different networks between TFs and other genes, we can use the BIRRA-computed Bayes factors were consistent in each case.

We plot the dataset-specific Bayes factors derived from the two methods (BIRRA and naive Bayes) side by side in Figure 7 and find that the relative dataset quality as judged by the self-contained BIRRA approach is in fairly good agreement with that obtained by using an independent standard. Importantly the true dataset quality (as estimated by direct comparison with ChIPseq data) varied for different TFs, and BIRRA was able to correctly identify which specific datasets were informative in each case. For example, based on comparisons with the gold standard, GSE4308 (black dashed lines) performs well at predicting targets for Myc, yet is completely uninformative for Smad1, and this trend if fully captured by BIRRA (Fig. 7).

While the ChIPseq-based validation relies only on the correlations between TFs and other genes, we can use the BIRRA-computed Bayes factors to produce integrations over all gene pair correlations. Importantly, however, focusing on a specific TF produces different relevance estimates for the same dataset, and the relative contribution of each dataset to the final result varies substantially (Fig. 7). Consequently, when extended to the complete correlation network, our method produces different networks depending on the TFs of interest. Thus, unlike other rank integration methods that can only output a single result, BIRRA can produce context-sensitive rank aggregations that maximize the agreement over a specific subset of the biological signal.

**5 METHODS**

**5.1 Rank aggregation methods**

The BIRRA method was programmed in R, and the source code is available in the supplement. The RRA and Stuart methods were called from the RobustRankAggreg R package.

**5.2 TF target networks**

Data for 14 ES-related studies were downloaded from InSilicoDB (Coletta et al., 2012), which provides Robust Multi-array Average (RMA)-processed Affymetrix files mapped to gene symbols. As in (Kolde et al., 2012) we created a standard for TF target interaction using the ChIPseq-determined TF-promoter mapping provided by (Chen et al., 2008). Following (Kolde et al., 2012), we considered all genes with a score of ≥0.6 as targets. We noted that for two of the TFs considered in the ChIPseq study, Smad1 and Suz12, the majority of individual expression datasets were predictive of their targets only when anti-correlation was considered, suggesting that these TFs act as repressors. Suz12 is known to be directly involved in gene repression (Cao and Zhang, 2004), and though Smad1 is not a dedicated repressor, in ESs it interacts with Nanog to repress its targets (Suzuki et al., 2006). Therefore, for these two TFs, we ranked genes in order of anti-correlation. For the remaining TFs, correlation was used.

**5.3 PD meta-analysis**

We collected the majority of the datasets used in a previously reported Parkinson’s meta-analysis (Zheng et al., 2010). For Affymetrix datasets, we downloaded the data from InSilicoDB. For Illumina datasets, the data was taken from the Gene Expression Omnibus database (GEO) (Edgar et al., 2002) and processed with background correction using the ‘bg.adjust’ function in the ‘affy’ Bioconductor package, log transformation and quantile normalization. Datasets were filtered by discarding the bottom 20% of low expression and low variance genes, and processed for differential expression with respect to disease status, including additional covariates, such as gender and brain region, whenever those were applicable and available in GEO.

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REFERENCES


