



Bioinformatics
doi.10.1093/bioinformatics/xxxxxx
Advance Access Publication Date:
Applications Note



Genetics and Population Analysis

skater: An R package for SNP-based Kinship Analysis, Testing, and Evaluation

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Associate Editor: XXX

Received on XXX; revised on XXX; accepted on XXX

Abstract

Motivation: SNP-based kinship analysis with genome-wide relationship estimation and IBD segment analysis methods produces results that often require further downstream processing and manipulation. A dedicated software package that consistently and intuitively implements this analysis functionality is needed.

Results: Here we present the skater R package for **S**NP-based **k**inship **a**nalysis, **t**esting, and **e**valuation with **R**. The skater package contains a suite of well-documented tools for importing, parsing, and analyzing pedigree data, performing relationship degree inference, benchmarking relationship degree classification, and summarizing IBD segment data.

Availability: The skater package is implemented as an R package and is released under the MIT license at https://github.com/signaturescience/skater. Documentation is available at https://signaturescience.github.io/skater.

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Supplementary information: Supplementary data are available at Bioinformatics Online.

1 Introduction

Inferring familial relationships between individuals using genetic data is a common practice in population genetics, medical genetics, and forensics. There are multiple approaches to estimating relatedness between samples, including genome-wide measures, such as those implemented in Plink (Purcell et al., 2007) or KING (Manichaikul et al., 2010), and methods that rely on identity by descent (IBD) segment detection, such as GERMLINE (Gusev et al., 2009), hap-IBD (Zhou et al., 2020), and IBIS (Seidman et al., 2020). Recent efforts focusing on benchmarking these methods (Ramstetter et al., 2017) have been aided by tools for simulating pedigrees and genome-wide SNP data (Caballero et al., 2019). Analyzing results from genome-wide SNP-based kinship analysis or comparing analyses to simulated data for benchmarking have to this point required writing one-off analysis functions or utility scripts that are seldom distributed with robust documentation, test suites, or narrative examples of usage. There is a need in the field for a well-documented software package

with a consistent design and API that contains functions to assist with downstream manipulation, benchmarking, and analysis of SNP-based kinship assessment methods. Here we present the skater package for SNP-based kinship analysis, testing, and evaluation with **R**.

2 The skater package

The skater package provides an intuitive collection of analysis and utility functions for SNP-based kinship analysis. Functions in the package include tools for importing, parsing, and analyzing pedigree data, performing relationship degree inference, benchmarking relationship degree classification, and summarizing IBD segment data. The package is designed to adhere to "tidy" data analysis principles, and builds upon the tools released under the tidyverse R ecosystem.



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2.1 Pedigree parsing, manipulation, and analysis

The skater package has several functions for importing, parsing, and analyzing pedigree data. Pedigrees define familial relationships in a hierarchical structure. Many genomics tools for working with pedigrees start with a .fam file, which is a tabular format with one row per individual and columns for unique IDs of the mother, father, and the family unit. The skater package contains the function <code>read_fam()</code> to read in a PLINK-formatted .fam file and another function <code>fam2ped()</code> to convert the content into a pedigree object as a nested tibble with one row per family. All pedigree processing from skater internally leverages a data structure from the kinship2 package (Sinnwell *et al.*, 2014). Further functions such as <code>plot_pedigree()</code> produce a multi-page PDF drawing a diagram of the pedigree for each family, while <code>ped2kinpair()</code> produces a pairwise list of relationships between all individuals in the data with the expected kinship coefficients for each pair (see Supplementary Material).

2.2 Relationship degree inference and benchmarking

The skater package includes functions to translate kinship coefficients to relationship degrees. The kinship coefficients could come from ped2kinpair() or other kinship estimation software.

The dibble() function creates a degree inference tibble, with degrees up to the specified maximum degree resolution, expected kinship coefficient, and lower and upper inference ranges as defined in Manichaikul et al. (2010). The kin2degree() function infers the relationship degree given a kinship coefficient and a maximum degree resolution (e.g., 7th-degree relatives) up to which anything more distant is classified as unrelated.

Once estimated kinship is converted to degree, it may be of interest to compare the inferred degree to known degrees of relatedness. When aggregated over many relationships and inferences, this can help benchmark performance of a particular kinship analysis method. The skater package adapts a confusion_matrix() function from Clark (2021) to provide standard contingency table metrics (e.g. sensitivity, specificity, PPV, precision, recall, F1, etc.) with a new reciprocal RMSE (R-RMSE) metric. The R-RMSE metric is defined more thoroughly in the skater package vignette (see Supplementary Material) and may be a preferable measure of classification accuracy when benchmarking relationship degree estimation. In many kinship benchmarking analyses, classification error is treated in a categorical manner (exact match plus or minus one degree), neglecting the true amount of sharing as a real number. Taking the reciprocal of the target and predicted degree in a typical RMSE calculation results in larger penalties for more egregious misclassifications (e.g., classifying a first-degree relative pair as second-degree) than misclassifications at more distant relationships (e.g., classifying a fourth-degree relative pair as fifth-degree).

2.3 IBD segment analysis

Tools such as hap-IBD (Zhou et~al., 2020), and IBIS (Seidman et~al., 2020) detect shared IBD segments between individuals. The skater package includes functionality to take those IBD segments, compute shared genomic centimorgan (cM) length, and converts that shared cM to a kinship coefficient. In addition to inferred segments, these functions can estimate "truth" kinship from simulated IBD segments (Caballero et~al., 2019). The read_ibd() function reads pairwise IBD segments from IBD inference tools and from simulated IBD segments. The read_map() function reads in genetic map in a standard format which is required to translate the total centimorgans shared IBD to a kinship coefficient using the ibd2kin() function.

3 Conclusion

The skater R package provides a robust software package for data import, manipulation, and analysis tasks typically encountered when working with SNP-based kinship analysis tools. All package functions are internally documented with examples, and the package contains a vignette demonstrating usage, inputs, outputs, and interpretation of all key functions (see Supplementary Material). The package contains internal tests that are automatically run with continuous integration via GitHub Actions whenever the package code is updated. The skater package is permissively licensed (MIT) and is easily extensible to accommodate outputs from new genome-wide relatedness and IBD segment methods as they become available.

Funding

This work was supported in part by award 2019-DU-BX-0046 (Dense DNA Data for Enhanced Missing Persons Identification) to B.B., awarded by the National Institute of Justice, Office of Justice Programs, U.S. Department of Justice and by internal funds from the Center for Human Identification. The opinions, findings, and conclusions or recommendations expressed are those of the authors and do not necessarily reflect those of the U.S. Department of Justice.

References

Caballero, M., Seidman, D. N., Qiao, Y., Sannerud, J., Dyer, T. D., Lehman, D. M., Curran, J. E., Duggirala, R., Blangero, J., Carmi, S., and Williams, A. L. (2019). Crossover interference and sex-specific genetic maps shape identical by descent sharing in close relatives. *PLOS Genetics*, **15**(12), e1007979.

Clark, M. (2021). https://github.com/m-clark/confusionmatrix.

Gusev, A., Lowe, J. K., Stoffel, M., Daly, M. J., Altshuler, D., Breslow, J. L., Friedman, J. M., and Pe'er, I. (2009). Whole population, genome-wide mapping of hidden relatedness. *Genome Research*, 19(2), 318–326.

Manichaikul, A., Mychaleckyj, J. C., Rich, S. S., Daly, K., Sale, M., and Chen, W.-M. (2010). Robust relationship inference in genomewide association studies. *Bioinformatics (Oxford, England)*, **26**(22), 2867–2873.

Purcell, S., Neale, B., Todd-Brown, K., Thomas, L., Ferreira, M. A. R., Bender, D., Maller, J., Sklar, P., de Bakker, P. I. W., Daly, M. J., and Sham, P. C. (2007). PLINK: A Tool Set for Whole-Genome Association and Population-Based Linkage Analyses. *The American Journal of Human Genetics*, 81(3), 559–575.

Ramstetter, M. D., Dyer, T. D., Lehman, D. M., Curran, J. E., Duggirala, R., Blangero, J., Mezey, J. G., and Williams, A. L. (2017). Benchmarking Relatedness Inference Methods with Genome-Wide Data from Thousands of Relatives. *Genetics*, **207**(1), 75–82.

Seidman, D. N., Shenoy, S. A., Kim, M., Babu, R., Woods, I. G., Dyer, T. D., Lehman, D. M., Curran, J. E., Duggirala, R., Blangero, J., and Williams, A. L. (2020). Rapid, Phase-free Detection of Long Identity-by-Descent Segments Enables Effective Relationship Classification. American Journal of Human Genetics, 106(4), 453–466.

Sinnwell, J. P., Therneau, T. M., and Schaid, D. J. (2014). The kinship2 R Package for Pedigree Data. *Human heredity*, **78**(2), 91–93.

Zhou, Y., Browning, S. R., and Browning, B. L. (2020). A Fast and Simple Method for Detecting Identity-by-Descent Segments in Large-Scale Data. *The American Journal of Human Genetics*, **106**(4), 426–437.



