Assessing sub-cellular resolution in spatial proteomics experiments

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Abstract

The sub-cellular localisation of a protein is paramount in defining its function, and a protein's mis-localisation is known to lead to adverse effect. As a result, numerous experimental techniques and datasets have been published, with the aim of deciphering the localisation of proteins at various scales and resolutions, including high profile mass spectrometry-based efforts. Here, we present a tool, termed QSep, and a meta-analysis assessing and comparing the sub-cellular resolution of 29 such mass spectrometry-based spatial proteomics experiments. Our goal is to provide a simple quantitative report of how well spatial proteomics resolve the sub-cellular niches they describe to inform and guide developers and users of such methods.

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1 Introduction

In biology, the localisation of a protein to its intended sub-cellular niche is a necessary condition for it to assume its biological function. Indeed, the localisation of a protein will determine its specific biochemical environment and its unique set of interaction partners. As a result, the same protein can assume different functions in different biological contexts. Protein mislocalisation can lead to adverse effects and have been implicated in multiple diseases [24, 5, 25].

Spatial proteomics is the systematic and high-throughput study of protein sub-cellular localisation. A wide range of techniques (reviewed in [10, 27]) and computational methods [11] have been documented, that confidently infer the localisation of thousands of proteins. Most techniques rely on some form of sub-cellular separation, many employing differential centrifugation or separation along density gradients, and the subsequent quantitative assessment of relative protein occupancy profiles in these sub-cellular fractions. Reciprocally, a broad array of computational methods have been applied, ranging from unsupervised learning e.g. clustering [29] and dimensionality reduction, and supervised learning such as classification (reviewed in [11]), semi-supervised learning and novelty detection [2] and, more recently, transfer learning [3] and Bayesian modelling [6].

Despite these advances, there is surprisingly little agreement in the community as to what constitutes a reliable spatial proteomics experiment, i.e a dataset that generates confident protein localisation results. Is is however implicit that reliability and trust in the results is dependent on adequate subcellular resolution, i.e. *enough* separation between the different sub-cellular niches being studied to be able to confidently discern protein profiles originating from different sub-cellular niches. And yet, every spatial proteomics publication will somehow arbitrarily claim to have obtained satisfactory resolution.

The importance of adequate sub-cellular resolution reaches beyond the generation of reliable static spatial maps. It is a necessary property of the data to consider tackling more subtle sub-cellular patterns such as multi-and trans-localisation, i.e. the localisation of proteins in multiple sub-cellular niches and the relocation of proteins upon perturbation [11].

In this work, we first describe how to understand and interpret widely used dimensionality reduction methods and visualise spatial proteomics data to critically assess their resolution. We then propose a simple, yet effective method, to quantitatively measure sub-cellular resolution and compare it across different experiments. Our recommendations should be useful to spatial proteomics practitioners, to assess the sub-cellular resolution of their experiments and compare it to similar studies while setting up and optimising their experiments, as well biologists interested in critically assessing spatial proteomics studies and their claims.

All the data and software used in this work is available in the pRoloc and pRolocdata packages [12]. The code to reproduce all results and figures presented here are available in the source of the document, available in the manuscript public repository [9].

2 Spatial proteomics datasets

For this meta-analysis, we make use of 29 spatial proteomics datasets, summarised in table 1. These data represent a diverse range of species, sample types, instruments and quantification methodologies.

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	Data	Proteins	Fractions	Clusters	PC var (%)	Title
	hyperLOPIT2015	5032	20	14	72.26	Protein and PMS-level hyperLOPIT datasets Mouse E14TG2a embryonic stem cells from Chriforou et al. (2016). [4]
	hyperLOPIT2015ms2	7114	10	14	74.72	Protein and PMS-level hyperLOPIT datasets Mouse E14TG2a embryonic stem cells from Chr forou et al. (2016). [4]
	HEK293T2011	1371	8	12	65.04	` ' ' ' ' ' ' ' ' ' ' ' ' ' ' ' ' ' ' '
	hirst2018	2046	15	12	82.50	Data from Hirst et al. 2018 [15]
	hyperLOPITU2OS2017	5020	40	12	63.29	2017 and 2018 hyperLOPIT on U2OS cells [28] fractions)
_	hyperLOPITU2OS2017b	5020	37	12	67.68	2017 and 2018 hyperLOPIT on U2OS cells (cleaned)
	itzhak2016stcSILAC	5265	30	12	70.61	Data from Itzhak et al. (2016) [17]
	itzhak2017	9201	9201 30 12 31.39 Dat	Data from Itzhak et al. 2017 [18]		
	rodriguez2012r1	2215	11	12	38.95	Spatial proteomics of human inducible goble LS174T cells from Rodriguez-Pineiro et al. (2 [23]
	$\tan 2009 r1$	888	4	11	88.49	LOPIT data from Tan et al. (2009) [26]
	E14TG2aS1	1109	8	10	65.98	LOPIT experiment on Mouse E14TG2a Embry Stem Cells from Breckels et al. (2016) [3]
	trotter 2010	347	16	10	81.14	LOPIT data sets used in Trotter et al. (2010)
	beltran2016HCMV120	2045	6	9	79.36	Data from Beltran et al. 2016 [19] HCMV infect 120 hpi (hours post-infection)
	beltran2016HCMV24	2196	6	9	77.18	Data from Beltran et al. 2016 [19] HCMV infect 24 hpi
	beltran 2016 HCMV 48	2206	6	9	74.95	Data from Beltran et al. 2016 [19] HCMV infect 48 hpi

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beltran 2016 HCMV 72	2062	6	9	75.32 Data from Beltran et al. 2016 [19] HCMV infecti	tion,
beltran 2016 HCMV 96	1868	6	9	72 hpi 76.59 Data from Beltran et al. 2016 [19] HCMV infecti 96 hpi	tion,
beltran2016MOCK120	1757	6	9	71.84 Data from Beltran et al. 2016 [19] MOCK, 120 h	hpi
beltran2016MOCK24	2220	6	9	80.12 Data from Beltran et al. 2016 [19] MOCK, 24 hp	-
beltran2016MOCK48	2181	6	9	68.90 Data from Beltran et al. 2016 [19] MOCK, 48 hp	-
beltran2016MOCK72	2161	6	9	68.83 Data from Beltran et al. 2016 [19] MOCK, 72 hp	ıpi
beltran2016MOCK96	1748	6	9	73.24 Data from Beltran et al. 2016 [19] MOCK, 96 hp	ıpi
dunkley2006	689	16	9	86.70 LOPIT data from Dunkley et al. (2006) [7]	
foster2006	1555	26	8	53.13 PCP data from Foster et al. (2006) [8]	
nikolovski2014	1385	20	8	67.97 LOPIMS data from Nikolovski et al. (2014) [22]]
groen2014cmb	424	18	7	64.30 LOPIT experiments on Arabidopsis thaliana roc	oots,
				from Groen et al. (2014) [13]	
nikolovski2012imp	1385	32	7	77.82 Meta-analysis from Nikolovski et al. (2012) [21]	
andreyev2010rest	2642	36	6	25.39 Six sub-cellular fraction data from mo	ouse
				macrophage-like RAW264.7 cells from Andreyev al. (2009) [1]	ev et
hall2009	1090	16	5	63.45 LOPIT data from Hall et al. (2009) [14]	

Table 1: Summary of the datasets used in this study. The percentage of variance along the principal components (PC) is related to the PCA plots on figure 12. All datasets are available in the pRolocdata package.

We have applied minimal post-processing to the data and have used, as far as possible, the data and annotation provided by the original authors. The data from Foster et al. [8] have been annotated using the curated marker list from Christoforou et al. [4], as only a limited number of markers were provided by the authors¹. Marker proteins are well-known and trusted, generally manually curated residents of sub-cellular niches, for the species and condition of interest, and are used to annotate the spatial proteomics experiment. These annotations are then used for visualisation and quality control (see section 3.2) and supervised machine learning.

We have also only considered sub-cellular niches (also referred to as protein clusters, or clusters) that were defined by at least 7 protein markers². We used combined dataset of multiple replicated experiments, when provided by the original authors, rather than individual replicates, as combining data often leads to better sub-cellular resolution [30]. In addition, for dimensionality reduction and visualisation, we have systematically replaced missing values by zeros. When calculating distances between protein profiles (see section 3.3), however, missing values were retained.

It is important to highlight that not all experiments used in this study have as main goal the generation of a global (or near global) sub-cellular map. While the works of Dunkley et al. [7] (Mapping the Arabidopsis organelle proteome), Tan et al. [26] (Mapping organelle proteins and protein complexes in Drosophila melanogaster) and more recently Christoforou et al. [4] (A draft map of the mouse pluripotent stem cell spatial proteome) and Itzhak et al. [17] (Global, quantitative and dynamic mapping of protein subcellular local*ization*) explicitly state such goal, other experiments such as Groen et al. [13] (Identification of trans-golgi network proteins in Arabidopsis thaliana root tissue) or Nikolovski et al. [22] (Label free protein quantification for plant Golqi protein localisation and abundance) have a more targeted goal (identification of trans-Golgi -network and Golgi apparatus proteins, respectively). When an experiment is targeted at resolving a particular niche, it is often the case that other sub-compartments are less well-resolved and hence, it is important to keep the overall aim of the studies in mind when assessing their overall resolution.

¹This results from the fact that they used a simple distance measurement, termed χ^2 , against very few markers to base their sub-cellular localisation prediction

²This number is relatively low, and we would typically recommend at least 13 markers per class to perform cross-validation when optimising classifier parameters. See Gatto et al. [11] and the main pRoloc tutorial for details.

3 Assessment

While never performed in a systematic way as in this work, authors have provided some metrics to demonstrate the quality of their data. One such metric is the macro-F1 score, calculated during classification parameter optimisation (see the pRoloc [12] documentation or [11] for details about this procedure). Briefly, the macro-F1 is computed as the harmonic mean of the precision (a measure of exactness) and recall (a measure of completeness) on marker proteins to infer a set of credible model parameters to be used subsequently, when inferring the localisation of proteins of unknown location. The scores are computed during a number of iteration, where some markers are used as validation test, and other are used for parameter selection using cross-validation. These optimisation metrics are computed over a range of parameters on different subsets of marker proteins, and optimal parameters are over-optimised for the subset of markers used at each iteration, and hence are likely to provide an over-fitted view of the data. Note that while each iteration is over-fitted, the goal of the iterative procedure is to identify recurring best parameters, in the hope that these will subsequently generalise to new, non-marker proteins.

Itzhak et al. [17] have used the Pearson correlation between replicates to demonstrate the replicability of their experiments. While useful within an experiment, correlation cannot be compared between experiments, as the precision of these values will depend on the number of proteins in the experiment. More generally, correlation isn't a good measure of reproducibility [16]; it focuses on values without context. A possible better approach would be to compare classification results and demonstrate that these agree across different replicates.

3.1 Sub-cellular diversity

A first assessment that provides an important indication of the resolution of the data concerns the number and diversity of sub-cellular niches that are annotated. In the 29 datasets used in this study, this number ranged from 5 (dataset hall2009) to 14 (dataset hyperLOPIT2015). These numbers should be assessed in the light of about 25 different sub-cellular niches that are documented in all 29 datasets, which are still underestimating the biological sub-cellular diversity.

3.2 Dimensionality reduction and visualisation

Principal component analysis (PCA) is a widely used dimensionality reduction technique in spatial proteomics. It projects the protein occupancy profiles into a new space in such a way as to maximise the spread of all points (i.e. marker proteins and proteins of unknown localisation) along the first new dimension (principal component, PC). The second PC is then chosen to be perpendicular to the first one while still maximising the remaining variability, and so on. Each PC accounts for a percentage of the total variability and it is not uncommon, in well executed experiments, that the two first PCs summarise over 70% of the total variance in the data, confirming that the resulting visualisation remains a reliable and useful simplification of the original, multidimensional data.

By firstly summarising the occupancy profiles along PC1 and PC2 (and, possibly, other PCs of interest if necessary), it becomes possible to visualise the complete dataset in a single figure (as opposed to individual sets of profiles - see for example figure 5 in Gatto et al. [10]). In a first instance, it is advised to visualise the data without marker annotation to confirm the presence of discrete clusters, i.e. dense clouds of points that are well separated from the rest of the data (see for example data from Christoforou et al. [4] on figure 1, left). Such patterns can further be emphasised by using transparency (figure 1, centre) or binned hexagon plots (figure 1, right) to highlight density.

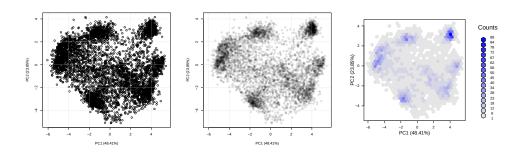


Figure 1: Unsupervised visualisation of spatial resolution using the plot2D function from the pRoloc package.

In figure 2 we compare three datasets to illustrate different levels of cluster density and separation. We see areas of high density (many proteins per hexagon) are highlighted by dark blue bins, and less dense areas are white/grey, as indicated by the count key on the right-hand side of each

plot. The figure on the left is the hyperLOPIT data from Christoforou et al. [4] (as on figure 1) that used synchronous precursor selection (SPS) MS³ on an Orbitrap Fusion. The middle figure represents the same experiment and same proteins, analysed using conventional MS², illustrating the effect of reduced quantitation accuracy. Finally, on the right, an experiment with considerable less resolution [14].

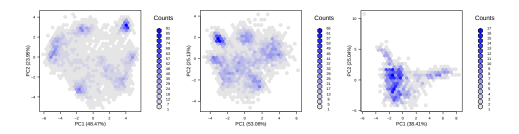


Figure 2: Comparing the cluster density and separation of experiments with excellent (left), intermediate (centre) and poor (right) resolution.

Considering that the aim of sub-cellular fractionation is to maximise separation of sub-cellular niches, one would expect sub-cellular clusters to be separated optimally in a successful spatial proteomics experiment. In PCA space, this would equate to the location of the marker clusters along the periphery of the data points. In other words, the maximum variability of a successful spatial proteomics experiments should be reflected by the separation of genuine (i.e. expected/annotated marker) spatial clusters, as illustrated on figure 3.

Another dimensionality reduction method that is worth mentioning here is linear discriminant analysis (LDA). LDA will project the protein occupancy profiles in a new set of dimensions using as a criterion the separation of marker classes by maximising the between class variance to the within class variance ratio. As opposed to the *unsupervised* PCA, the *supervised* LDA should not be used as an experiment quality control, but can be useful to assess if one or more organelles have been preferentially separated.

It is important to highlight that these representations, while generally reflecting a major proportion of the variability in the data, are only a summary of the total variability. Some sub-cellular niches that overlap in 2 dimensions

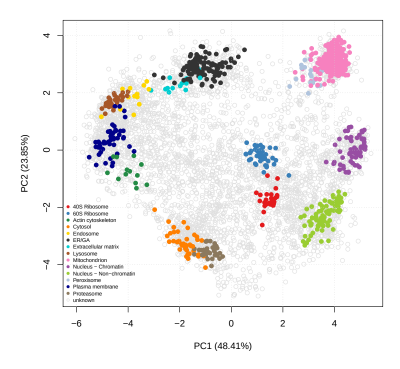


Figure 3: Annotated PCA plot of the hyperLOPIT2015 dataset.

can be separated along further components. It is sometimes useful to visualise data in three dimensions (using for example the plot3D function in the pRoloc package), which still, however, only reflect part of the total variability. When assessing the resolution of some organelles of interest, one should compare the full protein profiles of the marker proteins (pRoloc's plotDist function can be used for this, or the interactive application pRolocVis in the pRolocGUI package) or visualise a dendrogram representing the average distance between cluster profiles (the mrkHClust function from pRoloc offers this functionality). While detailed exploration of a dataset using these and other visualisation approaches is crucial before analysing and interpreting a new spatial proteomics experiment, a detailed exploration of each of the 29 datasets used in this meta-analysis is out of the scope of this work.

3.3 Quantifying resolution

While visualisation of spatial proteomics data remains essential to assess the resolution, and hence the success, of a spatial proteomics experiment, it is useful to be able to objectively quantify the resolution and directly compare different experiments. Here, we present a new infrastructure, termed QSep, available in the pRoloc package [12], to quantify the separation of clusters in spatial proteomics experiments. It relies on the comparison of the average euclidean distance within and between sub-cellular marker clusters. As illustrated on the heatmaps in figure 4 for the hyperLOPIT2015 data, these distances always refer to one reference marker cluster.

The raw distance matrix (figure 4, top-left) is symmetrical (i.e. the distance between cluster 1 and 2 is the same as between cluster 2 and 1). Within cluster distances, along the diagonal, are generally the smallest ones, except when two clusters overlap, as the lysosome and endosome in our example. To enable the comparison of these distances within and between experiments (see section 4 for the latter), we further divide each distance by the reference within cluster average distance (figure 4, top-right). This thus informs us as how much the average distance between cluster 1 and 2 is greater than the average distance within cluster 1 (i.e. the tightness of that cluster). At this stage, the distance matrix is no longer symmetrical. To facilitate the comparison of distances between organelles, the distance distributions can also be visualised as boxplots (figure 4, bottom).

The rational behind these measures is as follows. Intuitively, we assess resolution by contrasting the separation between clusters (formalised by the average distance between two clusters) and the tightness of single clusters (formalised by the average within cluster distance). Ideal sub-cellular fractionation would yield tight and distant clusters, represented by a large normalised between cluster distances on figure 4.

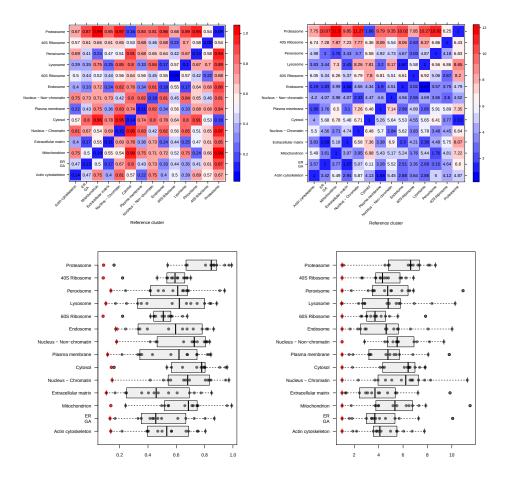


Figure 4: Quantifying resolution of the *hyperLOPIT2015* data Christoforou et al. [4]. The heatmaps at the top illustrate the raw (left) and average normalised (right) within (along the diagonal) and between euclidean cluster distances. The boxplots at the bottom summarise these same values (raw on the left, normalised on the right) to enable easier comparison between clusters, where the within distances are highlighted in red.

3.4 Application of the assessment criteria

To further demonstrate the interpretation of these resolution metrics, we directly compare the two recent global cell maps from [4] (dataset hyper-LOPIT2015) and [17] (dataset itzhak2016stcSILAC). Both feature high protein coverage (7114 and 5265 proteins respectively) and good sub-cellular diversity (14 and 12 annotated clusters respectively). The former contains duplicated experiments, each made of 10 fractions and the latter contains 6 replicates with 5 fractions each. Figure 5 shows the PCA plots applying transparency to identify the underlying structure in the quantitative data and the annotated versions using the markers provided by the respective authors.

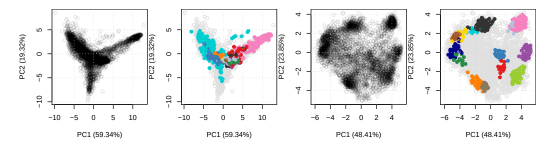


Figure 5: PCA plots for itzhak2016stcSILAC (left) and hyperLOPIT2015 (right). Here, we display PC 1 and 2 for both datasets for comparability. The original authors displayed PC 1 and 3 for the itzhak2016stcSILAC data (see figures 11 and 12 below).

Figure 6 illustrates the normalised distance heatmaps and boxplots for the two datasets (*itzhak2016stcSILAC* at the top and *hyperLOPIT2015* at the bottom). The two heatmaps display strikingly different colour patterns. The top heatmap shows a majority of small normalised distances (blue cells) and with only a limited number of large distances (red cells), along the mitochondrial reference cluster. Conversely, the bottom heatmap displays a majority of average (white cells) and large distances (red cells) across all sub-cellular clusters. The boxplots allow a more direct comparison of the distances across the two datasets. On the top boxplot, we detect relatively short distances for most clusters, with most large distances stemming from the mitochondria, leading to a median distance of 2.48. The distributions on the bottom boxplot show larger distances, equally spread among all clusters, with an median distance of 4.91.

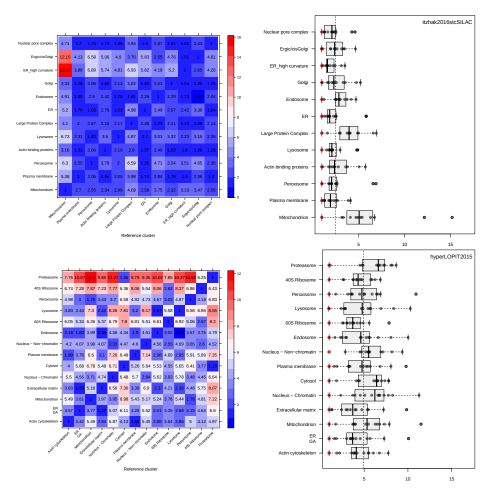


Figure 6: Contrasting quantitative separation assessment between the itzhak2016stcSILAC [17] (top) and hyperLOPIT2015 [4] (bottom) datasets. The dashed vertical lines on the boxplots represent the overall media between cluster distance, 2.48 and 4.91 for itzhak2016stcSILAC and hyperLOPIT2015 respectively.

4 Comparative study

We next apply the quantitative assessment of spatial resolution described in section 3.3 to compare the 29 experiments presented in section 2. Figure 7 shows, for each dataset, a boxplot illustrating the distribution of the global average normalised distances for all spatial clusters. The datasets have been ordered using the experiment-wide median between distance. It is important to always refer back to the original data when considering summarising metrics like these, to put the resolution into context; the density and annotated PCA plots discussed in section 3.2 are provided in figures 11 and 12 and the quantitative assessment boxplots and heatmaps are shown in figures 13 and 14.

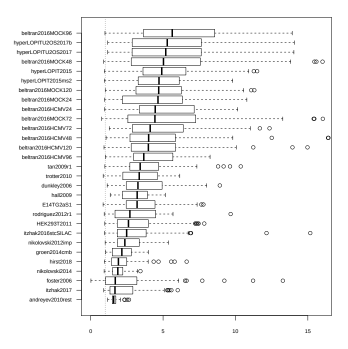


Figure 7: Quantitative separation assessment using experiment-wide normalised distances between cluster distances. The vertical line represents the normalised intra-cluster distance of 1.

The individual LOPIT-based experiments from Jean Beltran et al. [19] and the embryonic mouse stem cell [4] and U2OS [28] hyperLOPIT experiments (using SPS MS³ and conventional MS²) show the best global,

experiment-wide resolution. The next set of experiment are tan2009r1, trotter2010, dunkley2006, hall2009 and E14TG2aS1. It is important to highlight that most of datasets (as well as HEK293T2011, discussed later) have either been directly re-analysed using a semi-supervised novelty detection algorithm phenoDisco [2] (the only exception here being hall2009), or, in the case of trotter 2010, have been annotated using markers based on the phenoDisco re-analysis. The novelty detection algorithm, phenoDisco, searches for new clusters of unlabelled proteins, using the marker proteins to guide the clustering of unlabelled features. These new clusters, termed phenotypes, are then validated by the user for coherence with known sub-cellular niches. This re-analysis has proven successful [2] and has identified previously undetected sub-cellular niches that form tight and well-resolved clusters (see for example ribosomial and trans-Golgi network (TGN) in dunkley2006, or proteasome and nucleus in tan2009r1 to cite only a few), which in turn favour good resolution scores. The hall 2009 dataset is relatively poorly annotated (only 5 sub-cellular clusters, which is the lowest in all test datasets). As long as these few clusters are well separated, poor annotation will however not negatively influence the resolution scoring.

The next set of experiments that show comparable resolution profiles are HEK293T2011, itzhak2016stcSILAC and nikolovski2012imp. Note that the quantitative separation measurement is robust to questionable marker annotation. For example, the $Large\ Protein\ Complex$ class defined by the original authors in the itzhak2016stcSILAC data could be dropped as it loosely defines many niches and thus lacks resolution. This omission only marginally influences the overall assessment metrics as only the distances to/from that class are affected (i.e. 23 out of 144 distances) and as such it would not change its rank among the test datasets.

As mentioned earlier, the *groen2014cmb* and *nikolovski2014* are targeted experiments, focusing on the trans-Golgi-network and Golgi niches respectively. Such experiments do not aim for the best global resolution, which is reflected by relatively low resolution.

The foster2006 experiment displays relatively poor separation. This might be due to the relatively high number of missing values (42.4 %). Finally, the andreyev2010rest dataset suffers from very broad sub-cellular clusters (compared to separation between clusters).

The PCA plots and QSep heatmaps for all datasets are provided in the appendix, section A.

5 Assessing the resolution metric

In this section, we assess the resolution metric, and how the annotation of the spatial proteomics data influences the metric itself.

We find that the number of classes does not have any effect the resolution assessment scoring. Indeed, dropping any class will result in a sub-sample of normalised inter-cluster distances, with random variations around the overall median inter-cluster distance. On figure 8, we show the distribution of the resolution metrics when removing all possibly combinations of 1 to 3 sub-cellular classes for the E14TG2aS1 dataset, that displays an average overall resolution, and hyperLOPIT2015, that has among the highest resolution. In both cases, we see that the number of removed classes does not influence the overall score distributions. When modelling the linear relation between the median scores and the number of removed classes, the slopes are 0.036 and 0.01 respectively.

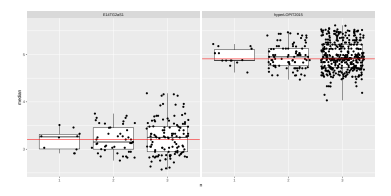


Figure 8: Effect of removing sub-cellular clusters on the resolution metric for the E14TG2aD1 (left) and hyperLOPIT2015 experiments (right). Each dot represents a median resolution score for the experimental setting (i.e missing n classes). The horizontal lines represents the median resolution metrics for the complete dataset. Note the overall higher median assessment scores for the better hyperLOPIT2015 experiment

The definition of marker proteins has of course an effect on the assessment metric. Tighter clusters will result in smaller intra-class distances and, as a result, in larger normalised inter-class distances. To illustrate the effect of marker definition, we transferred the marker annotation between the hyperLOPIT2015 and itzhak2016stcSILAC datasets (see PCA plots on fig-

ure 9, left) and calculated the quantitative resolution metrics (see boxplots on figure 9, right).

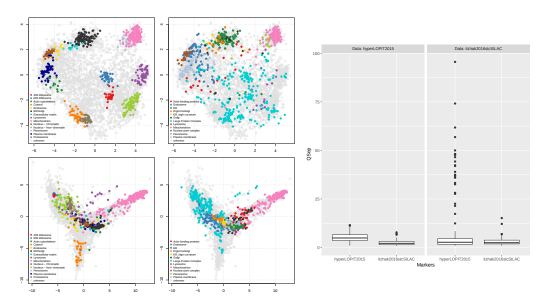


Figure 9: Marker transfer between hyperLOPIT2015 and itzhak2016stcSILAC. On the left, the 4 data/marker combinations are displayed on PCA plots: the top and bottom row contains the hy-perLOPIT2015 and itzhak2016stcSILAC data respectively, while the left and right columns display the markers from hyperLOPIT2015 and itzhak2016stcSILAC respectively. On the right, the resolution scores have been calculated for the same markers (along the x axis) data (left and right panels).

As expected from the PCA plots on figure 9 and the diffuse Large Protein Complex cluster, and as can be seen on QSep distribution boxplots, transferring the itzhak2016stcSILAC markers to the hyperLOPIT2015 dataset (distributions in the left panel) has a detrimental effect on the separation (testing the log-transformed distributions with a t-test produces a p-value of 4.3×10^{-32}). Annotating itzhak2016stcSILAC with the hyperLOPIT2015 markers (distributions in the right panel) hardly improves its resolution metric (p-value of 0.0015, log-transformed QSep scores). The main effect here is to emphasise the separation between the mitochondrion and other spatial niches, in particular the very tight 40S and 60S ribosomal clusters. These examples illustrate the importance and impact of marker curation and anno-

tation for individual experiments. In particular, the *Large Protein Complex* cluster from *itzhak2016stcSILAC*, while also diffuse in its original dataset, has a severe effect on a dataset that it was not curated for.

6 Conclusions

In this manuscript, we have described in great detail how to assess and quantify the resolution of spatial proteomics experiments. We have applied dimensionality reduction and visualisation, as well as a simple and intuitive quantitative metric to explore and compare a variety of publicly available spatial proteomics datasets using the annotation provided by the original authors. We have also assessed the resolution metric itself and observed that it was immune to the number of clusters used for its computation and showed the possible influence of different marker annotation on the metric itself.

The ordering of the quantitative resolution detailed in section 4 should not be taken as absolute. Its main purpose is to provide a guide to compare different experiments. It will be useful for laboratories that do spatial studies on different models and with different fractionation and/or quantitation methods, to assess the impact of these variables (such as, for example hyperLOPIT MS² and MS³). It is also useful to compare separation between different labs, as demonstrated in our comparative study (section 4). We anticipate that it will also prove useful for the researcher wanting to assess the resolution of newly published studies, and put them into a wider context. It is necessary to emphasise that the importance and effect of marker definition on estimating and assessing the resolution of spatial proteomics experiments (section 5) and, of course, the subsequent assignment of proteins to their most likely sub-cellular compartments. Sub-cellular resolution is of course only one aspect of spatial proteomics, albeit am important one, that critically determines the reliability of protein assignments to spatial niches as well as the identification of multi- and trans-localisation events.

Finally, we reflect on the implications of this work on the spatial proteomics community, and more generally the cell biology community that relies on protein localisation data. We have assessed dataset spanning 12 years of spatial proteomics. Since 2006, the community has seen many important improvements: tremendous advances in mass spectrometry, improvements in spatial proteomics designs, and considerable breakthroughs in data anal-

ysis. One might then wonder whether these benefits have lead to tangible improvements in resolution over time?

On figure 10, we have ordered the datasets' resolution metric according to their publication year. We can see that a set of recent datasets, including the mouse stem cell [4] and U2OS [28] hyperLOPIT experiments (published in 2016 and 2017 respectively), and a variation thereof, where Jean Beltran et al. [19] (published in 2016), incorporating a temporal component in their experimental design, show a consistent superior resolution.

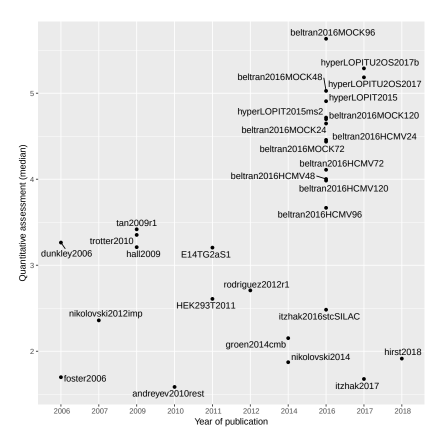


Figure 10: Resolution of spatial proteomics experiments over time. The assigned dates match either the original publication or, when know, the actual date the data was generated. For publications that re-analysed data, the date of the original publication of generation was used. When publications used several datasets, the date of the most recent one was used.

While the definition of sub-cellular resolution, as defined by the QSep

measure, is only one aspect of spatial proteomics, one could argue that the community at large would benefit from a more systematic approach when considering the resolution of spatial proteomics experiments. Indeed, there are various aspects that can be worked on to improve resolution: quantitation accuracy at the mass spectrometry level (see figures 1 and 2 comparing SPS MS³ and conventional MS² for an example), optimisations in sub-cellular fractionation (as exemplified by the substantial improvement obtained by [4]), improved data annotation (see for example figure 9), as well as superior data analysis (for example using semi-supervised learning Breckels et al. [2]).

As already suggested [20], there is arguably a need for standardisation, or for general guidelines in assessing spatial proteomics data in the community? Shouldn't the community as a whole aim for collective improvement and some agreement as to what constitutes a good spatial proteomics experiment and a reliable protein sub-cellular assignment? The latter can assessed using improved probabilistic classifiers such as the Bayesian mixture modelling approach proposed by Crook et al. [6]. In this work, we propose the QSep metric to assess the former. Better spatial proteomics data and more reliable interpretation will be of direct benefit to the spatial proteomics researchers themselves, and will increase the trust and reliance of the cell biology community.

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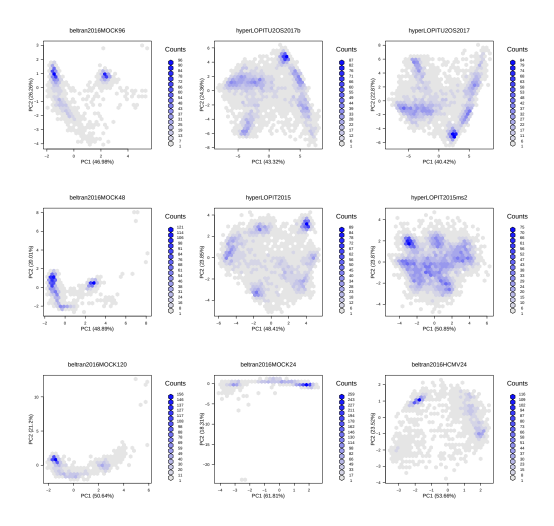
Author contributions

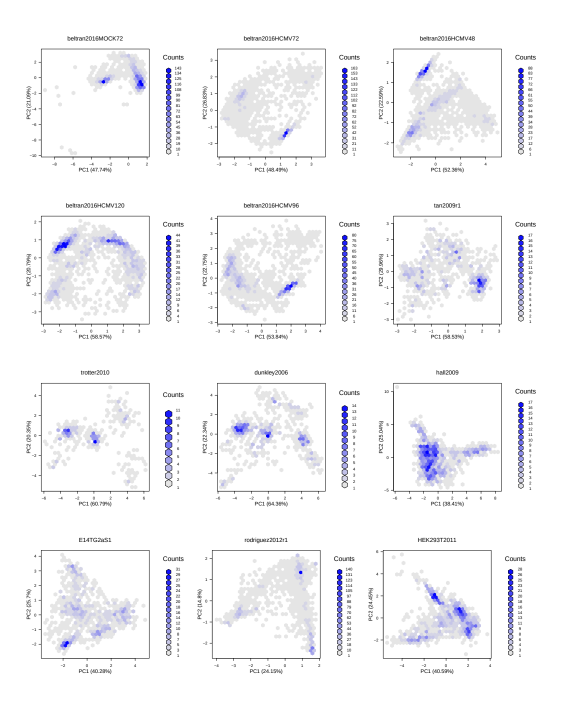
LG conceptualised the method and wrote the initial manuscript draft. LG and LMB developed the QSep code. KSL contributed datasets and feedback. All authors read and approved the manuscript.

Appendices

A Additional figures

This section shows the density and annotated PCA and QSep plots for the 29 datasets showcased in this manuscript.





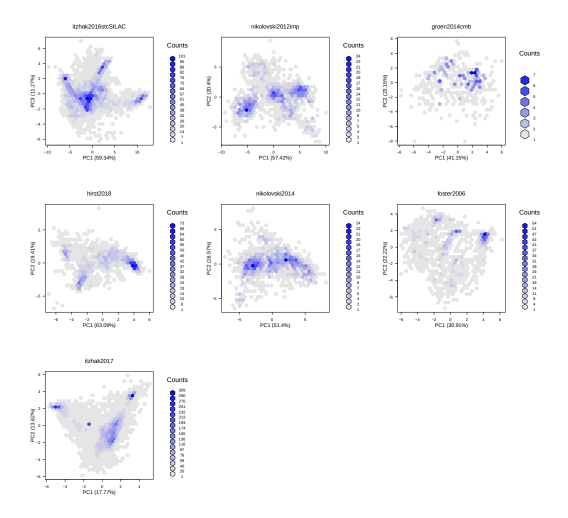
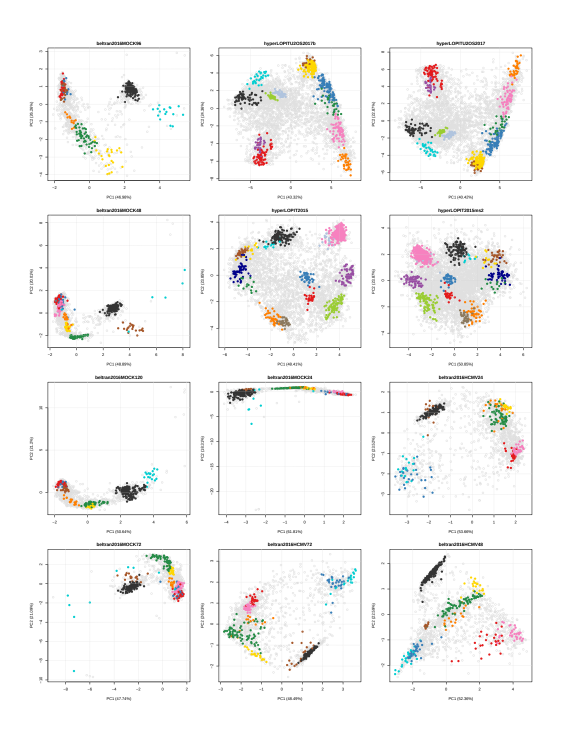
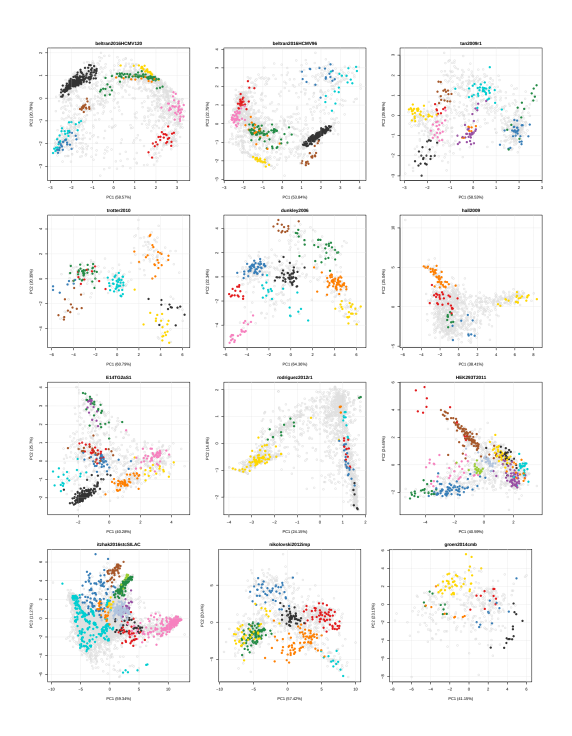


Figure 11: Density PCA plots for the 29 experiments used in this study. PC 1 and 2 were used except for *itzhak2016stcSILAC*, where PC 1 and 3 were used to conform to the original authors figures. The experiments are ordered according to the median average between cluster distance (see figure 7). Figures have been generated using the plot2D function from the pRoloc package.





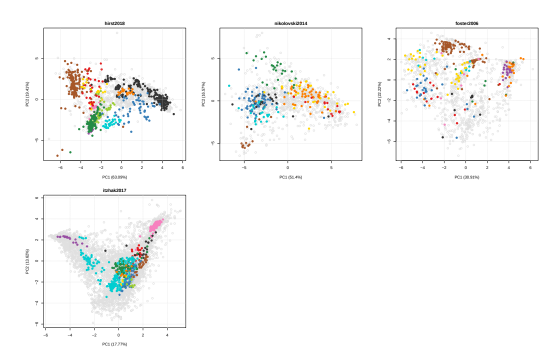
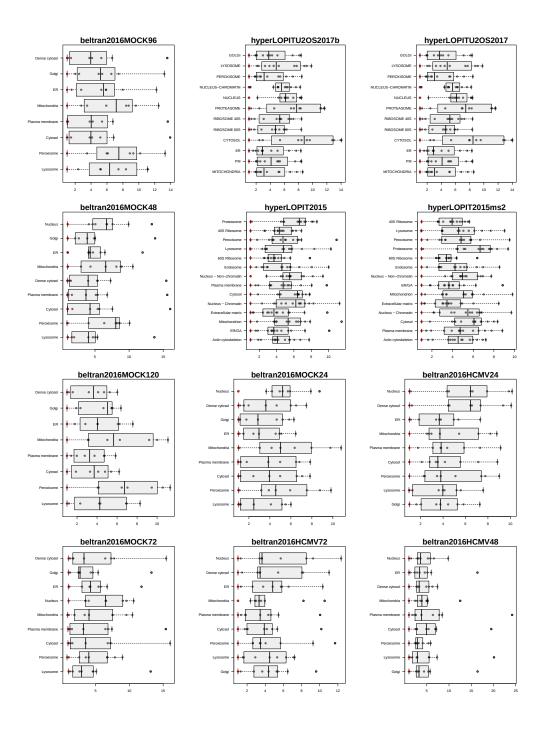


Figure 12: PCA plots for the 29 experiments used in this study. PC 1 and 2 were used except for itzhak2016stcSILAC, where PC 1 and 3 were used to conform to the original authors figures. The experiments are ordered according to the median average between cluster distance (see figure 7). The percentage of variance explained along the 2 PCs on the plots can be found in table 1. Figures have been generated using the plot2D function from the pRoloc package.





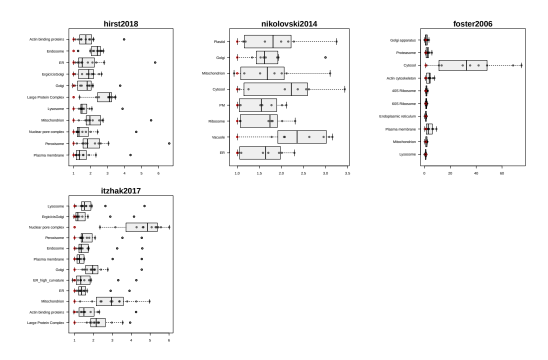
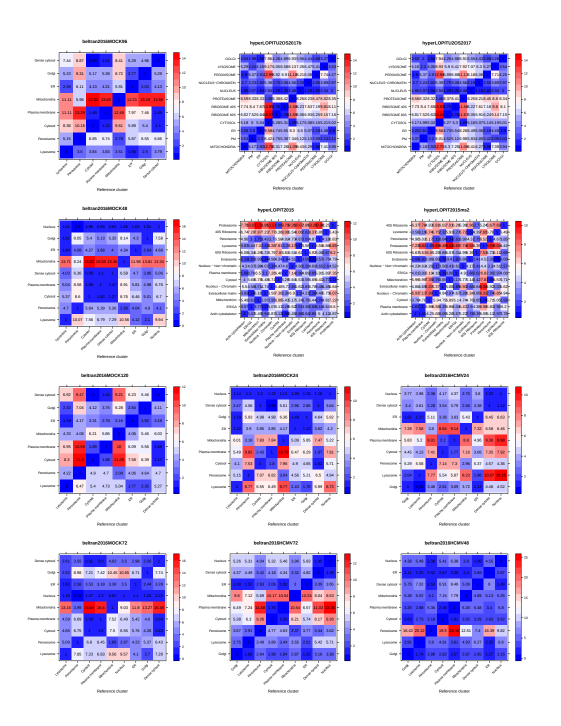
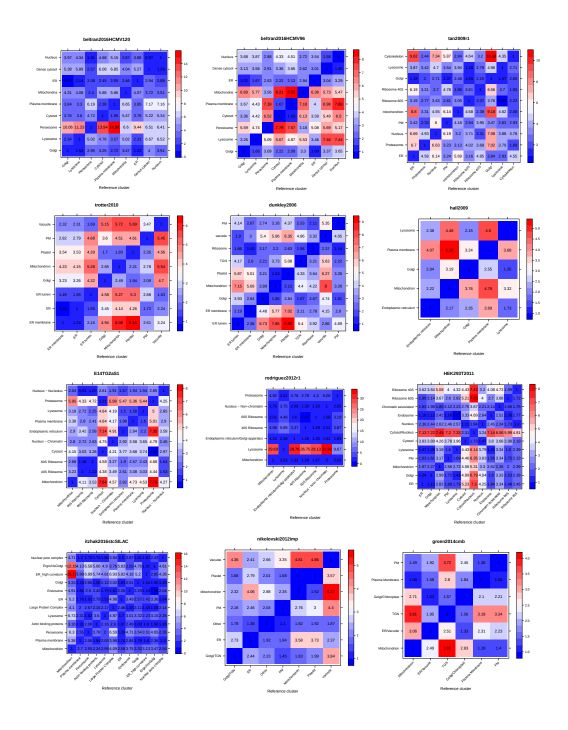


Figure 13: Quantitative separation boxplot for the 29 experiments used in this study. The experiments are ordered according to the median average between cluster distance (see figure 7).





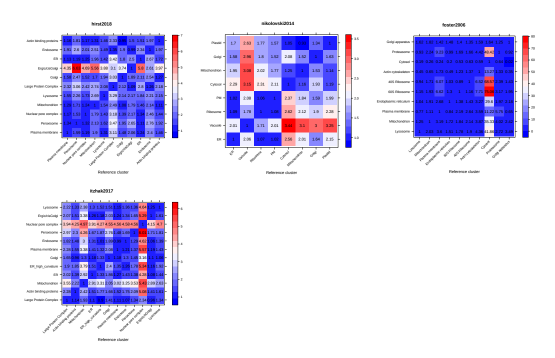


Figure 14: Quantitative separation heatmaps for the 29 experiments used in this study. The experiments are ordered according to the median average between cluster distance (see figure 7).

B Session information

The software and versions used to produce this document are summarised below. The source of this document enabling to reproduce all results and figures is available in the source of this document in the public manuscript repository [9] available at https://github.com/lgatto/QSep-manuscript/.

- R Under development (unstable) (2018-04-02 r74505), x86_64-pc-linux-gnu
- Running under: Ubuntu 14.04.5 LTS
- Matrix products: default
- BLAS: /usr/lib/atlas-base/atlas/libblas.so.3.0
- LAPACK: /usr/lib/lapack/liblapack.so.3.0
- Base packages: base, datasets, graphics, grDevices, methods, parallel, stats, stats4, utils
- Other packages: annotate 1.58.0, AnnotationDbi 1.42.1, Biobase 2.40.0, BiocGenerics 0.26.0, BiocParallel 1.14.2, cluster 2.0.7-1, ggplot2 3.0.0, ggrepel 0.8.0, hexbin 1.27.2, IRanges 2.14.10, MLInterfaces 1.60.1, MSnbase 2.7.3, mzR 2.15.2, pRoloc 1.21.7, pRolocdata 1.19.3, ProtGenerics 1.12.0, Rcpp 0.12.18, S4Vectors 0.18.3, XML 3.98-1.12, xtable 1.8-2
- Loaded via a namespace (and not attached): abind 1.4-5, affy 1.58.0, affyio 1.50.0, assertthat 0.2.0, backports 1.1.2, base64enc 0.1-3, bindr 0.1.1, bindrcpp 0.2.2, BiocInstaller 1.30.0, biomaRt 2.36.1, bit 1.1-14, bit64 0.9-7, bitops 1.0-6, blob 1.1.1, broom 0.5.0, caret 6.0-80, class 7.3-14, coda 0.19-1, codetools 0.2-15, colorspace 1.3-2, compiler 3.6.0, crayon 1.3.4, crosstalk 1.0.0, CVST 0.2-2, DBI 1.0.0, ddalpha 1.3.4, dendextend 1.8.0, DEoptimR 1.0-8, digest 0.6.15, dimRed 0.1.0, diptest 0.75-7, doParallel 1.0.11, dplyr 0.7.6, DRR 0.0.3, e1071 1.6-8, evaluate 0.11, flexmix 2.3-14, FNN 1.1, foreach 1.4.4, fpc 2.1-11.1, gbm 2.1.3, gdata 2.18.0, genefilter 1.62.0, geometry 0.3-6, ggvis 0.4.3, glue 1.3.0, gower 0.1.2, grid 3.6.0, gridExtra 2.3, gtable 0.2.0, gtools 3.8.1,

highr 0.7, hms 0.4.2, htmltools 0.3.6, htmlwidgets 1.2, httpuv 1.4.5, httr 1.3.1, hwriter 1.3.2, igraph 1.2.1, impute 1.54.0, ipred 0.9-6, iterators 1.0.10, kernlab 0.9-26, knitr 1.20, labeling 0.3, Laplaces Demon 16.1.1, later 0.7.3, lattice 0.20-35, lava 1.6.2, lazyeval 0.2.1, limma 3.36.2, lpSolve 5.6.13, lubridate 1.7.4, magic 1.5-8, magrittr 1.5, MALDIquant 1.18, MASS 7.3-50, Matrix 1.2-14, mclust 5.4.1, memoise 1.1.0, mime 0.5, mixtools 1.1.0, mlbench 2.1-1, ModelMetrics 1.1.0, modeltools 0.2-22, munsell 0.5.0, mytnorm 1.0-8, mzID 1.18.0, nlme 3.1-137, nnet 7.3-12, pcaMethods 1.72.0, pillar 1.3.0, pkgconfig 2.0.1, pls 2.6-0, plyr 1.8.4, prabclus 2.2-6, preprocessCore 1.42.0, prettyunits 1.0.2, prodlim 2018.04.18, progress 1.2.0, promises 1.0.1, proxy 0.4-22, purrr 0.2.5, R6 2.2.2, randomForest 4.6-14, RColorBrewer 1.1-2, ReppRoll 0.3.0, RCurl 1.95-4.11, rda 1.0.2-2.1, recipes 0.1.3, reshape2 1.4.3, rlang 0.2.1, robustbase 0.93-1.1, rpart 4.1-13, RSQLite 2.1.1, sampling 2.8, scales 0.5.0, segmented 0.5-3.0, sfsmisc 1.1-2, shiny 1.1.0, splines 3.6.0, stringi 1.2.4, stringr 1.3.1, survival 2.42-4, threejs 0.3.1, tibble 1.4.2, tidyr 0.8.1, tidyselect 0.2.4, timeDate 3043.102, tools 3.6.0, trimcluster 0.1-2.1, viridis 0.5.1, viridisLite 0.3.0, vsn 3.48.1, whisker 0.3-2, with 2.1.2, zlibbioc 1.26.0

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