A comparative study of evaluating missing value imputation methods in label-free proteomics

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The presence of missing values (MVs) in label-free quantitative proteomics greatly reduces the completeness of data. Imputation has been widely utilized to handle MVs, and selection of the proper method is critical for the accuracy and reliability of imputation. Here we present a comparative study that evaluates the performance of seven popular imputation methods with a large-scale benchmark dataset and an immune cell dataset. Simulated MVs were incorporated into the complete part of each dataset with different combinations of MV rates and missing not at random (MNAR) rates. Normalized root mean square error (NRMSE) was applied to evaluate the accuracy of protein abundances and intergroup protein ratios after imputation. Detection of true positives (TPs) and false altered-protein discovery rate (FADR) between groups were also compared using the benchmark dataset. Furthermore, the accuracy of handling real MVs was assessed by comparing enriched pathways and signature genes of cell activation after imputing the immune cell dataset. We observed that the accuracy of imputation is primarily affected by the MNAR rate rather than the MV rate, and downstream analysis can be largely impacted by the selection of imputation methods. A random forest-based imputation method consistently outperformed other popular methods by achieving the lowest NRMSE, high amount of TPs with the average FADR < 5%, and the best detection of relevant pathways and signature genes, highlighting it as the most suitable method for label-free proteomics.

Large-scale protein quantitation of biospecimens is critical for biological, biomedical and pharmaceutical researches. Liquid chromatography-mass spectrometry (LC–MS) based proteomics technology is a popular approach in profiling global protein expressions. Thousands of proteins can be identified and quantified in a single MS injection. Two common quantification methods have been developed in bottom-up proteomics: label-based and label-free1. Label-free proteomics is more commonly employed in large-scale biological studies because of its cost-effective sample processing and capacity of quantifying a very large number of samples in one batch. Key applications of label-free proteomics include the discovery of biomarkers and new drug targets, but a major issue is that the power of statistical inference and downstream functional analysis is greatly impacted by the presence of missing values (MVs) in the protein abundance data. Multiple factors contribute to MVs in LC–MS based proteomics, including biological factors, such as proteins do not exist and protein abundances are below the instrument detection limit, as well as analytical factors, such as sample loss in preparation, mis-cleavage of peptides during digestion, poor ionization efficiency and bad peptide-spectrum matches2. Missing values in proteomic data can be generally characterized into missing at random (MAR) and missing not at random (MNAR)2,3. MAR missing values mostly result from technical limitations and stochastic fluctuations in an abundance-independent manner, and oppositely, MNAR missing values are more abundance-dependent that can be explained by the measurability of the corresponding peptides. Missing values in proteomic data are a mixture of MAR and MNAR. Although the real proportion is difficult to determine, it is believed that MNAR plays a dominant role in producing missing values4.

To minimize the constraints of MVs, imputation methods are used to replace missing values with estimated values. Skewed distribution of protein abundances is often observed in proteomic data because of the existence of MNAR missing values2. Therefore, left-censored methods are applied as they replace MVs with small values closed to or smaller than the globally minimum quantitative value to mimic MNAR2–6. Imputation methods that are commonly utilized in other biological expression data (microarray, etc.)7,8, such as local similarity methods

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and global structure methods, have also been introduced to proteomics because they can handle mixed types of MVs2,3,5.

To date, what is the best method to handle MVs in proteomic data remains controversial. One big challenge is the lack of a high-quality, large-scale dataset to benchmark protein abundance that enables a more systematic and comprehensive evaluation. Proteomic datasets with spike-in controls were used in some evaluation studies but they either were relatively small-scale or contained only a few spike-in proteins5,6, which were not comparable to real-world proteomic data. Evaluation works using these datasets could be inaccurate and unreliable. Models were developed to generate artificial peptide and protein abundance for evaluating imputation methods2,3, but real proteomic data is still preferable because whether the simulated datasets bring systematic bias is not clear yet. Furthermore, the lack of a representative collection of imputation methods in one study also prevents the selection of the best method. For example, the top-ranked imputation methods from one study7 were not included in the other1, resulting in difficult cross comparisons. In addition, simulated missing values have been commonly used for evaluating the performance of imputation3,9,10,18, but testing with real missing values is still highly desirable. However, without knowing the real values, differences between imputed and real values cannot be calculated, thus a better strategy of comparison is needed.

A novel MS1-based label-free proteomics workflow, named IonStar, was developed recently to generate high-quality quantitative proteomic data for large-scale studies. It remarkably outperforms other popular proteomics software for its high quantitation accuracy and extremely low missing value rate11. In this study, we utilized a benchmark spike-in dataset generated by the IonStar workflow to explore the performance of imputation methods. Quantitative protein abundances were measured across 32 samples that were grouped with different portions of E. coli and yeast proteome spiked into the human proteome. Intragroup variations were added on purpose to mimic biological replicates of large-cohort studies. MAR and MNAR missing values were then incorporated with designated proportions to simulate different mechanisms of missingness. Protein ratios after imputation were compared with real spike-in ratios. Detection of differentially expressed proteins was also assessed in terms of true positives (TPs) and false altered-protein discovery rate (FADR). To allow an insightful yet not exhaustive comparison, we intentionally selected seven imputation methods: lowest of detection (LOD), random drawing of a left-censored normal distribution (ND), k-nearest neighbors (kNN), local least squares (LLS), random forest (RF), singular value decomposition (SVD) and Bayesian principal component analysis (BPCA). These methods cover left-censored, local similarity, and global-structure methods, and have demonstrated robust performance in previous studies3,5,6,12,13. Moreover, we utilized a label-free proteomic dataset of immune cells sorted by flow cytometry in this study. This dataset contains both steady and activated states of each cell type, and the alterations of signaling pathways regarding the activations have already been well understood14. Therefore, whether these pathways are detected after imputation can be used to evaluate the performance of imputation methods.

Results

Two large-scale label-free proteomic datasets were selected to evaluate the performance of imputation methods. In the benchmark dataset, small and variable amounts of E. coli and yeast proteins were spiked into the human proteome with designated ratios. The proportion of human proteins in all samples remained constantly 70% as background. In group A to D, the proportion of E. coli proteins was 5%, 7.5%, 10%, and 12.5%, and the proportion of yeast proteins was 25%, 22.5%, 20%, and 17.5%, respectively. Therefore, between Group B-D and Group A, ratios of E. coli proteins were 1.5, 2, and 2.5, and ratios of yeast proteins were 0.9, 0.8, and 0.7 (Table S1). In addition, proportions of E. coli and yeast proteins were variable in eight individual samples of each group to mimic variations in biological replicates, while the mean values of each group were as designated and differences between Group B-D and Group A were statistically significant (p-value < 0.05) (Table S1). The benchmark dataset contains 1,184 E. coli proteins, 1,081 yeast proteins, and 4,424 human proteins without MVs. In another study, Rieckmann and colleagues profiled the proteomes of steady and activated states of 28 different types of blood cells sorted by flow cytometry from human donors14. The data of steady-state and activated B cells, T4 cells, T8 cells, and monocytes were selected for evaluating the performance of imputation methods. The immune cell dataset contains 10,081 proteins with 27% values are missing, within which 3,963 protein contain zero MVs.

To test the performance of imputation methods, simulated MVs were incorporated into the complete datasets at three different rates (10%, 20%, 30%). Three rates of MNAR (20%, 50%, 80%) were further combined with each MV rate to mimic low, middle, and high levels of MNAR. Ten dataset repeats were generated with the same missingness condition to test the reproducibility of imputation methods, resulting in ninety test datasets in total from each original dataset (Fig. 1). To ensure unbiased comparison, the key parameters of imputation methods were optimized. The parameters were tested with the benchmark dataset in three conditions (20%MV-20%MNAR, 20%MV-50%MNAR, 20%MV-80%MNAR). Normalized Root Mean Square Error (NRMSE) between imputed and real protein abundances was utilized to determine the accuracy of imputation. For the local similarity methods kNN and LLS, it is critical to select a k-value that determines the number of most similar proteins for estimation15. We chose k (number of nearest neighbors) = 6 for kNN and k (number of similar proteins used for regression) = 150 for LLS (Fig. S1A,B). The RF-based imputation method does not require tuning specific parameters16. The different number of trees to grow in each forest (ntree) did not show a significant change in imputation accuracy, thus default setting (ntree = 100) was applied (Fig. S1C). For global-structure methods, the accuracy of imputation largely depends on the number of principal components used for regression17,18. The number of principal components (nPCs) was set to 2 for both SVD and BPCA based on the overall best performance (Fig. S1D,E).

NRMSE has been widely used to evaluate the difference between imputed values and real values9,10,18. Therefore, we used NRMSE to compare the imputed and original protein abundances with both datasets (Fig. 1).
Additional comparisons were performed with the benchmark dataset. First, the accuracy of the intergroup protein ratio was compared between the designated ratios and observed ratios after imputation by calculating NRMSE as well. Second, detection of DE proteins was compared by counting how many E. coli and yeast proteins (true positives, TPs) were significantly altered between Group B-D and Group A. Third, false altered-protein discovery rate (FADR), which was defined by the proportion of human proteins (false positives, FPs) in the total positives, was used to assess the false positive rate of DE protein detection after imputation11. The whole immune cell dataset consists of four cell types in steady and activated states with approximately 27% missing values. After imputation, Gene Ontology (GO) biological processes enrichment was analyzed with DE proteins from activation of each cell type to test if relevant proteins, activation processes and signaling pathways were detected (Fig. 1).

**Imputation accuracy of protein abundances and intergroup protein ratios with the benchmark datasets.** Accuracy of imputation was first evaluated by comparing NRMSE of imputed and original protein abundance values using the benchmark dataset. Overall, the left-censored methods, LOD and ND, had the highest average NRMSE of all test datasets, and RF and LLS estimated protein abundances with the highest accuracy as they displayed the lowest average NRMSE. Between the global-structure methods, BPCA performed better than SVD (Fig. 2A). Furthermore, NRMSE of protein abundances was compared in terms of different rates of MV and MNAR. When the MV rate increased, NRMSE of LOD, ND, and kNN increased as well. NRMSE of LLS, RF, SVD, and BPCA remained constant with low or middle level of MNAR regardless of the level of MV and slightly dropped when the MV rate increased with 80% MNAR in the datasets (Fig. 2B). Left-censored methods performed better with higher MNAR rates as expected. Interestingly, a remarkable boost of NRMSE was observed for non-left-censored methods when the MNAR rate increased from 50 to 80% (Fig. 2B). Particularly, NRMSE of SVD imputation displayed a larger increase between 50% MNAR and 80% MNAR than other methods, suggesting that SVD was the most sensitive method to high MNAR rate.

Besides of accuracy of protein abundance values, we also investigated whether relative protein ratios between groups could be accurately estimated after imputation. The protein ratios between Group B-D and Group A after imputation were compared with designated spike-in protein ratios and the differences were evaluated by NRMSE as well. RF and BPCA had the lowest average NRMSE of all test datasets and their NRMSE of protein ratios were consistent regardless of different rates of MV and MNAR (Fig. 2C,D), indicating that these two methods performed the best in estimating intragroup protein ratios. Of note, although LLS had similarly low level of NRMSE as RF in imputing protein abundances (Fig. 2A), it showed significantly higher NRMSE of protein ratios with extremely large variations among datasets (Fig. 2C), and the high NRMSE of LLS mostly resulted from 30%MV-80%MNAR datasets (Fig. 2D). Similar to LLS, SVD imputation performed better than ND and kNN in estimating protein abundances but had much higher NRMSE when estimating protein ratios (Fig. 2A,C) especially with datasets containing 80% MNAR missing values (Fig. 2D). Between the two left-censored methods, ND significantly outperformed LOD in estimating protein ratios with around 30-fold lower NRMSE. Together, these results suggest that RF imputation is the most accurate method in terms of protein abundance values and relative protein ratios, and the robustness of imputation is primarily affected by the rate of MNAR.

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**Figure 1.** Analysis workflow. Missing values were incorporated into the benchmark dataset and complete part of the immune cell dataset in nine MV-MNAR combinations. Ten repeats were created for each MV-MNAR situation based on different random seeds. Datasets with missing values were imputed separately with seven methods: lowest of detection (LOD), random drawing from a left-censored normal distribution (ND), k-nearest neighbors (kNN), local least squares (LLS), random forest (RF), singular value decomposition (SVD) and Bayesian principal component analysis (BPCA). Accuracy of proteins abundances and intergroup protein ratios were compared based on normalized root mean square error (NRMSE) between real and after imputation. DE protein detection between Group B-D and Group A of the benchmark dataset was compared based on the amounts of true positives (TPs), false altered-protein discovery rate (FADR), and receiver operating characteristic (ROC) curves. The whole immune cell dataset with real missing values was imputed using seven methods, and enriched pathways of immune cell activation were compared.
Detection of differentially expressed proteins with the benchmark datasets. In the benchmark dataset, the spiked-in *E. coli* and yeast proteins were designed to be significantly different between Group B-D and Group A with intragroup variations. Therefore, the accuracy of DE protein detection can be assessed by the number of non-human proteins (TPs) that are significantly altered after imputation and the proportion of human proteins (FPs) in all significantly altered proteins (FADR). In general, imputation using LLS, RF, SVD, and BPCA detected more TPs than LOD, ND, and kNN (Fig. 3A–C). Between Group D and Group A, over 90% of non-human proteins displayed significant alteration after imputation by LLS, RF, SVD, and BPCA (Fig. 3C). Increasing of the MNAR rate or decreasing of the MV rate led to more TPs detected by LOD, ND, and kNN but less TPs detected by SVD and BPCA, while LLS and RF detected consistent amounts of TPs across different MV-MNAR conditions (Fig. 3A–C). Although SVD and BPCA appeared to detect more TPs than RF and LLS, especially in the comparison between Group B and Group A (Fig. 3A), it was notably that SVD and BPCA also introduced more FPs (Fig. 3D). The average FADR of RF and LLS (4.9% and 5.7%, respectively) were nearly half of those of SVD and BPCA (10.8% and 9.0%, respectively). Particularly for Group B/Group A comparison in datasets with 30%MV-80% MNAR, the average FADR of SVD was 23.8%, which was 6.3-fold higher than the average FADR of RF (3.8%) (Fig. 3D). Interestingly, although the amounts of TPs detected by LLS and RF imputation were somewhat constant with increasing MNAR, the FADR increased markedly. For example, in the situation of 20% MV in Group B/Group A comparison, RF and LLS detected 1% and 0% more TPs when the MNAR rate raised from 50 to 80%, respectively, but FADR of RF increased twofold and FADR of LLS increased 2.5-fold (Fig. 3A,D). The boost of FADR with a high MNAR rate was observed for most non-left-censored methods, including RF, LLS, SVD, and BPCA (Fig. 3D–F). These results suggest that the accuracy of DE protein detection of imputation methods is mainly determined by the rate of MNAR.

To further compare the accuracy of DE protein detection, we plotted the average true positive rate (TPR) and false-positive rate (FPR) on receiver operating characteristic (ROC) curves for different MV-MNAR conditions.

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**Figure 2.** NRMSE of protein abundance and intergroup protein ratios with the benchmark dataset. (A) Overall NRMSE of protein abundances of seven imputation methods. (B) NRMSE of protein abundances in nine MV-MNAR conditions. (C) Overall NRMSE of intergroup protein ratios of seven imputation methods. (D) NRMSE of intergroup protein ratios in nine MV-MNAR conditions. In graph (A) and (C), bars represent average values of all ninety repeats, and error bars represent standard deviations. In graph (B) and (D), bars represent average values of ten repeats in each MV-MNAR condition, and error bars represent standard deviations. Nine MV-MNAR conditions are (from left to right): 10%MV-20%MNAR, 10%MV-50%MNAR, 10%MV-80%MNAR, 20%MV-20%MNAR, 20%MV-50%MNAR, 20%MV-80%MNAR, 30%MV-20%MNAR, 30%MV-50%MNAR, 30%MV-80%MNAR.
For Group B/Group A comparison, RF, LLS, SVD, and BPCA explicitly demonstrated the best performance with 20% MNAR missing values (Fig. 4A–C). RF outperformed other methods by showing relatively higher TPR and lower FPR in most conditions with 50% and 80% MNAR (Fig. 4D–I). In the most extreme condition (30% MV-80% MNAR), RF achieved approximately 74% TPR with 5% FPR, which was roughly 10% higher than LLS and 20% higher than BPCA (Fig. 4I). Of note, the performance of SVD imputation was comparable with RF when the MNAR rate was low but dropped dramatically while the MNAR rate increased (Fig. 4), suggesting that the accuracy of SVD was largely dependent on the mechanism of missingness. Similar results were observed from the comparisons between Group C-D and Group A as well, that RF imputation performed equally well or better than other methods consistently (Fig. S2A,B). In summary, DE protein detection was the most accurate if RF imputation was employed.

Evaluating imputation methods with the immune cell dataset. Accuracy of imputing protein abundance values was also evaluated using the complete part of the immune cell dataset, which contained 3,963 proteins. Missing values were incorporated with the same MV-MNAR conditions as described before for the benchmark dataset (Fig. 1). RF imputation again achieved the lowest average NRMSE of all test datasets while left-censored methods showed the highest average NRMSE (Fig. 5A). Unlike the benchmark dataset, with which SVD displayed much higher NRMSE than BPCA (Fig. 2A), SVD and BPCA had similar average NRMSE with the immune cell datasets (Fig. 5A). Furthermore, we compared the NRMSE across different MV-MNAR conditions. Accuracy of left-censored methods, LOD and ND, rose with the increase of MNAR rate and fell with the increase of MV rate. The non-left-censored methods shared similar patterns that increasing MV rate with 20% or 50% MNAR did not affect NRMSE much and increasing MV rate with 80% MNAR slightly reduced NRMSE (Fig. 5B). Additionally, for non-left-censored methods, a much greater increase of NRMSE was observed when the MNAR rate increased from 50 to 80% than that from 20 to 50%, which was consistent with observations from the benchmark dataset (Figs. 2B, 5B).

The whole immune cell dataset containing 10,081 proteins and 27% missing values was imputed using different methods to test their performance on real missing values. For each immune cell type, DE proteins between activated and steady-state were determined by BH-adjusted p-value < 0.05. The over-represented biological processes from Gene Ontology (GO) annotation within the DE proteins were then determined by enrichment analysis and BH-adjusted p-value < 0.05 was considered as statistically significant. The enriched activation processes and signaling pathways used for comparison were selected based on the biological relevance to the specific stimulation of each cell type. Most B cell activation-related and monocyte activation-related pathways showed enrichment by all imputation methods but with different levels of significance (Figs. 6A, S3A).
when B cells were activated, imputation using LLS and RF resulted in more significant enrichment of regulation of lymphocyte activation, and more proteins related to this regulation process were characterized as DE proteins as well (Fig. 6A). Moreover, the TOR signaling pathway was essential in regulating B cell activation, but significant enrichment of TOR signaling was only detected by RF and LLS imputation. The outperformance of RF and LLS was more apparent when comparing enriched pathways of activated T4 cells. T cell activation process was more significantly enriched by LLS and RF imputation, and more T cell activation DE proteins were detected as well (Fig. 6B). In addition, enrichment of T cell activation involved in immune response, I-kappaB kinase/NF-kappaB signaling and integrin-mediated signaling pathway were uniquely detected by RF and LLS imputation (Fig. 6B). For T8 cell activation, LLS, RF, and BPCA imputation could detect most of the relevant pathways. Notably, only the T cell activation process was enriched after LOD and ND imputation, and no pathway was significantly enriched with kNN imputation (Fig. S3B). Together, RF and LLS imputation consistently demonstrate the best performance in detecting enriched biological pathways in terms of the relevance to specific immune cell activations.

We further investigated whether specific gene signatures of T4 cell activation can be detected by different imputation methods. From a signature gene list of T4 cell activated by anti-CD3 and anti-CD28, 14 genes were identified in the immune cell proteomic dataset with at least one missing value across all samples and detected as DE by at least one imputation method. Without imputation, eight proteins had sufficient values for statistical analysis and only one protein were determined DE protein in activated T4 cells. 14 and 15 DE proteins were detected by RF and LLS imputation, respectively, which markedly improved the detection of the signatures of T4 activation. Meanwhile, no obvious improvement was observed when imputation was performed using other methods (Fig. S5). These results suggested that RF and LLS outperformed other imputation methods at the individual protein level in terms of accurate detection of known gene signatures.
Missing value imputation is a key preprocessing step in large-scale label-free proteomic studies to improve proteome coverage and statistical power. A comprehensive evaluation of imputation methods is critical for selecting the most suitable one by understanding the advantages and disadvantages of each method. Our results revealed that the RF and LLS imputation methods consistently performed better than other methods, and RF slightly outperformed LLS in terms of protein ratio estimation and DE protein detection. Besides the consistently robust performance across all MV-MNAR situations with both datasets, RF imputation does not require parameter adjustment, together suggest that RF is the most suitable method for label-free proteomic studies without a comprehensive understanding of the mechanism of the missingness. Though RF imputation takes relatively longer time to process (Table S1), probably due to the intensive iterations of the algorithm, it's considerably
shorter than time-consuming steps in proteomics, such as data acquisition and database search, and could be
shortened by high performance computers.

Performance of LLS imputation was highly comparable to RF-based on our results, however, when the MV rate
and MNAR rate were both high, the intergroup protein ratios could be unreliable (Fig. 2D). In fact, the
exceptionally high NRMSE of protein ratios was only observed in one dataset repeat in 30%MV-80%MNAR
condition while NRMSE of protein abundances appeared normal. Though the reason for this phenomenon is
not clear yet, it raises concern that protein ratios obtained after LLS imputation may be misleading.

Recent evaluation studies with metabolomic data showed contradictory results for SVD imputation9,10.
Between the two global-structure imputation methods, our results have clearly shown that BPCA outperformed
SVD, and the accuracy of SVD was greatly influenced by the high rate of MNAR. In addition, global structure
methods are less preferable because of the high false discovery rate of DE protein detection, although BPCA
achieved similar accuracy to RF in terms of NRMSE of protein intensities and protein ratios.

The proportion of MNAR missing values appears to be the primary determinant of imputation accuracy.
Left-censored methods were devoted to handling MNAR as they replaced missing values with relatively small
values. As expected, only left-censored methods showed improvement while the MNAR rate increased. On the
other hand, it was surprising that the NRMSE of LLS, RF, SVD, and BPCA decreased slightly when the MV rate
increased. This phenomenon was described for BPCA imputation with metabolomic data as well12. Contradicto-
riely, Lazar et al. employed RMSE-observations standard deviation ratio (RSR), which was a normalized version
of RMSE as well, for evaluation and demonstrated that performance of SVD imputation decreased with increasing
MV rate in proteomic data1. Although how the MV rate affects imputation accuracy still needs further investiga-
tion, our results suggest that better imputation performance can be achieved by minimizing the MNAR rate.

If missing values in real datasets can be categorized into MAR and MNAR, it would be beneficial to apply
imputation methods based on the mechanism of missingness. Left-censored methods, such as LOD or ND,
can be used to impute MNAR missing values, and RF or LLS can be used to handle MAR missing values. This
hybrid method handles every missing value individually based on its mechanism, which in theory provides the
best estimation of missing values. However, missing values in real-world proteomic datasets are mixed MAR
and MNAR that can hardly be characterized individually. For proteomic studies with multiple sample groups,
particularly with different cell types, stimulations or drug treatments, it is common that certain proteins are only
detectable in specific groups. For instance, the T-cell surface glycoprotein CD4 was completely missing in T8
cells, and the T-cell surface glycoprotein CD8B was completely missing in B cells and monocytes14. Therefore, if
a protein is completely missing in one sample group, these missing values could be more likely MNAR. Previous
studies have applied a similar method that one MV was considered MAR only if it appeared once in a sample
group. Proteins with MAR missing values were filtered out from statistical analysis and MNAR missing values
were replaced by zeros14,22. Nonetheless, whether the hybrid method is more accurate needs more investigation.

Our results of the enriched pathways of immune cell activation clearly showed that downstream analysis
could be greatly impacted by imputation. For example, no pathway was significantly enriched for T8 cell activa-
tion if kNN imputation was applied (Fig. S3B). Interestingly, SVD and BPCA detected the largest amount of
DE proteins in the benchmark dataset, but RF and LLS detected much more DE proteins than SVD and BPCA
in the immune cell dataset (Fig. S4). This could be due to the SVD and BPCA algorithms are more robust with
structured missing values, suggesting that RF and LLS perform better in handling real missing values. Analysis
at the single protein level using signature genes of T4 cell activation further confirmed that RF and LLS detected
DE proteins more accurately (Fig. S5), which could lead to better biological data interpretation.

There are some limitations in this study that need to be taken into consideration in future works. First, other
preprocessing steps, such as log transformation, data filtering and normalization, were not considered in this
study but would contribute to the overall accuracy of the data analysis framework. Thus, a more systematic com-
parison that combines all these steps will be highly desired. Second, we only imputed protein-level quantitative
data for comparison. A previous study has indicated that imputation protein-level and peptide-level may result
in different conclusions3, therefore, peptide-level data, as well as different approaches of aggregating peptide-
level data to protein-level data, need to be assessed together with imputation methods. Third, we intentionally
controlled the rates of MV and MNAR to simulate different mechanisms of missingness, however, the approach
we applied tended to underestimate the MNAR rate. As MAR missing values were randomly selected after incor-
porating MNAR missing values, MNAR patterns could still be created by random chance, especially when the MV
rate was high. Fourth, normalization is preferably performed before imputation3, thus an optimal normalization
approach could improve the performance of RF imputation but needs further investigation.

The performance of imputation is greatly impacted by the rate and type of missing values in label-free
proteomics. Proper selection of imputation methods markedly reduces error rate and improves downstream
analysis. Our comparative study demonstrates that an RF-based imputation method has consistently the most
robust performance and does not require a clear understanding of the mechanism of missingness, which is the
most suitable for proteomic data. Our strategy to evaluate the detection of DE proteins and performance with
real missing values can also be implemented in future studies.

Methods

Datasets. Two label-free proteomic datasets were used in this study. The first is a spike-in benchmark data-
set where E. coli and yeast protein digest was mixed with human protein digest in different ratios. The second
dataset is protein expression data of steady-state and activated immune cells.

The benchmark dataset contains seven groups with differential percentages of E. coli and yeast protein digest
spiked in human protein digest. Homo sapiens colon cancer cell line SW620, Saccharomyces cerevisiae strain
BY4741, and E.coli strain ATCC 25,922 were used to generate this dataset. We selected group A to D for this study.

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Each group has eight replicates in this dataset and the composition of *E. coli* and yeast proteins within a group are varied to mimic biological replicates (Table S2). Samples were analyzed by LC-MS and IonStar workflow as described previously14. Raw files and quantitative results are available on ProteomeXchange Consortium via PRIDE repository (https://www.ebi.ac.uk/pride/) with identifier PXD017915.

The immune cell dataset consists of protein intensities of seven cell lineages isolated from human blood14. Protein abundances were measured for both steady-state and activated states by the specific stimulus for each immune cell type. We selected the steady-state and activated data of B cells, T4 cells, T8 cells and monocytes in this study, which had four biological replicates (different blood donors) in each group, except that activated monocytes had three biological replicates. LFQ protein intensities were used as these values were more accurate23.

Intergroup protein ratios were calculated using Group A as the control for the benchmark dataset and the corresponding steady-state group as the control for each immune cell type. Adjusted p-values were calculated by two-sample equal variance Student’s t-test with Benjamini–Hochberg adjustment. An adjusted p-value lower than 0.05 was considered statistically significant.

**Missing value simulation.** To incorporate missing values into the complete datasets with desired rates of MV and MNAR, we modified and implemented an approach described previously3. In brief, with α represents the rate of total missing values in the dataset D and β represents the ratio of MNAR missing values, if the dataset D contains N total values, N × α × β values will be replaced by missing values to simulate MNAR as follows: (1). A threshold matrix T with the same dimensions as D is generated from a normal distribution with μ = αth quantile of all values in D and σ = 0.3. (2). A probability matrix P with the same dimensions as D is generated from a Bernoulli distribution using β as the probability of success. (3). For each value D_{i,j} (i and j represents the ith row and jth column) in the complete dataset, if D_{i,j} < T_{i,j} and P_{i,j} = 1, D_{i,j} is selected as MNAR and replaced by a missing value. After MNAR incorporation, N × α × (1 − β) values are randomly chosen from the remaining values of the dataset and replaced by missing values to simulate MAR. Rows with only missing values were removed from the dataset.

**Missing value imputation.** Instead of an exhaustive comparison, we picked two commonly used left-censored approaches: lowest of detection (LOD) and random drawing from a left-censored normal distribution (ND); three local similarity methods: k-nearest neighbors (kNN), local least squares (LLS), and random forest (RF); and two global-structure methods: Singular value decomposition (SVD) and Bayesian principal component analysis (BPCA). LOD imputation was performed by simply replacing missing values with the lowest value in the dataset. For ND imputation, missing values were replaced by random drawn values from a normal distribution on the left tail of the abundance distribution of the complete dataset. The mean and standard deviation of this left-censored normal distribution was determined as: μ = m̄ − 2σm̄ and σ = 0.3σm̄ (μ = mean, σ = standard deviation, i = values for imputation, m = measured values). The parameters were identical to previously mentioned14. The kNN imputation was implemented by using the kNN function of the “VIM” R package24. LLS imputation was implemented by using LLSImpute function of the “pcamethods” package35. RF imputation was performed with missForest function of the “missForest” R package16. SVD and BPCA imputation were performed using pca function from the “pcamethods” R package as well.

**Performance evaluation.** Lower NRMSE indicates higher imputation accuracy. The nrmse function from “missForest” R package16 was applied to calculate NRMSE of protein intensities and relative protein ratios between imputed and real datasets. In the benchmark dataset, yeast and *E. coli* proteins with adjusted p-values less than 0.05 were considered true positives (TP) and human proteins with adjusted p-values less than 0.05 were considered false positives (FP). Therefore, the false altered-protein discovery rate (FADR) was calculated as FADR = FP/(TP + FP). In the immune cell dataset, differentially expressed proteins (adjust p-value < 0.05) in the activated cells were used for the over-representation test of Gene Ontology (GO) biological processes. The analysis was conducted using “clusterProfiler” R package23.

**Data availability**

The benchmark proteomic data is available on ProteomeXchange Consortium via PRIDE repository (https://www.ebi.ac.uk/pride/) with identifier PXD017915. The immune cell dataset is available from Supplementary data of Rieckmann et al.14. All codes were written with R (Version 3.6.1), and R session information is available in Additional File 1. R codes are available at GitHub (https://github.com/liangjin0912/proteomics_imputation), and all packages used in this study are available at CRAN (https://cran.r-project.org/) or Bioconductor (https://www.bioconductor.org/).

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Author contributions

L.J. wrote the code for missing value simulation and imputation, analyzed the results, and drafted the manuscript. L.J., Y.B., and Y.T. designed the study and interpreted the results. X.W. and S.S. prepared and analyzed samples for the benchmark study. Y.B., C.H., J.Q., and Y.T. critically read and edited the manuscript. All authors read and approved the final manuscript.

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Competing interests

The authors declare no competing interests.

Additional information

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