Machine learning methods for rare diseases

This manuscript (permalink) was automatically generated from jaybee84/ml-in-rd@d2b651c on July 29, 2020.

Authors

• Jineta Banerjee

(D 0000-0002-1775-3645 ⋅ **(7** jaybee84

Sage Bionetworks · Funded by Neurofibromatosis Therapeutic Acceleration Program; Children's Tumor Foundation

Robert J Allaway

© 0000-0003-3573-3565 · ♠ allaway · ❤ allawayr

Sage Bionetworks · Funded by Neurofibromatosis Therapeutic Acceleration Program; Children's Tumor Foundation

Jaclyn N Taroni

© 0000-0003-4734-4508 · ♥ jaclyn-taroni

Childhood Cancer Data Lab, Alex's Lemonade Stand Foundation

Casey Greene

© 0000-0001-8713-9213 · ♥ cgreene

Department of Systems Pharmacology and Translational Therapeutics, Perelman School of Medicine, University of Pennsylvania; Childhood Cancer Data Lab, Alex's Lemonade Stand Foundation

Justin Guinney

© 0000-0003-1477-1888 · ♥ jguinney

 $Sage\ Bionetworks\cdot Funded\ by\ Neurofibromatosis\ The rapeutic\ Acceleration\ Program;\ Children's\ Tumor\ Foundation$

Synopsis

(Instructions: Describe the background, basic structure of the article, list material to be covered indicating depth of coverage, how they are logically arranged, include recent pubs in the area, 300-500 words)

Substantial technological advances have dramatically changed biomedicine by making deep characterization of patient samples routine. These technologies provide a rich portrait of genes, cellular pathways, and cell types involved in complex phenotypes. Machine learning is often a perfect fit for the types of data now being generated, and Nature Methods routinely has reports of machine learning methods that extract disease-relevant patterns from these high dimensional datasets. Often, these methods require a large number of samples to identify reproducible and biologically meaningful patterns. With rare diseases, biological specimens and consequently data, are limited due to the rarity of the condition. In this perspective, we outline the challenges and emerging solutions for using machine learning in these settings. We aim to spur the development of powerful machine learning techniques for rare diseases. We also note that precision medicine presents a similar challenge, in which a common disease is partitioned into small subsets of patients with shared etiologies and treatment strategies. Advances from rare disease research are likely to be highly informative for other applications as well.

Introduction

Machine learning is gaining momentum in biomedical data analysis as data collection is increasingly high-throughput and algorithms or approaches become more transparent and interpretable.

Application of machine learning to any dataset poses challenges, but the application to biomedical data and subsequent interpretation requires depth of knowledge not only in the biomedical domain but also a clear understanding of the methods and their underlying assumptions.

Rare disease research has not yet significantly benefited from machine learning applications for various reasons, including lack of statistical power in dataset size, heterogeneity in available data, and sensitivity of machine learning methods to misinterpretation in view of small datasets. We anticipate higher occurrence of such applications in the near future and aim to highlight the current state-of-art in this perspective. However recent advances in the methodologies to accommodate rarity of samples and increased transparency in model outputs have encouraged application of machine learning in rare disease.

Application of machine learning to any kind of data consists of the following major steps: (1) data evaluation and question formulation, (2) selection of normalization/dimension reduction to mitigate technical differences, (3) selection of appropriate algorithms which select features to answer the formulated question, (4) evaluation of the answers generated by the algorithm. Each of these steps require the practitioner to choose from a variety of methodologies to apply. The selection of the methodologies at each of these steps need to be based upon robust reasoning to ensure stability of the results. Moreover, in the context of rare diseases, special considerations need to be made at each of the above mentioned steps to safeguard against misinterpretation of data. Such considerations include incorporation of techniques that build upon prior domain-specific knowledge, methods that are resilient to challenges posed by small datasets, and methods that can mitigate technical disparities in the data.

Techniques that build on prior knowledge and indirectly related data are necessary for many rare disease applications

This section will highlight promising approaches for analyzing rare disease data to extract biological insights. We will discuss techniques like transfer learning, representation learning, cascade learning, integrative analysis, and knowledge-graph creation and use that leverage other knowledge and data sources to construct testable hypotheses from rare diseases datasets with limited sample sizes.

Ensemble Learning

Implementing machine learning on data with low sample size and high label uncertainty can lead to unstable predictions. In such cases where single predictors fail, various machine learning methods together or *ensemble learning* may help increase accuracy of prediction. *Ensemble learning* methods like random forests use bootstrap aggregation (or bagging) of independent decision trees that use similar parameters but different paths for the selection of features to form a consensus about the important predictive features[1]. However, successful application of consensus based ensemble learning requires "gold standard" data where the diagnosis or label of a data point in the training dataset has very little uncertainty (or "label-noise") associated with it [6]. In most cases of rare disease, due to the inherent nature of being less defined, the symptoms as well as any underlying biology comes with a reasonable amount of uncertainty (or "label-noise") leading to a silver standard dataset[7]. In such cases, ensemble learning with multiple methods leveraging distinct underlying assumptions are used in tandem to capture stable patterns existing in the silver standard data and reduce uncertainty. Such cascade learning classifiers have been widely used in image recognition where initially a small subset of image features are used to classify images (e.g. features identifying a face like eyes, nose, mouth). The initial classification is then augmented by more complex features and algorithms like AdaBoost (boosting) that weight the various features implemented to detect the content of the image (e.g. features identifying a human face like relative distance between eyes etc.) [<u>8</u>].

In rare diseases, a variant of *cascade learning* that showed robustness in view of uncertainty in the data was implemented to identify rare disease patients from electronic health records from the general population [11]. This implementation consisted of three steps each employing a different independent learning algorithm: (1) feature extraction to assign text words (from Pubmed literature) to diagnosis using word2vec [12], (2) preliminary prediction using an ensemble of decision trees with penalization for excessive tree-depth, (3) prediction refinement using similarity of data points to resolve sample labels and reiterating step (2). In this implementation the algorithm was able to identify rare disease patients due to the robustness conferred by the independence of the feature extraction step and the prediction refinement step from the preliminary classification of the labeled dataset. The classification step capitalized upon the information learned by the label prediction step preceding it and the prediction refinement step following it, and was able to perform better over other ensemble methods when implemented on silver standard data.

Most cascade classifiers follow *one-classifier-at-a-time* approach where algorithms at each level predict all classes involved. But scenarios where the need for high prediction accuracy for one class outweighs other classes (e.g. malignant tumor-types, or severe psychiatric cases) require further modification of the cascade learning efforts. An example of this was seen implemented for triaging psychiatric patients where the identification of one class of psychiatric patients ("severe") far outweighed the need for optimized overall classification accuracy[13]. Due to the requirements of the problem, they developed a *one-class-at-a-time* approach for cascade learning, where at each stage a binary classifier is used to predict a specific class against all others. The final model implemented all models together each identifying one class sequentially and the final prediction was the union of the

predictions at all the different models. The cascade classifiers using the *one-class-at-a-time* approach were found to perform better than multi-class ensemble classifiers in most cases.

Thus ensemble learning can be helpful in producing stable predictions from data that is limited in quality or quantity, where single algorithms would otherwise produce unstable predictions. However, the choice of using *bagging*, *boosting*, independent algorithmic steps, or *one-class-at-a-time* approach would strictly depend on the nature of the prediction problem. In most cases involving rare disease data, it seems that *bagging* has had limited success, which has necessitated various modifications of the approaches as discussed above.

Techniques and procedures must be implemented to manage model complexity without sacrificing the value of machine learning

Inherent challenges posed by low sample numbers in rare diseases are further aggravated by disease heterogeneity, poorly defined disease phenotypes, and often a lack of control (i.e. normal) data. Machine learning approaches must be carefully designed to address these challenges. We discuss how to implement methodological solutions like bootstrapping sample data, regularization methods for deep learning, and hyper-ensemble techniques to minimize misinterpretation of the data.

Bootstrapping

Bootstrap or resampling computation is a powerful statistical technique that can be used for estimating population values from datasets of limited sample size [14]. The technique utilizes random sampling of data points from a dataset of limited sample size with replacement to approximate a larger population and estimate various population statistics (e.g. mean). Subsequent iterations of resampling generates a distribution of the statistical value (mean) which minimizes the error of the estimate. Bootstrap based techniques are used in conjunction with various learning methods to find the most informative models given a specific dataset (e.g. bootstrap aggregating or bagging used in random forests [2,15], bootstrap in neural networks [16], or regression models [17,18]).

While most datasets in practice are of finite sample size and can benefit from bootstrapping, rare disease datasets with limited number of samples necessitate the use of bootstrap to form an informative dataset in addition to model selection [19]. In this study, bootstrapping the training sample without replacement simulated formation of different incomplete datasets that helped expose the learning models (in this case random forests) to the incompleteness of the data. Such additional bootstrapping of the training data helped create confidence intervals for the predictions and the important predictors originating from unstable ensemble models run on the incomplete training data

Regularization

Machine learning algorithms are optimized to find patterns among data points and prioritizes the strongest patterns that exist in a dataset. Given a limited dataset with strong pre-existing technical differences between groups of samples, this optimization may lead to the model learning technical differences thus lowering its predictive accuracy [20]. For example, in a set of 1000 samples where 700 samples are from one healthcare site and 300 from another, it is likely that there will remain site-specific differences between them even after normalization of the samples. If the site-specific differences are more pronounced than the underlying patterns differentiating the samples, any machine learning model trained with these data will preferentially learn the site-specific differences to classify the samples, and rank them higher than the underlying patterns leading to a model showing high prediction accuracy of training data (termed low bias in model). When new test data points are introduced to the model, possibly coming from a third site, the model is unable to locate the earlier differences in the new data points and fails to classify them accurately causing a significant drop in

accuracy of the model (termed high variance in model prediction). Such a model is termed "overfit" to its training data. Overfitting can lead to misinterpretation of the site-specific differences as true patterns in the limited data points and thus needs to be minimized. Minimization of overfitting can be accomplished by cross-validation and regularization methodologies.

While cross-validation aims to reduce the variance in prediction, regularization adds a small amount of bias to the initial model to minimize its dependence and sensitivity to training data. Regularization makes models less reliant on training data by adding a penalty (determined by cross-validation), and then minimizes the error between the model's prediction and ground truth of the test data. Regularization can not only minimize overfitting but can additionally help in predicting outcomes using a limited number of samples.

Regularization can be of three main types, each with their particular strengths and weaknesses. (1) Ridge regression aims to minimize the magnitude of the features, but in models that try to select the most important features for accurate prediction of sample labels, ridge regression shrinks all features equally, but cannot completely remove unimportant features. Thus in presence of many correlated parameters (e.g. gene expression networks), ridge regression may not be ideal in reducing the feature space. (2) LASSO or least absolute shrinkage and selection operator regression on the other hand works well for selecting few important features since its effect can minimize the magnitude of some features more than the others. Thus it helps in selecting most important features while the magnitude of irrelevant features are shrunk to 0 and eventually removed. This selection attribute of LASSO (in a sample set of size "n", LASSO can select "n" features for the model) may be an advantage in reducing model complexity, but a disadvantage in cases where identification of all possible collinear features is important (e.g. all biomarkers correlating to a particular disease phenotype) [21]. (3) Elastic-Net regression is a combination of LASSO and ridge regression[22]. Both of the methodologies when applied together helps to select most useful features, specially where there are a lot of correlated features. In this setup, LASSO leads to selection of one of the correlated features and reduces the others to 0 (grouping of features), and the magnitude of the selected features are then minimized through ridge regression.

Any supervised learning implementation in rare disease would require robustness towards feature selection from a small number of samples, i.e. the features selected by a model as important should be stable in view of new data points added to a dataset, even though their relative importance may change due to additional evidence. This robustness is mostly acquired through the combination of various regression strategies. Since machine learning applications in rare disease are infrequent, combination strategies used for rare variant discovery and immune cell signature discovery can serve as good case studies to examine. Many deleterious genomic variants can be extremely rare due to the constant selection pressure working against them. Since the frequency of a rare variant is so low (less than 1%) applying routine statistical procedures that were extensively developed for common variant association, to analyze a low minor allele frequency (MAF) seem inappropriate [23]. For its feature selection attribute, LASSO has been widely applied in microarray and GWAS data for common variants. But since LASSO by itself is too stringent for rare variants, it has been employed along with group penalties to help identify rare variants/ low frequency predictors [24]. Variations of LASSO have also been implemented to aggregate or group the occurrence of rare variants together by gene or chromosome location [25,26,27]. In this strategy, a 0-1 dummy variable was created for each SNP based on the presence or absence of the rare variant. Then linear combinations of the selected dummy variables were considered by using the LASSO procedure. Even though most of the dummy variables were 0, their linear combination was more likely to be nonzero thus leading to increased signal to noise ratio for the rare variants. Only those linear combinations that were non-zero in at least 5% of the subjects were then included to ensure that the new markers were not rare [26,28]. While ridge regression is not usually utilized for feature selection, adaptive ridge regression has been utilized to help combine rare variants into a single score analogous to feature engineering for increasing the signal of rare variants[29]. Another variation of LASSO included its integration with the

probabilistic logistic bayesian approach to identify a protective rare variant in lung cancer[30]. Xu et al. on the other hand combined the feature selection methods with a generalized pooling strategy, and evaluated the performance of these hybrid approaches for detection of rare genetic variants[31]. Another interesting approach is the sparse-group LASSO approach which incorporates prior knowledge into the regularization[32]. This approach works well for a scenario where only few genes in a pathway are true predictors of a phenotype, where it helps select the driving genes in a pathway of interest.

Alternatively, Elastic-net regression (a combination of LASSO and ridge regression) has also been used to reduce the feature space in various types of cancer datasets [33,34]. In cases where the number of features were far greater than the number of samples, elastic-net has usually been found to outperform the other regression approaches [22]. A variation of the elastic-net regression was used for identifying immune cell signatures in an RNA-seq dataset where the number of cells sampled were far fewer than number of genes profiled [??? 10.1186/s12859-019-2994-z]. This two-step regularized logistic regression technique included a pre-filtering phase to select the optimal number of genes and then implemented elastic-net regularization for gene selection. The second step generated gene signatures for individual cell types using selected genes from first step and then implemented a binary regularized logistic regression for each cell type against all other samples

Still to add: techniques in deep learning e.g. Deep and shallow architecture: https://ieeexplore.ieee.org/document/7863293

Techniques to manage disparities in data generation are required to power robust analyses in rare diseases

As with common diseases, genomic and transcriptomic data from rare diseases can suffer from artifacts introduced by batch, processing methodology, sequencing platform, specimen/data quality or other non-biological phenomena. The consequences of these non-biological artifacts are amplified in rare diseases which often have few samples and heterogeneous phenotypes. Furthermore, because datasets are many times pieced together from multiple small studies, in which disease phenotypes or other important biological characteristics are often confounded by the previously mentioned "batch" factors. A key consideration here is, if possible, active dialogue with the data generators or experts in the field who may have unexpected insights into potential sources of variation. One example of the value of this, experienced by the authors, occurred when studying tumors associated with the disease neurofibromatosis type 1. These datasets were, unbeknownst to the computational biologists, generated from samples obtained with vastly different surgical techniques (laser ablation and excision vs standard excision), resulting in substantial biological differences that are a consequence of process, not reality. One might expect, in this example, that this technical decision would result in profound changes in the underlying biology, such as the activation of heat shock protein related pathways, unfolded protein responses, and so on. Consequently, careful assessment of confounding factors and implementation of normalization methods is important to identifying biologically meaningful features within a dataset. Assessment of confounding factors and heterogeneity in rare disease datasets is perhaps most easily performed using unsupervised learning approaches such as clustering and dimensionality reduction. Clustering methods like k-means clustering or hierarchical clustering can be used to characterize the structure present in many different types of data such as genomic or imaging data. [35,36]. Similarly, a variety of dimensionality reduction methods are can be used to visualize sample heterogeneity and potential confounding variables, including multidimensional scaling (MDS), principal components analysis (PCA), t-distributed stochastic neighbor embedding (t-SNE), and uniform manifold approximation and projection (UMAP), among many others. [37,38,39,40] All of these methods can be effectively used to identify batch effects and other structure in the data, though some, like t-SNE and UMAP, have parameters, such as perplexity (number of nearest neighbors), that can substantially affect the output, and thus the interpretation, of the analysis [40,41]. Therefore, successful application of these methods requires a

sufficient understanding of the underlying method and parameter sweeping to get a clear picture of the structure of the underlying data. Another important consideration is that, as discussed by Way, et. al. [42], a single dimensionality reduction method alone may not be sufficient to reveal all of the technical or biological heterogeneity; testing multiple methods may result in a more comprehensive portrait of the data Dimensionality reduction techniques are not restricted to 'omic' data - they can also be used in rare disease applications to characterize the structure and heterogeneity of imaging data [43], mass cytometry data [44], and others. Once the nature of the non-biological heterogeneity has been established, different techniques can be used to correct the differences. Common approaches to ameliorate non-biological effects include the assessment of data quality using robust metrics, reprocessing the raw data using a single analysis pipeline if the data are obtained from different sources, application of batch correction methods [45,46], normalization of raw values (e.g. zscores, trimmed mean of M-values [47]). It can also be helpful to be fatalistic, in some sense, when working with rare disease data. For various reasons including ethical considerations, limited funding, and limited biospecimen availability, experimental design and the resulting data will be less-than-ideal - for example - when batch variables and biological variables are confounded. In these cases, it may be prudent to take a step back, re-evaluate the data, and identify methods that can operate within these constraints.

Conclusions

We will conclude by discussing the potential of the above-mentioned approaches in rare diseases and other biomedical areas where data is scarce.

draft

this is a test file to differentiate draft-branch from master

References

1. Evaluating predictive modeling algorithms to assess patient eligibility for clinical trials from routine data

Felix Köpcke, Dorota Lubgan, Rainer Fietkau, Axel Scholler, Carla Nau, Michael Stürzl, Roland Croner, Hans-Ulrich Prokosch, Dennis Toddenroth

BMC Medical Informatics and Decision Making (2013-12-09) https://doi.org/f5jqvh

DOI: <u>10.1186/1472-6947-13-134</u> · PMID: <u>24321610</u> · PMCID: <u>PMC4029400</u>

2.:{unav)

Leo Breiman

Machine Learning (2001) https://doi.org/d8zjwq

DOI: 10.1023/a:1010933404324

3. Analyzing bagging

Peter Bühlmann, Bin Yu

The Annals of Statistics (2002-08) https://doi.org/btmtjp

DOI: 10.1214/aos/1031689014

4. Utilising artificial intelligence to determine patients at risk of a rare disease: idiopathic pulmonary arterial hypertension

David G. Kiely, Orla Doyle, Edmund Drage, Harvey Jenner, Valentina Salvatelli, Flora A. Daniels, John Rigg, Claude Schmitt, Yevgeniy Samyshkin, Allan Lawrie, Rito Bergemann

Pulmonary Circulation (2019-11-20) https://doi.org/gg4jc7

DOI: <u>10.1177/2045894019890549</u> · PMID: <u>31798836</u> · PMCID: <u>PMC6868581</u>

5. Double-bagging: combining classifiers by bootstrap aggregation

Torsten Hothorn, Berthold Lausen

Pattern Recognition (2003-06) https://doi.org/btzfh6

DOI: <u>10.1016/s0031-3203(02)00169-3</u>

6. Learning statistical models of phenotypes using noisy labeled training data

Vibhu Agarwal, Tanya Podchiyska, Juan M Banda, Veena Goel, Tiffany I Leung, Evan P Minty, Timothy E Sweeney, Elsie Gyang, Nigam H Shah

Journal of the American Medical Informatics Association (2016-11) https://doi.org/f9bxf9

DOI: 10.1093/jamia/ocw028 · PMID: 27174893 · PMCID: PMC5070523

7. Classification in the Presence of Label Noise: A Survey

Benoit Frenay, Michel Verleysen

IEEE Transactions on Neural Networks and Learning Systems (2014-05) https://doi.org/f5zdgg

DOI: <u>10.1109/tnnls.2013.2292894</u> · PMID: <u>248</u>08033

8. Component-based face detection

B. Heiselet, T. Serre, M. Pontil, T. Poggio

Institute of Electrical and Electronics Engineers (IEEE) (2005-08-25) https://doi.org/c89p2b

DOI: <u>10.1109/cvpr.2001.990537</u>

9. The Architecture of the Face and Eyes Detection System Based on Cascade Classifiers

Andrzej Kasinski, Adam Schmidt

Advances in Soft Computing (2007) https://doi.org/cbzq9n

DOI: 10.1007/978-3-540-75175-5 16

10. Real time facial expression recognition with AdaBoost

Yubo Wang, Haizhou Ai, Bo Wu, Chang Huang
Institute of Electrical and Electronics Engineers (IEEE) (2004) https://doi.org/crv3sq

DOI: 10.1109/icpr.2004.1334680

11. Learning to Identify Rare Disease Patients from Electronic Health Records.

Rich Colbaugh, Kristin Glass, Christopher Rudolf, Mike Tremblay Volv Global Lausanne Switzerland *AMIA ... Annual Symposium proceedings. AMIA Symposium* (2018-12-05)

https://www.ncbi.nlm.nih.gov/pubmed/30815073

PMID: 30815073 · PMCID: PMC6371307

12. Efficient Estimation of Word Representations in Vector Space

Tomas Mikolov, Kai Chen, Greg Corrado, Jeffrey Dean *arXiv* (2013-01-16) https://arxiv.org/abs/1301.3781v3

13. Machine learning for psychiatric patient triaging: an investigation of cascading classifiers.

Vivek Kumar Singh, Utkarsh Shrivastava, Lina Bouayad, Balaji Padmanabhan, Anna Ialynytchev, Susan K Schultz

Journal of the American Medical Informatics Association: JAMIA (2018-11-01)

https://www.ncbi.nlm.nih.gov/pubmed/30380082

DOI: 10.1093/jamia/ocy109 · PMID: 30380082 · PMCID: PMC6213089

14. Improvements on Cross-Validation: The 632+ Bootstrap Method

Bradley Efron, Robert Tibshirani

Journal of the American Statistical Association (1997-06) https://doi.org/gfts5c

DOI: <u>10.1080/01621459.1997.10474007</u>

15. Bootstrap Methods for Developing Predictive Models

Peter C Austin, Jack V Tu

The American Statistician (2004-05) https://doi.org/bzjjxt

DOI: <u>10.1198/0003130043277</u>

16. Bootstrap for neural model selection

Riadh Kallel, Marie Cottrell, Vincent Vigneron

Neurocomputing (2002-10) https://doi.org/c8xpqz

DOI: <u>10.1016/s0925-2312(01)00650-6</u>

17. Fast bootstrap methodology for regression model selection

A. Lendasse, G. Simon, V. Wertz, M. Verleysen

Neurocomputing (2005-03) https://doi.org/dx5c3p

DOI: 10.1016/j.neucom.2004.11.017

18. A bootstrap resampling procedure for model building: Application to the cox regression model

Willi Sauerbrei, Martin Schumacher

Statistics in Medicine (1992) https://doi.org/cnpg3d

DOI: 10.1002/sim.4780111607 · PMID: 1293671

19. Integrative Analysis Identifies Candidate Tumor Microenvironment and Intracellular Signaling Pathways that Define Tumor Heterogeneity in NF1

Jineta Banerjee, Robert J Allaway, Jaclyn N Taroni, Aaron Baker, Xiaochun Zhang, Chang In Moon, Christine A Pratilas, Jaishri O Blakeley, Justin Guinney, Angela Hirbe, ... Sara JC Gosline

Genes (2020-02-21) https://doi.org/gg4rbj

DOI: 10.3390/genes11020226 · PMID: 32098059 · PMCID: PMC7073563

20. Definitions, methods, and applications in interpretable machine learning

W. James Murdoch, Chandan Singh, Karl Kumbier, Reza Abbasi-Asl, Bin Yu *Proceedings of the National Academy of Sciences* (2019-10-29) https://doi.org/ggbhmq

DOI: 10.1073/pnas.1900654116 · PMID: 31619572 · PMCID: PMC6825274

21. Regularization

Jake Lever, Martin Krzywinski, Naomi Altman *Nature Methods* (2016-09-29) https://doi.org/gf3zrr

DOI: 10.1038/nmeth.4014

22. Regularization and variable selection via the elastic net

Hui Zou, Trevor Hastie

Journal of the Royal Statistical Society: Series B (Statistical Methodology) (2005-04)

https://doi.org/b8cwwr

DOI: <u>10.1111/j.1467-9868.2005.00503.x</u>

23. Statistical analysis strategies for association studies involving rare variants

Vikas Bansal, Ondrej Libiger, Ali Torkamani, Nicholas J. Schork *Nature Reviews Genetics* (2010-10-13) https://doi.org/dn4jtz

DOI: 10.1038/nrg2867 · PMID: 20940738 · PMCID: PMC3743540

24. Association screening of common and rare genetic variants by penalized regression

H. Zhou, M. E. Sehl, J. S. Sinsheimer, K. Lange

Bioinformatics (2010-08-06) https://doi.org/c7ndkx

DOI: 10.1093/bioinformatics/btq448 · PMID: 20693321 · PMCID: PMC3025646

25. Identification of Grouped Rare and Common Variants via Penalized Logistic Regression

Kristin L. Ayers, Heather J. Cordell

Genetic Epidemiology (2013-09) https://doi.org/f5cw72

DOI: 10.1002/gepi.21746 · PMID: 23836590 · PMCID: PMC3842118

26. A LASSO-based approach to analyzing rare variants in genetic association studies

Jennifer S Brennan, Yunxiao He, Rose Calixte, Epiphanie Nyirabahizi, Yuan Jiang, Heping Zhang *BMC Proceedings* (2011-11-29) https://doi.org/bjcndj

DOI: <u>10.1186/1753-6561-5-s9-s100</u> · PMID: <u>22373373</u> · PMCID: <u>PMC3287823</u>

27. Methods for Detecting Associations with Rare Variants for Common Diseases: Application to Analysis of Sequence Data

Bingshan Li, Suzanne M. Leal

The American Journal of Human Genetics (2008-09) https://doi.org/d4jpcb

DOI: 10.1016/j.ajhg.2008.06.024 · PMID: 18691683 · PMCID: PMC2842185

28. Comparison of statistical approaches to rare variant analysis for quantitative traits

Han Chen, Audrey E Hendricks, Yansong Cheng, Adrienne L Cupples, Josée Dupuis, Ching-Ti Liu *BMC Proceedings* (2011-11-29) https://doi.org/b9mf4x

DOI: <u>10.1186/1753-6561-5-s9-s113</u> · PMID: <u>22373209</u> · PMCID: <u>PMC3287837</u>

29. Adaptive Ridge Regression for Rare Variant Detection

Haimao Zhan, Shizhong Xu

PLoS ONE (2012-08-28) https://doi.org/f36tm5

DOI: 10.1371/journal.pone.0044173 · PMID: 22952918 · PMCID: PMC3429469

30. An Improved Version of Logistic Bayesian LASSO for Detecting Rare Haplotype-Environment Interactions with Application to Lung Cancer

Yuan Zhang, Swati Biswas

Cancer Informatics (2015-02-09) https://doi.org/ggxxfp

DOI: 10.4137/cin.s17290 · PMID: 25733797 · PMCID: PMC4332044

31. Multiple Regression Methods Show Great Potential for Rare Variant Association Tests

ChangJiang Xu, Martin Ladouceur, Zari Dastani, J. Brent Richards, Antonio Ciampi, Celia M. T. Greenwood

PLoS ONE (2012-08-08) https://doi.org/f35726

DOI: <u>10.1371/journal.pone.0041694</u> · PMID: <u>22916111</u> · PMCID: <u>PMC3420665</u>

32. A Sparse-Group Lasso

Noah Simon, Jerome Friedman, Trevor Hastie, Robert Tibshirani Journal of Computational and Graphical Statistics (2013-04) https://doi.org/gcvjw8

DOI: <u>10.1080/10618600.2012.681250</u>

33. Regularized logistic regression with adjusted adaptive elastic net for gene selection in high dimensional cancer classification

Zakariya Yahya Algamal, Muhammad Hisyam Lee

Computers in Biology and Medicine (2015-12) https://doi.org/f73xvj

DOI: 10.1016/j.compbiomed.2015.10.008 · PMID: 26520484

34. Sparse logistic regression with a L1/2 penalty for gene selection in cancer classification

Yong Liang, Cheng Liu, Xin-Ze Luan, Kwong-Sak Leung, Tak-Ming Chan, Zong-Ben Xu, Hai Zhang *BMC Bioinformatics* (2013-06-19) https://doi.org/gb8v2x

DOI: <u>10.1186/1471-2105-14-198</u> · PMID: <u>23777239</u> · PMCID: <u>PMC3718705</u>

35. Clustering cancer gene expression data: a comparative study

Marcilio CP de Souto, Ivan G Costa, Daniel SA de Araujo, Teresa B Ludermir, Alexander Schliep *BMC Bioinformatics* (2008-11-27) https://doi.org/dggbn6

DOI: <u>10.1186/1471-2105-9-497</u> · PMID: <u>19038021</u> · PMCID: <u>PMC2632677</u>

36. Removing Batch Effects From Histopathological Images for Enhanced Cancer Diagnosis

Sonal Kothari, John H. Phan, Todd H. Stokes, Adeboye O. Osunkoya, Andrew N. Young, May D. Wang

IEEE Journal of Biomedical and Health Informatics (2014-05) https://doi.org/gdm9jd

DOI: 10.1109/jbhi.2013.2276766 · PMID: 24808220 · PMCID: PMC5003052

37. Multidimensional Scaling

Michael A. A. Cox, Trevor F. Cox

Springer Berlin Heidelberg (2008) https://doi.org/dg9m4f

DOI: 10.1007/978-3-540-33037-0 14

38. Principal component analysis: a review and recent developments

Ian T. Jolliffe, Jorge Cadima

Philosophical Transactions of the Royal Society A: Mathematical, Physical and Engineering Sciences (2016-04-13) https://doi.org/gcsfk7

DOI: 10.1098/rsta.2015.0202 · PMID: 26953178 · PMCID: PMC4792409

40. **UMAP: Uniform Manifold Approximation and Projection for Dimension Reduction** Leland McInnes, John Healy, James Melville arXiv (2018-12-07) https://arxiv.org/abs/1802.03426

41. How to Use t-SNE Effectively

Martin Wattenberg, Fernanda Viégas, lan Johnson *Distill* (2016-10-13) https://doi.org/gffk7g

DOI: 10.23915/distill.00002

42. Compressing gene expression data using multiple latent space dimensionalities learns complementary biological representations

Gregory P. Way, Michael Zietz, Vincent Rubinetti, Daniel S. Himmelstein, Casey S. Greene *Genome Biology* (2020-05-11) https://doi.org/gg2mjh

DOI: <u>10.1186/s13059-020-02021-3</u> · PMID: <u>32393369</u> · PMCID: <u>PMC7212571</u>

43. Automatic detection of rare pathologies in fundus photographs using few-shot learning

Gwenolé Quellec, Mathieu Lamard, Pierre-Henri Conze, Pascale Massin, Béatrice Cochener *Medical Image Analysis* (2020-04) https://doi.org/ggsrc7

DOI: <u>10.1016/j.media.2020.101660</u> · PMID: <u>32028213</u>

44. Sensitive detection of rare disease-associated cell subsets via representation learning

Eirini Arvaniti, Manfred Claassen

Nature Communications (2017-04-06) https://doi.org/gf9t7w

DOI: <u>10.1038/ncomms14825</u> · PMID: <u>28382969</u> · PMCID: <u>PMC5384229</u>

45. Adjusting batch effects in microarray expression data using empirical Bayes methods

W. Evan Johnson, Cheng Li, Ariel Rabinovic

Biostatistics (2007-01) https://doi.org/dsf386

DOI: <u>10.1093/biostatistics/kxj037</u> · PMID: <u>16632515</u>

46. svaseq: removing batch effects and other unwanted noise from sequencing data

Jeffrey T. Leek

Nucleic Acids Research (2014-12-01) https://doi.org/f8k8kf

DOI: <u>10.1093/nar/gku864</u> · PMID: <u>25294822</u> · PMCID: <u>PMC4245966</u>

47. A scaling normalization method for differential expression analysis of RNA-seq data

Mark D Robinson, Alicia Oshlack

Genome Biology (2010) https://doi.org/cq6f8b

DOI: <u>10.1186/gb-2010-11-3-r25</u> · PMID: <u>20196867</u> · PMCID: <u>PMC2864565</u>