Integrated view and comparative analysis of baseline protein expression in mouse and rat tissues

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33 Keywords

- 34 Mass spectrometry, quantitative proteomics, comparative protein expression, public data
- 35 reuse, rat proteome, mouse proteome, PRIDE database

36 Abstract

37	The increasingly large amount of proteomics data in the public domain enables, among other
38	applications, the combined analyses of datasets to create comparative protein expression
39	maps covering different organisms and different biological conditions. Here we have
40	reanalysed public proteomics datasets from mouse and rat tissues (14 and 9 datasets,
41	respectively), to assess baseline protein abundance. Overall, the aggregated dataset contained
42	23 individual datasets, including a total of 211 samples coming from 34 different tissues
43	across 14 organs, comprising 9 mouse and 3 rat strains, respectively.
44	
45	In all cases, we studied the distribution of canonical proteins between the different organs.
46	The number of canonical proteins per dataset ranged from 273 (tendon) and 9,715 (liver) in
47	mouse, and from 101 (tendon) and 6,130 (kidney) in rat. Then, we studied how protein
48	abundances compared across different datasets and organs for both species. As a key point
49	we carried out a comparative analysis of protein expression between mouse, rat and human
50	tissues. We observed a high level of correlation of protein expression among orthologs
51	between all three species in brain, kidney, heart and liver samples, whereas the correlation of
52	protein expression was generally slightly lower between organs within the same species.
53	Protein expression results have been integrated into the resource Expression Atlas for
54	widespread dissemination.
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56 Author summary

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58 We have reanalysed 23 baseline mass spectrometry-based public proteomics datasets stored 59 in the PRIDE database. Overall, the aggregated dataset contained 211 samples, coming from 60 34 different tissues across 14 organs, comprising 9 mouse and 3 rat strains, respectively. We 61 analysed the distribution of protein expression across organs in both species. We also studied 62 how protein abundances compared across different datasets and organs for both species. Then 63 we performed gene ontology and pathway enrichment analyses to identify enriched biological 64 processes and pathways across organs. We also carried out a comparative analysis of baseline 65 protein expression across mouse, rat and human tissues, observing a high level of expression 66 correlation among orthologs in all three species, in brain, kidney, heart and liver samples. To 67 disseminate these findings, we have integrated the protein expression results into the resource 68 Expression Atlas.

69 **1. Introduction**

70

71	High-throughput mass spectrometry (MS)-based proteomics approaches have matured
72	significantly in recent years, becoming an essential tool in biological research [1]. This has
73	been the consequence of very significant technical improvements in MS instrumentation,
74	chromatography, automation in sample preparation and computational analyses, among other
75	areas. The most used MS-based experimental approach is Data Dependent Acquisition
76	(DDA) bottom-up proteomics. Among the main quantitative proteomics DDA techniques,
77	label-free intensity-based approaches remain very popular, although labelled-approaches,
78	especially those techniques based on the isotopic labelling of peptides (MS ² labelling), such
79	as iTRAQ (Isobaric tag for relative and absolute quantitation) and TMT (Tandem Mass
80	Tagging), are becoming increasingly used as well.
81	
82	Following the steps initiated by genomics and transcriptomics, open data practices in the field
83	have become embedded and commonplace in proteomics in recent years. In this context,

84 datasets are now commonly available in the public domain to support the claims published in

85 the corresponding manuscripts. The PRIDE database [2], located at the European

86 Bioinformatics Institute (EBI), is currently the largest resource worldwide for public

87 proteomics data deposition. PRIDE is also one of the founding members of the global

88 ProteomeXchange consortium [3], involving five other resources, namely PeptideAtlas,

89 MassIVE, iProX, jPOST and PanoramaPublic. ProteomeXchange has standardised data

90 submission and dissemination of public proteomics data worldwide.

91

As a consequence, there is an unprecedented availability of data in the public domain, which
is triggering multiple applications [4], including the joint reanalysis of datasets (so-called

94	meta-analysis studies) [5-7]. Indeed, public proteomics datasets can be systematically
95	reanalysed and integrated e.g., to confirm the results reported in the original publications,
96	potentially in a more robust manner since evidence can be strengthened if it is found
97	consistently across different datasets. Potentially, new insights different to the aims of the
98	original studies can also be obtained by reanalysing the datasets using different strategies, this
99	includes repurposing of public datasets [8], including for instance approaches such as
100	proteogenomics studies for genome annotation purposes [9-12].

101

102 In this context of reuse of public proteomics data, PRIDE has started to work on developing 103 data dissemination and integration pipelines into popular added-value resources at the EBI. 104 This is perceived as a more sustainable approach in the medium-long term than setting up 105 new independent bioinformatics resources. One of them is Expression Atlas [13], a resource 106 that has enabled over the years easy access to gene expression data across species, tissues, 107 cells, experimental conditions and diseases. Only recently, protein expression information 108 coming from reanalysed datasets has been integrated in the 'bulk' section of Expression 109 Atlas. As a result, proteomics expression data can be integrated with transcriptomics 110 information, mostly coming from RNA-Seq experiments. So far, we have performed two 111 meta-analysis studies involving the reanalysis and integration of: (i) 11 public quantitative 112 datasets coming from cell lines and human tumour samples [13]; and (ii) 24 human baseline 113 datasets coming from 31 different organs [14].

114

The next logical step is to perform an analogous study of baseline protein expression in two of the main model organisms: *Mus musculus* and *Rattus norvegicus*. To date, there are only a small number of bioinformatics resources providing access to reanalysed MS-based quantitative proteomics datasets, and even fewer if one considers only mouse and rat data. In

119 this context, at the end of 2020, ProteomicsDB [15] released a first version of the mouse 120 proteome, based on the reanalysis of five label-free datasets. To the best of our knowledge, 121 there is no such public resource storing accurate MS-derived data for rat data yet. PaxDB is a 122 resource [16] that provides protein expression information coming from many species 123 (including mouse and rat) but the reported data relies on spectral counting, a technique that 124 generally does not provide the same level of accuracy than intensity-based label-free 125 approaches. Additionally, although antibody-based human protein expression information is 126 provided via the Human Protein Atlas [17], their efforts are focused on human protein 127 expression. 128 129 Here, we report the reanalysis and integration of 23 public mouse (14 datasets) and rat (9 130 datasets) label-free datasets, and the incorporation of the results into the resource Expression 131 Atlas as baseline studies. Additionally, we report a comparative analysis of protein 132 expression across mouse, rat and human (in this case using the results reported at [14] using 133 the same methodology).

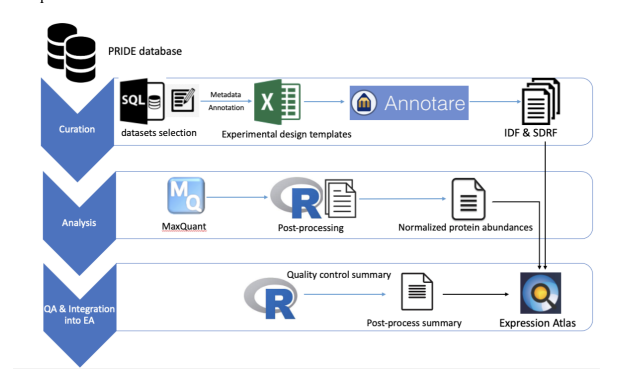
135 **2. Results**

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137 **2.1. Baseline proteomics datasets**

138 Overall, we quantified protein expression from 34 healthy tissues in 14 organs coming from 139 23 datasets. The analyses covered a total of 1,173 MS runs from 211 samples that were 140 annotated as healthy/control/non-treated samples, thus representing baseline protein 141 expression. Non-control/disease samples associated with these datasets were also reanalysed 142 but are not discussed here. Normalised protein abundances values (as ppb, parts per billion, 143 see Methods for calculation) from both control/healthy/non-treated and disease/treated tissue 144 samples are available to view as heatmaps in Expression Atlas. The protein abundances along 145 with sample annotations, sample quality assessment summary and experimental parameter 146 inputs for MaxQuant can be downloaded from Expression Atlas as text files. A summary of 147 the data selection and reanalysis protocols is shown in Fig. 1. The total number of peptides 148 and proteins identified in these datasets are shown in Table 1.



150 Figure 1. An overview of the study design and reanalysis pipeline. QA: Quality assessment.

Expression Atlas accession numbers	PRIDE dataset identifiers	Tissues	Organs	Species	Strains	Fractionation	Number of MS runs	Number of samples	Number of protein groups [†]	Number of peptides †	Number of unique peptides [†]	Number of unique genes mapped [†]
E-PROT-7§	PXD000867 ^[18]	Liver	Liver	Mus musculus	C57BL/6J	Yes	24	4	12,792	246,738	167,725	9,715
E-PROT-10 [§]	PXD000288 ^[19]	Triceps muscles	Triceps Muscles	Mus musculus	C57BL/6	Yes	36	3	10,870	189,553	126,670	6,421
E-PROT-16	PXD003155 ^[20]	Cerebellum, Liver	Brain, Liver	Mus musculus	C57BL/6	No	24	12	4,508	59,696	45,728	3,797
E-PROT-74	PXD004612 ^[21]	Achilles and Plantaris tendon	Tendon	Mus musculus	C57BL/6	No	8	8	457	6,643	3,271	273
E-PROT-75	PXD005230 ^[22]	Hippocampus, Cerebellum, Cortex	Brain	Mus musculus	C57BL/10J	Yes	72	36	7,663	63,479	41,683	6,037
E-PROT-76	PXD009909 ^[23]	Retina	Еуе	Mus musculus	ND4 Swiss Webster	Yes	12	1	5,002	29,454	24,961	3,686
E-PROT-77	PXD012307 ^[24]	Lung	Lung	Mus musculus	C57BL/6	No	32	2	6,809	106,391	73,950	5,795
E-PROT-78	PXD009639 ^[25]	Lens	Eye	Mus musculus	CD1	Yes	10	1	4,519	20,779	18,006	3,064
E-PROT-79	PXD019394 ^[26]	Heart, Kidney, Liver, Lung, Brain, Spleen, Testis, Pancreas	Heart, Kidney, Liver, Lung, Brain, Spleen, Testis, Pancreas	Mus musculus	Swiss- Webster	Yes	96	8	9,853	141,506	105,701	8,185
E-PROT-81	PXD012636 ^[27]	Left atrium, Left ventricle, Right atrium, Right ventricle	Heart	Mus musculus	C57BL/6	Yes	120	4	7,772	146,966	99,577	6,435
E-PROT-82	PXD019431 ^[28]	Articular cartilage	Articular cartilage	Mus musculus	BALB_c	No	72	6	1,815	17,695	15,191	1,518
E-PROT-83	PXD022614 ^[29]	Brain	Brain	Mus musculus	C57BL/6J: Rj C57BL/6JR ccHsd	Yes	120	6	6,645	97,443	69,884	5,673

			r	1		1	1		1	1		
E-PROT-84	PXD004496 ^[30]	Hippocampus	Brain	Mus musculus	C57BL/6J	Yes	204	17	4,192	37,363	30,100	3,424
E-PROT-85	PXD008736 ^[31]	Right atrium, Sinus node	Heart	Mus musculus	C57BL/6J	Yes	143	6	7,906	144,926	94,379	6,554
E-PROT-86 [§]	PXD012677 ^[32]	Amygdala	Brain	Rattus norvegicus	Sprague Dawley	No	3	3	1,872	15,326	12,367	1,382
E-PROT-87 [§]	PXD006692 ^[33]	Lung	Lung	Rattus norvegicus	Sprague Dawley	No	10	10	2,079	14,440	11,696	1,398
E-PROT-88 [§]	PXD016793 ^[34]	Liver	Liver	Rattus norvegicus	Sprague Dawley	No	8	8	4,787	57,998	46,411	3,743
E-PROT-89 [§]	PXD004364 ^[35]	Testis	Testis	Rattus norvegicus	Sprague Dawley	No	3	3	2,351	15,880	13,674	1,601
E-PROT-91	PXD001839 ^[36]	Left ventricle	Heart	Rattus norvegicus	F344/BN	No	12	12	1,345	10,310	8,804	925
E-PROT-92 [§]	PXD013543 ^[37]	Left ventricle	Heart	Rattus norvegicus	Wistar	No	8	8	1,858	17,303	13,622	1,340
		First segment of proximal tubule, second segment of proximal tubule, third segment of proximal tubule, medullary thick ascending limb, cortical thick ascending limb, distal convoluted tubule, connecting tubule, cortical collecting duct, outer medullary		Rattus	Sprague							
E-PROT-93	PXD016958 [38]	collecting duct	Kidney	norvegicus	Dawley	Yes	132	32	7,846	103,886	83,662	6,130

E-PROT-94	PXD003375 ^[39]	Caudal and rostral segments of spinal cord	Spinal cord	Rattus norvegicus	Wistar	Yes	21	18	2,477	29,213	22,025	1,926
E-PROT-95 [§]	PXD015928 ^[40]	Tendon	Tendon	Rattus norvegicus	Wistar	No	3	3	199	1,253	1,063	101
TOTAL	23 datasets (Mouse: 14, Rat: 9)	34 tissues (Mouse: 21, Rat: 18)	14 organs (Mouse: 12, Rat: 8)				1,173 MS runs (Mouse: 973, Rat: 200)	211 samples (Mouse: 114, Rat: 97)				

Table 1. List of mouse and rat proteomics datasets that were reanalysed. [§]Only normal/untreated samples within this dataset are reported in this

154 study. However, results from both normal and disease samples are available in Expression Atlas. † Numbers after post-processing.

155 **2.2. Protein coverage across organs and datasets**

One of our main aims was to study protein expression across various organs. To enable a simpler comparison [14] we first grouped 34 different tissues into 14 distinct organs, as discussed in 'Methods'. We defined 'tissue' as a distinct functional or structural region within an 'organ'. We estimated the number of 'canonical proteins' identified across organs by first mapping all members of each protein group to their respective parent genes. We defined the parent gene as equivalent to the UniProt 'canonical protein' and we will denote the term 'protein abundance' to mean 'canonical protein abundance' from here on in the manuscript.

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164 **2.2.1. Mouse proteome**

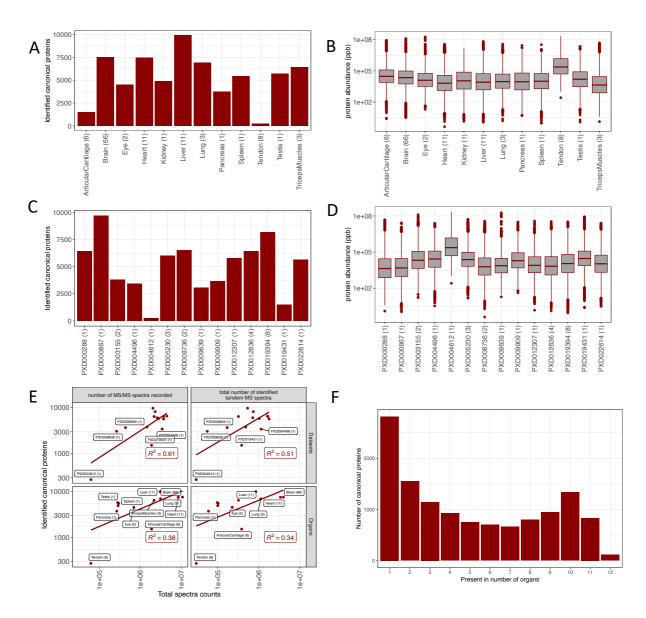
165 A total of 21,274 protein groups were identified from mouse datasets, among which 8,176 166 protein groups (38.4%) were uniquely present in only one organ and 70 protein groups 167 (0.3%) were ubiquitously observed (see the full list in Supplementary File 2). This does not 168 imply that these proteins are unique to these organs. Merely, this is the outcome considering 169 the selected datasets. Mouse protein groups were mapped to 12,570 genes (canonical 170 proteins) (Supplementary File 3). We detected the largest number of canonical proteins in 171 samples coming from liver (9,920, 78.9% of the total) and the lowest numbers in samples from tendon (273, 2.2%) and articular cartilage (1,519, 12.1%) (Fig. 2A). In the case of 172 173 tendon and articular cartilage, both experiments did not include sample fractionation in their 174 sample preparation methodology, which can also explain the lower number of detected 175 proteins. The comparatively even lower number of proteins identified in tendon could be 176 attributed to the smallest sample size (only one sample out of 114, 0.9%). Also, tendon is a 177 relatively hypocellular tissue, which has a low protein turnover rate. Dataset PXD000867, 178 containing mouse liver samples, had the highest number of canonical proteins detected

(9,715, 77.3%), while the smallest number of proteins was detected in dataset PXD004612
(tendon, 273, 2.2%), as highlighted above (Fig. 2C).

181

182 We studied the normalised protein abundance distribution in organs (Fig. 2B) and found that 183 all organs, except tendon, had similar median abundances. However, one cannot attribute 184 further biological meaning to these observations, since by definition the method of 185 normalisation fixes each sample to have the same "total abundance", which then gets shared 186 out amongst all proteins. The normalised protein abundance distribution in datasets indicated 187 a higher than median abundances detected in datasets PXD004612 (tendon) and PXD003164 188 (testis) (Fig. 2D). A linear relationship was observed between the number of canonical 189 proteins detected in datasets and organs, when compared to the relative amount of their 190 spectral data (Fig. 2E). We found a significant number of proteins uniquely detected in one 191 organ (Fig. 2F). However, the list of concrete canonical proteins that were detected in just 192 one organ should be taken with caution since the list is subjected to inflated False Discovery 193 Rate (FDR), due to the accumulation of false positives when analysing the datasets 194 separately. 195 Some of the organs (liver, heart and brain) were represented across multiple mouse studies in 196 the aggregated dataset. A pairwise comparison of protein abundances in these organs generally showed a good correlation in expression (heart: R² values ranged from 0.54 to 0.83; 197 brain: R² from 0.28 to 0.72; and liver: R² from 0.59 to 0.74) (Figure S1 in Supplementary File 198 199 4).

200



202

Figure 2. (A) Number of canonical proteins identified across different mouse organs. The 203 204 number within the parenthesis indicates the number of samples. (B) Range of normalised 205 iBAQ protein abundances across different organs. The number within the parenthesis 206 indicates the number of samples. (C) Canonical proteins identified across different datasets. 207 The number within the parenthesis indicate the number of unique tissues in the dataset. (D) 208 Range of normalised iBAQ protein abundances across different datasets. The number within 209 parenthesis indicate the number of unique tissues in the dataset. (E) Comparison of total 210 spectral data with the number of canonical proteins identified in each dataset and organ. (F) 211 Distribution of canonical proteins identified across organs.

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213 **2.2.2. Rat proteome**

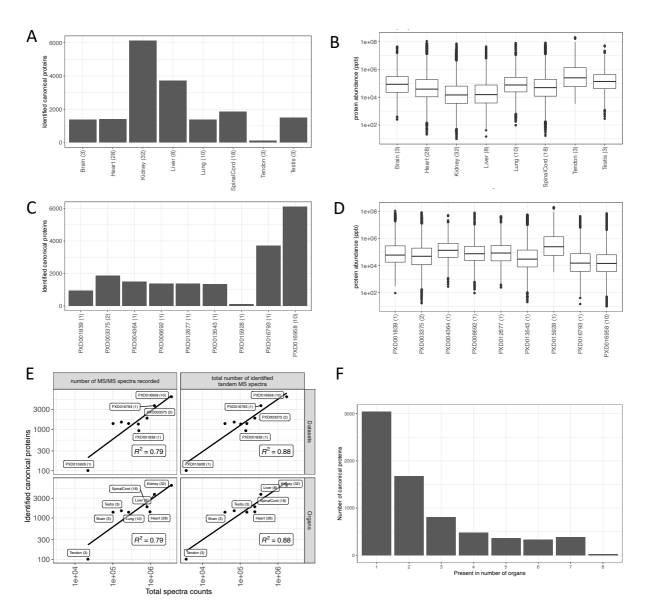
214 A total of 7,769 protein groups were identified across 8 different rat organs among which 215 3.649 (46.9%) protein groups were unique to one specific organ while 13 (0.16%) protein 216 groups were present among all organs (see full list in Supplementary File 2). The protein 217 groups were mapped to 7,116 genes (canonical proteins) (Supplementary File 3). The highest 218 number of canonical proteins (6,106, 85.1%) was found in rat kidney samples. The lowest 219 number of canonical proteins (101, 1.4%) was found in samples from tendon, as shown in 220 Fig. 3A. The largest number of canonical proteins identified in kidney is likely because of the 221 relatively large number of samples (32 samples), when compared to other organs. However, it 222 is interesting to note that large numbers of canonical proteins were detected in liver samples, 223 which relatively had fewer number of samples, when compared to the total number of 224 samples in heart and spinal cord.

225

226 Datasets PXD016958 and PXD016793 consisted entirely of kidney (where fractionation was 227 performed) and liver (no fractionation) samples, respectively, and as mentioned above had 228 the largest number of canonical proteins identified (Fig. 3C). The normalised protein 229 abundances were similar among the various organs and datasets (Fig. 3B, D). We also 230 observed a linear relation between the number of canonical proteins identified and the MS 231 spectra identified (Fig. 3E). As seen in the mouse datasets, we also observed a large number 232 of proteins uniquely detected in one organ (Fig. 3F). As highlighted above, the list of 233 concrete canonical proteins that were detected in just one organ should be taken with caution 234 since the list is subjected to inflated False Discovery Rate (FDR). 235 In the case of rat datasets, left ventricle heart samples were the only ones represented in more 236 than one study (PXD001839 and PXD013543) in the aggregated dataset. A pairwise

- 237 comparison of protein abundances of heart between these two datasets was performed,
- showing a strong correlation in protein expression ($R^2 = 0.9$) (Figure S1D in Supplementary
- 239 File 4).

240



241

Figure 3. (A) Number of canonical proteins identified across different rat organs. The
number within the parenthesis indicates the number of samples. (B) Range of normalised
iBAQ protein abundances across different organs. The number within the parenthesis
indicates the number of samples. (C) Canonical proteins identified across different datasets.
The number within the parenthesis indicate the number of unique tissues in the dataset. (D)

Range of normalised iBAQ protein abundances across different datasets. The number within
parenthesis indicate the number of unique tissues in the dataset. (E) Comparison of total
spectral data with the number of canonical proteins identified in each dataset and organ. (F)
Distribution of canonical proteins identified across organs.

251

252 **2.3.** Protein abundance comparison across organs

Next, we studied how protein abundances compared across different datasets and organs. The presence of batch effects between datasets makes this type of comparisons challenging. To aid comparison of protein abundances between datasets we transformed the normalised iBAQ intensities into ranked bins as explained in 'Methods', i.e., proteins included in bin 5 are highly abundant whereas proteins in bin 1 are expressed in the lowest abundances (among the detected proteins).

259

260 **2.3.1. Mouse proteome**

We found that 1,086 (8.6%) proteins were found with their highest level of expression in at least 3 organs, with a median bin value greater than 4 (Supplementary File 3). On the other end of the scale, 138 (1.1%) canonical proteins were found with their lowest expression in at least 3 organs, with a median bin value of less than 2. The bin transformed abundances in all organs are provided in Supplementary File 3.

266

267 To compare protein expression across all organs, we calculated pairwise Pearson correlation

268 coefficients across 117 samples (Fig. 4A). We observed some correlation in protein

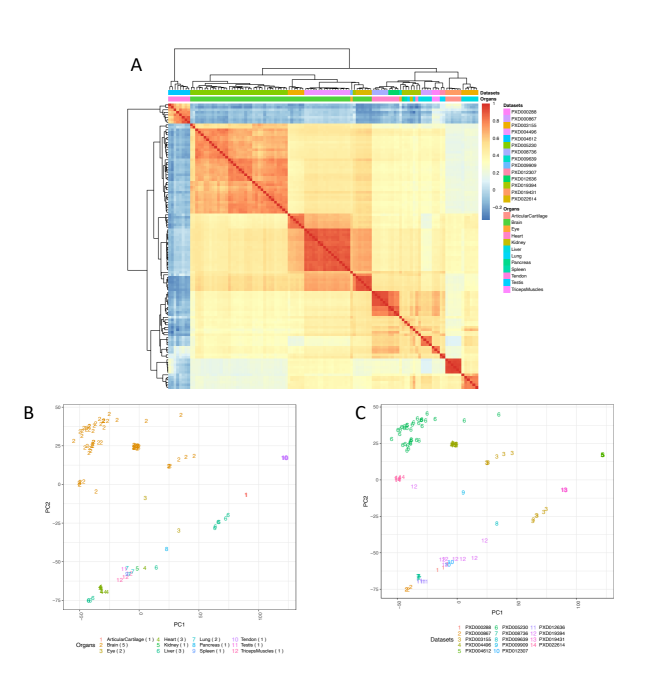
269 expression within brain (median $R^2 = 0.31$) and a higher one in heart (median $R^2 = 0.67$)

270 samples. We performed Principal Component Analysis (PCA) on all samples from mouse

271 datasets for testing the effectiveness of the bin transformation method in reducing batch

effects. Fig. 4B shows the clustering of samples from various organs of mouse. We observed
samples from the same organ generally clustered together. For example, we observed that
brain samples all clustered together in one group, even though they come from different
datasets, indicating decent removal of batch effects (Fig. 4C). However, we also observed
that samples from other organs such as liver did not cluster according to their organ types but
clustered together within the dataset they were part of, indicating some residual batch effects,
which are hard to remove completely.





281 Figure 4. (A) Heatmap of pairwise Pearson correlation coefficients across all mouse samples. 282 The colour represents the correlation coefficient and was calculated using the bin transformed 283 iBAQ values. The samples were hierarchically clustered on columns and rows using 284 Euclidean distances. (B) PCA of all samples, using the binned protein abundances as input, 285 coloured by the organ types. (C) PCA of all samples coloured by their respective dataset 286 identifiers. The numbers in parenthesis indicate the number of datasets for each organ. 287 Binned values of canonical proteins quantified in at least 50% of the samples were used to 288 perform the PCA.

289

290 In addition, we compared the protein abundances generated in this study with the data 291 available in the resource PaxDB generated using spectral counting across different mouse 292 organs. We observed generally a strong correlation of protein abundances calculated using 293 iBAQ from this study (fraction of total (FOT) normalised ppb) and spectral counting methods 294 (Figure S2 in Supplementary File 4). However, the expression of low abundant proteins 295 seemed to be underestimated in PaxDB when compared with our results, as shown by a S-296 shaped curve in the scatterplot in organs such as brain, heart, liver and lung. The 'dynamic 297 exclusion' [41] setting used by modern mass spectrometers prevents the instrument from 298 fragmenting abundant peptides multiple times when they are repeatedly observed in scans 299 nearby in time. This has the effect that spectral counting approaches will limit the dynamic 300 range observed, as high abundant proteins will be under sampled. This is a limitation when 301 using spectral counting methods, and these days spectral counting is not commonly used as a 302 truly quantitative data type in proteomics.

303

304 2.3.2. Rat proteome

305 Next, we studied the distribution of protein abundances across organs in rat. On one hand, 306 311 (4.3%) proteins were found with their highest expression in at least 3 organs with a 307 median bin value greater than 4. On the other hand, 27 (0.37%) canonical proteins were 308 found with their lowest expression in at least 3 organs, with a median bin value of less than 2. 309 The bin transformed abundances in all organs are provided in Supplementary File 3. 310 Overall, the samples from rat datasets showed a better correlation in protein expression (Fig. 311 5A) than in the case of mouse. We observed generally a strong correlation of protein expression within samples from liver (median Pearson's correlation $R^2 = 0.85$), lung (median 312 $R^2 = 0.71$), spinal cord (median $R^2 = 0.65$), heart (median $R^2 = 0.71$) and brain (median $R^2 = 0.71$) 313 314 0.86). We also observed the clustering in the PCA of samples coming from the same organ 315 (Fig. 5B). Kidney, lung, spinal cord and heart samples all clustered together according to 316 their organ type. Fig. 5C shows the samples based on the dataset they were part of. However, 317 most organ samples were part of individual datasets except in the case of samples from heart, 318 which came from two datasets (PXD001839 and PXD013543). Fig. 5C shows that the heart 319 samples clustered into two nearby groups (bottom left two clusters on Fig. 5B and 5C), 320 wherein each cluster included samples from a different dataset, indicating the presence of 321 small batch effects.

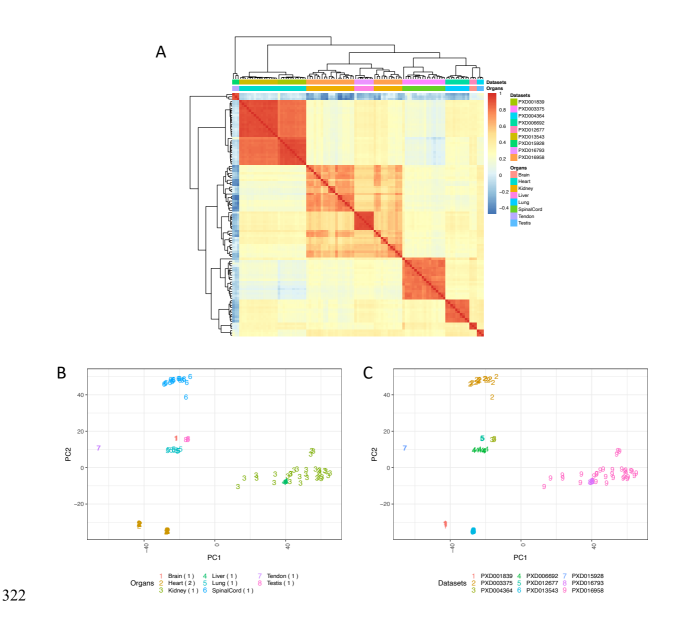


Figure 5. (A) Heatmap of pairwise Pearson correlation coefficients across all rat samples.
The colour represents the correlation coefficient and was calculated using the bin transformed
iBAQ values. The samples were hierarchically clustered on columns and rows using
Euclidean distances. (B) PCA of all samples coloured by the organ types. (C) PCA of all
samples coloured by their respective dataset identifiers. The numbers in parenthesis indicate
the number of datasets for each organ. Binned values of canonical proteins quantified in at
least 50% of the samples were used to perform the PCA.

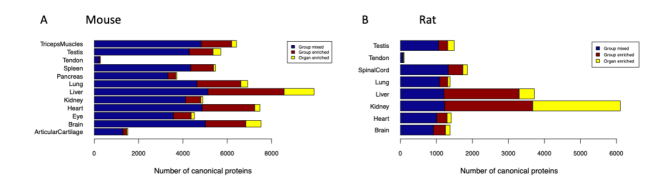
330

2.4. The organ elevated proteome and the over-representative biological processes

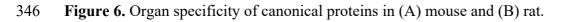
332	Based on their expression, canonical proteins were classified into three different groups based
333	on their organ specificity: "mixed", "group-enriched" and "organ-enriched" (see
334	Supplementary File 5). We considered over-expressed canonical proteins in each organ as
335	those which were in "group-enriched" and "organ-enriched". The analysis showed that on
336	average, 20.8% and 26.0% of the total elevated canonical proteins were organ group-specific
337	in mouse and rat, respectively (Fig. 6). In addition, 4.3% and 14.2% were unique organ-
338	enriched in mouse and rat, respectively. The highest ratio of organ-enriched in mouse was
339	found in liver (13.6%), whereas in rat, it was found in kidney (39.8%).

- 340 We then performed a gene ontology (GO) enrichment analysis of those proteins that were
- 341 'organ-enriched' and group-enriched' using GO terms associated with biological processes.
- 342 We found 1,036 GO terms to be statistically significant in all organs, as seen in
- 343 Supplementary File 6. The most significant GO terms for each organ are shown in Table 2.









347

Organ	Species	GO ID	Description	adjusted p- value
Articular cartilage	$ M_{us} _{musculus} (+()\cdot)043062$		Extracellular matrix organization Extracellular structure organization External encapsulating structure organization	8.94*10 ⁻³⁸ 8.94*10 ⁻³⁸ 8.94*10 ⁻³⁸
Brain	rain Mus musculus GO:0050804 GO:0099177 GO:0050808		Modulation of chemical synaptic transmission Regulation of trans-synaptic signalling Synapse organization	7.03*10 ⁻⁶⁵ 7.03*10 ⁻⁶⁵ 1.41*10 ⁻⁴⁸

Heart	t Mus musculus		Heart contraction Regulation of heart contraction Muscle tissue development	$\begin{array}{c c} 7.10^{*10^{-11}} \\ 4.43^{*10^{-10}} \\ 6.16^{*10^{-10}} \end{array}$
Kidney	Mus musculus	GO:0015711 GO:0044282 GO:0016054	Organic anion transport Small molecule catabolic process Organic acid catabolic process	4.59*10 ⁻¹⁹ 4.91*10 ⁻¹⁵ 6.25*10 ⁻¹⁵
Eye	Mus musculus	GO:0007601 GO:0001654 GO:0099504	Visual perception Eye development Synaptic vesicle cycle	7.54*10 ⁻⁵⁰ 5.31*10 ⁻³¹ 8.36*10 ⁻¹⁸
Liver	Mus musculus	GO:0016569 GO:0016570 GO:0019369	Covalent chromatin modification Histone modification Arachidonic acid metabolic process	6.26*10 ⁻¹⁰ 1.71*10 ⁻⁰⁸ 1.71*10 ⁻⁰⁸
Lung	Mus musculus	GO:0120031 GO:0030031 GO:0044782	Plasma membrane bounded cell projection assembly Cell projection assembly Cilium organization	3.61*10 ⁻¹⁴ 3.61*10 ⁻¹⁴ 9.83*10 ⁻¹⁴
Pancreas	Mus musculus	GO:0007586 GO:0032328	Digestion Alanine transport	0.005 0.018
Spleen	Mus musculus	GO:0046649 GO:0050776 GO:0045087	Lymphocyte activation Regulation of immune response Innate immune response	4.12*10 ⁻²² 2.00*10 ⁻²⁰ 2.23*10 ⁻²⁰
Tendon	ndon Mus musculus GO:00 GO:00 GO:00		Muscle system process Multicellular organismal movement Musculoskeletal movement	$\begin{array}{c} 1.46^{*}10^{-25}\\ 3.14^{*}10^{-19}\\ 1.46^{*}10^{-25}\end{array}$
Testis	Testis Mus musculus G		Male gamete generation Cilium movement Cilium organization	8.75*10 ⁻⁴⁹ 3.04*10 ⁻³⁸ 6.78*10 ⁻³⁷
Triceps muscles	Mus musculus	GO:0061061 GO:0055002 GO:0003009	Muscle structure development Striated muscle cell development Skeletal muscle contraction	$\begin{array}{c} 1.56^{*}10^{-14}\\ 2.41^{*}10^{-14}\\ 3.53^{*}10^{-14}\end{array}$
Brain	Rattus norvegicus	GO:0099537 GO:0007268 GO:0098916	Trans-synaptic signalling Chemical synaptic transmission Anterograde trans-synaptic signalling	1.79*10 ⁻⁶⁰ 1.79*10 ⁻⁶⁰ 1.79*10 ⁻⁶⁰
Heart	Rattus norvegicus	GO:0061061 GO:0003012 GO:0055001	Muscle structure development Muscle system process Muscle cell development	$\begin{array}{c} 2.94^{*}10^{-17} \\ 6.30^{*}10^{-16} \\ 4.00^{*}10^{-15} \end{array}$
Kidney	Rattus norvegicus	GO:0006396 GO:0045944 GO:0006260	RNA processing positive regulation of transcription by RNA polymerase II DNA replication	6.19*10 ⁻¹³ 7.29*10 ⁻⁰⁶ 1.74*10 ⁻⁰⁵
Liver	Rattus norvegicus	GO:0008202 GO:0016054 GO:0032787	Steroid metabolic process Organic acid catabolic process Monocarboxylic acid metabolic process	$2.74^{*}10^{-10} \\ 1.61^{*}10^{-09} \\ 1.64^{*}10^{-09}$
Lung	Rattus norvegicus GO:0031589 GO:0009617 GO:0030036		Cell-substrate adhesion Response to bacterium Actin cytoskeleton organization	7.62*10 ⁻⁰⁸ 7.62*10 ⁻⁰⁸ 1.40*10 ⁻⁰⁷
Spinal cord	pinal cord Rattus norvegicus GO:0061564 GO:0099537 GO:0007268		Axon development Trans-synaptic signalling Chemical synaptic transmission	4.26*10 ⁻¹⁸ 5.93*10 ⁻¹⁶ 5.93*10 ⁻¹⁶
Tendon	Rattus norvegicus	GO:0030199 GO:0061448 GO:0001501	Collagen fibril organization Connective tissue development Skeletal system development	1.23*10 ⁻¹³ 2.31*10 ⁻⁰⁹ 3.39*10 ⁻⁰⁹

Testis	Rattus norvegicus	GO:0019953 GO:0051704 GO:0007018	Sexual reproduction Multi-organism process Microtubule-based movement	3.98*10 ⁻²⁴ 1.61*10 ⁻¹⁸ 4.00*10 ⁻¹²
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348

349 Table 2. Analysis of the top three GO terms for each organ in mouse and rat using the 350 elevated organ-specific and group-specific canonical proteins as described in the 'Methods' 351 section. 352 353 2.5. Protein abundances across orthologs in three species In a previous study, we analysed 25 label-free proteomics datasets from healthy human 354 355 samples to assess baseline protein abundances in 14 organs following the same analytical 356 methodology [14]. We compared the expression of canonical proteins identified in all three 357 species (rat, mouse and human). Overall, 13,248 detected human genes (corresponding to the 358 canonical proteins) were compared with 12,570 genes detected in mouse and 7,116 genes 359 detected in rat. The number of orthologous mappings (i.e., "one-to-one" mappings, see 360 'Methods') between rat, mouse and human genes are listed in table 3. We only considered 361 one-to-one mapped orthologues for the comparison of protein abundances. 362

		Orthologs of human genes identified	ith different mapping against identified human genes				
Species	Identified genes	in [14]	one-to-one	one-to- many	many-to- many	many-to- one	not mapped
Mus musculus	12,570	10,601	80.4%	1.9%	0.56%	1.46%	15.7%
Rattus norvegicus	7,116	6,058	82.0%	2.2%	0.70%	0.25%	14.9%

363

364 **Table 3.** Homologs identified in mouse and rat datasets when compared with the background

365 list of genes (corresponding to canonical proteins) identified in human datasets

366 (Supplementary File 2 in [14]).

367

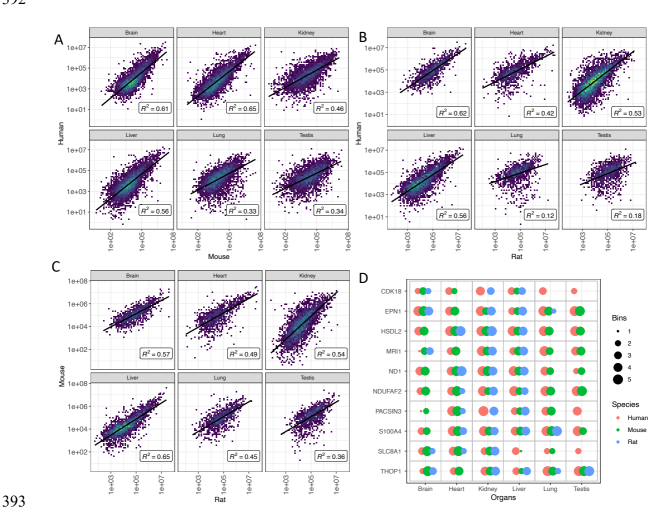
368 Among human and mouse orthologues we observed relatively high levels of correlation of protein abundances in brain ($R^2 = 0.61$), heart ($R^2 = 0.65$) and liver ($R^2 = 0.56$) (Fig. 7A). 369 Human and rat orthologs showed also relatively high levels of correlation in brain ($R^2 =$ 370 0.62), kidney ($R^2 = 0.53$) and liver ($R^2 = 0.56$), but almost no correlation in lung ($R^2 = 0.12$) 371 and testis ($R^2 = 0.18$) (Fig. 7B). Between mouse and rat orthologs, the correlation of protein 372 abundances was higher in liver ($R^2 = 0.65$), kidney ($R^2 = 0.54$) and brain ($R^2 = 0.57$) samples, 373 when compared to the samples coming from the rest of the organs (Fig. 7C). Fig. 7D shows 374 375 an illustration of some example comparisons of individual orthologs using binned protein 376 abundances. 377

For the same corresponding subsets, we also investigated the correlation of protein
expression between various organs within each organism. We observed that in general the
correlation of protein expression was slightly lower between organs within the same species,
when compared to a higher correlation, which was observed among orthologs (Figure S3 in
Supplementary File 4). The found lower correlation of protein expression between different
organs was more apparent in mouse and rat.

384

Among the orthologs expressed in all organs in all three species, 747 (12.3%) orthologs were detected with a median bin expression value of more than 4, i.e., proteins that appear to have conserved high expression in all organs and all tissues. Additionally, 13 (0.2%) orthologs were found with a median bin expression value less than 2 in all organs, although, it is harder to detect consistently proteins with low abundances across all organs. A full list of the binned protein abundances of orthologs is available in Supplementary File 7. The illustration of all binned protein abundances across the three species is shown in Supplementary File 8.

392



394 Figure 7. Comparison of protein abundances (in ppb) between one-to-one mapped orthologs 395 of mouse, rat and human in various organs. (A) Pairwise correlation using normalised protein 396 abundances of human and mouse orthologues. (B) Human and rat orthologs. (C) Mouse and 397 rat orthologs. (D) As an example, the comparisons of binned protein expression of ten 398 randomly sampled orthologs are shown. Data corresponding to all cases (as reported in panel 399 D) are available in Supplementary File 7 and the corresponding illustration of binned values 400 is available in Supplementary File 8. Orthologs in (D) are shown using their human gene 401 symbol.

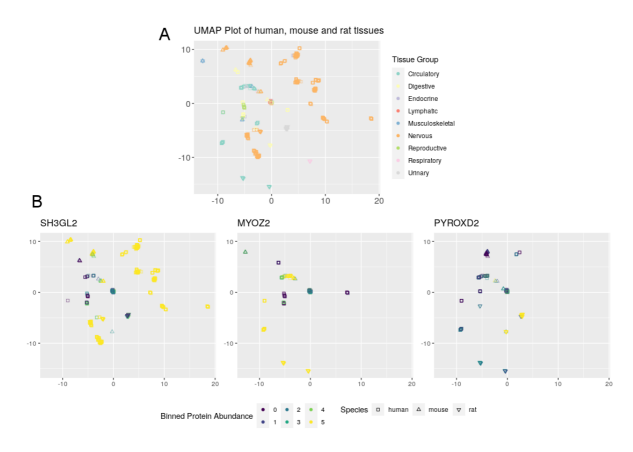
402 Since each sample contains potentially thousands of protein values this creates a high level of403 dimensionality within the data. To reduce this, we used the non-linear dimension reduction

404 algorithm, Uniform Manifold Approximation and Projection (UMAP) (see Section 4.7 in the 405 'Methods' section). The UMAP algorithm enables the reduction of multidimensional data to 406 a two-dimensional space upon which the relationship between each sample can be visualised. 407 Specifically, it enables the visualisation of the relationships of proteins across individual 408 samples and organs. Should multiple samples be positioned near to each other, it allows for 409 us to predict that these samples shared similar properties (in this case, similar protein 410 abundance values). Consequently, by overlaying samples from various species UMAP 411 representations can be used to visualise the relationship of various orthologs across similar 412 organs.

Using the UMAP algorithm, we were able to visualise the relationships between individual 413 414 organs regardless of the involved species (human, mouse, rat) and to identify similar genes 415 (corresponding to canonical proteins) within those organs. The overall view of all samples 416 labelled by their respective organ is shown as Figure 8A. We chose to use the biological 417 system as the basis for the colouring scheme for each sample to reduce the overall complexity 418 of the visualisation, due the high number of organs included. By using this labelling scheme, 419 we could see that the clustering of each sample was deterministic. Each sample was 420 positioned within a clear region for the corresponding organs, despite the original layout being unaware of this information. This indicates that not only do the samples within those 421 422 organs share common protein abundance values, but furthermore, that samples that come 423 from the same organs share similar protein expression (as three species are present).

Furthermore, in Figure 8B we show the representation of binned protein abundance values for
three example genes (SH3GL2, MYOZ2 and PYROXD2), providing information on the
abundance of them across different biological systems. These visualisations use the same
layout than within Figure 8A. In the example of SH3GL2, it can be seen that Figure 8B

428 shows multiple values that have been scored as bin 5. By referring to Figure 8A, we can see 429 that those points corresponding to highly abundant proteins, come from samples from the 430 nervous system (in all three species). Furthermore, using the same method, it can be seen that 431 MYOZ2 is highly abundant in the circulatory system, and that PYROXD2 is highly abundant 432 in the urinary system. The UMAP coordinates and our binned protein abundance data that is 433 used in these plots to allow for the generation of similar visualisations are provided in 434 Supplementary File 9.





436 **Figure 8:** Visualisations generated using the UMAP algorithm to show the relationships

437 between human, mouse, and rat samples. (A) Shows the relationship of all samples,

438 particularly showing strong relationship between biological systems. (B) Shows the protein

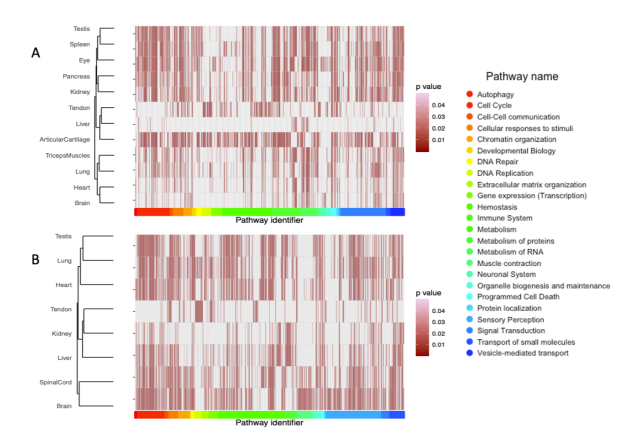
439 abundancy of 3 example gene orthologs (SH3GL2, MYOZ2 and PYROXD2), within each

sample. Human baseline protein expression data was generated in [14].

442 **2.6. Pathway enrichment analysis**

443 Based on the ortholog protein expression analysis described above, we mapped canonical proteins from mouse and rat to the corresponding ortholog human proteins, which were 444 445 subsequently subjected to pathway-enrichment analysis using Reactome (Fig. 9). After 446 filtering out the disease and statistically insignificant pathways, there were 2,990 pathways 447 found in all the organs of mouse and 2,162 pathways in all the organs of rat. In mouse 448 samples, the largest number of pathways (367) were found in articular cartilage, and the 449 lowest number of pathways was found in liver (44). We also observed that Neuronal System-450 related pathways were predominantly present in the brain and eye, which is consistent with expectations. In rat samples, brain included the largest number of pathways (387), while the 451 452 lowest number of pathways was found in tendon, with 117.





- 455 **Figure 9.** Pathway analysis performed using the canonical proteins, showing the statistically
- 456 significant representative pathways (p-value < 0.05) in (A) mouse and (B) rat organs.

457 **3. Discussion**

458 We have previously reported two meta-analysis studies involving the reanalysis and 459 integration in Expression Atlas of public quantitative datasets coming from cell lines and 460 human tumour samples [13], and from human baseline tissues [14], respectively. In this 461 study, we reanalysed mouse and rat baseline proteomics datasets representing protein 462 expression across 34 healthy tissues and 14 organs. We have used the same methodology as 463 in the study involving baseline human tissues, which enabled a comparison of protein 464 expression levels across the three species. Our main overall aim was to provide a system-465 wide baseline protein expression catalogue across various tissues and organs of mouse and rat 466 and to offer a reference for future related studies. 467 468 We analysed each dataset separately using the same software (MaxQuant) and the same 469 search protein sequence database. The disadvantage of this approach is that the FDR 470 statistical thresholds are applied at a dataset level and not to all datasets together as a whole. 471 However, as reported before [14], using a dataset per dataset analysis approach is in our view 472 the only sustainable manner to reanalyse and integrate quantitative proteomics datasets, at

473 least at present. The disadvantage of this approach is that the FDR statistical threshold are 474 applied at a dataset level and not to all datasets together as a whole, with the potential 475 accumulation of false positives across datasets. However, it is important to highlight that the 476 number of commonly detected false positives is reduced in parallel with the increase in the 477 number of common datasets where a given protein is detected. As also reported in previous 478 studies, one of the major bottlenecks was the curation of dataset metadata, consisting of 479 mapping files to samples and biological conditions. Very recently, the MAGE-TAB-480 Proteomics format has been developed and formalised to enable the reporting of the 481 experimental design in proteomics experience, including the relationship between samples

and raw files, which is recorded in the SDRF-Proteomics section of the file [42]. Submission
of the SDRF-Proteomics files to PRIDE is now supported. The more well-annotated datasets
in the public domain, the easier these data reuse activities will become.

485

486 The generated baseline protein expression data can be used with different purposes such as the generation of protein co-expression networks and/or the inference of protein complexes. 487 488 For the latter application, expression data can be alone or for potentially refining predictions 489 obtained using different methods such as the recently developed AlphaFold-based protein 490 complexes predictions [43]. Mouse and rat are widely used species in the context of drug 491 discovery, the latter especially, to undertake regulatory pre-clinical safety studies. Therefore, 492 it is important to know quantitative protein expression distribution in these species in 493 different tissues [44] to assist in the selection of species for such studies and also for the 494 interpretation of the final results.

495

496 In addition to the analyses reported, it would have also been possible to perform correlation 497 studies between gene and protein expression information. However, we did not find any 498 relevant public datasets in the context of this manuscript where the same samples were 499 analysed by both techniques, which is the optimal way to perform these studies. Future 500 directions in analogous studies will involve: (i) additional baseline protein expression studies 501 of other species, including other model organisms or other species of economic importance; 502 (ii) the inclusion of differential proteomics datasets (e.g. using TMT and/or iTRAO); and (iii) 503 include relevant proteomics expression data coming from the reanalysis of Data Independent 504 Acquisition (DIA) datasets [45].

505

506 As mentioned above, we performed a comparative analysis of baseline protein expression 507 across human, mouse and rat. It was possible to perform this analysis for six common organs 508 (brain, heart, kidney, liver, lung and testis). Ortholog expression across species is useful to 509 infer protein function across experimentally studied proteins. This is particularly useful as 510 evolutionarily closely related species are likely to conserve protein function. We could not 511 find in the literature an analogous comparative study performed at the protein level. 512 However, expression from closely related orthologs across tissues or organs has been 513 compared at the transcriptomics level, providing a complete picture of gene expression. In 514 this context, many studies have compared gene-expression in mouse, rat and human 515 orthologues and found that orthologues had generally a highly correlated expression tissue 516 distribution profile in baseline conditions [46-50]. Gene expression levels among orthologs 517 were found to be highly similar in muscle and heart tissues, liver and nervous system and less 518 similar in epithelial cells, reproductive systems, bone and endocrine organs [48]. Studies have 519 also shown that variability of gene expression between homologous tissues/organs in closely 520 related species can be lower than the variability between unrelated tissues within the same 521 organism [46, 47], in agreement with the results reported here at the protein level. 522 Additionally, we showed an initial analysis of protein expression of orthologs across the three 523 species using UMAP.

524

525 In conclusion we here present a meta-analysis study of public mouse and rat baseline 526 proteomics datasets from PRIDE. We demonstrate its feasibility, perform a comparative 527 analysis across the three species and show the main current challenges. Finally, the data is 528 made available *via* Expression Atlas. Whereas there are several analogous studies performed 529 at the gene expression level for mouse and rat tissues, to the best of our knowledge this is the 530 first of this kind at protein expression level.

531 4. Materials and Methods

532

533 **4.1. Datasets**

- As of May 2021, there were 2,060 mouse (*Mus musculus*) and 339 rat (*Rattus norvegicus*)
- 535 MS proteomics datasets publicly available in the PRIDE database
- 536 (<u>https://www.ebi.ac.uk/pride/</u>). Datasets were manually selected based on the selection
- 537 criteria described previously [14]. Briefly, we selected datasets where baseline expression
- 538 experiments were performed on (i) label-free samples from tissues not enriched for post-
- 539 translational modifications; (ii) Thermo Fisher Scientific instruments such as LTQ Orbitrap,
- 540 LTQ Orbitrap Elite, LTQ Orbitrap Velos, LTQ Orbitrap XL ETD, LTQ-Orbitrap XL ETD,
- 541 Orbitrap Fusion and Q-Exactive, since they represent a large proportion of datasets in PRIDE
- and to avoid heterogeneity introduced by data from other vendor instruments; (iii) had
- 543 suitable sample metadata available in the original publication or it was possible to obtain it by
- 544 contacting the authors; and (iv) our previous experience in the team of some datasets
- 545 deposited in PRIDE, which were discarded because they were not considered to be useful.
- 546 Overall, 14 mouse and 9 rat datasets were selected from all mouse and rat datasets for further
- analysis. Table 1 lists the selected datasets. The 23 datasets contained a total of 211 samples
- 548 from 34 different tissues across 14 organs (meaning groups of related tissues, more details
- 549 below), comprising 9 different mouse and 3 rat strains, respectively.
- 550 The sample and experimental metadata were manually curated using the information
- 551 provided in the respective publications or by contacting the original authors/submitters.
- 552 Annotare [51] was used for annotating the metadata and stored using the Investigation
- 553 Description Format (IDF) and Sample-Data Relationship Format (SDRF) file formats [42],
- s54 which are required for integration in Expression Atlas. An overview of the experimental
- 555 design including experimental factors, protocols, publication information and contact

information are present in the IDF file, and the SDRF includes sample metadata describing
the relationship between the various sample characteristics and the data files contained in the
dataset.

559

560 **4.2. Proteomics raw data processing**

561 All datasets were analysed with MaxQuant (version 1.6.3.4) [52, 53] on a Linux high-

562 performance computing cluster for peptide/protein identification and protein quantification.

563 Input parameters for each dataset, such as MS¹ and MS² tolerances, digesting enzymes, fixed

and variable modifications, were set as described in their respective publications, with two

565 missed cleavage sites. The FDR at the PSM (peptide spectrum match) and protein levels were

set to 1%. The MaxQuant parameters were otherwise set to default values: the maximum

number of modifications per peptide was 5, the minimum peptide length was 7, the maximum

568 peptide mass was set to 4,600 Da, and for the matches between runs the minimum match time

569 window was set to 0.7 seconds and the minimum retention time alignment window was set to

570 20 seconds. The MaxQuant parameter files are available for downloading from Expression

571 Atlas. The *Mus musculus* UniProt Reference proteome release-2021_04 (including isoforms,

572 63,656 sequences) and *Rattus norvegicus* UniProt Reference proteome release-2021_04

573 (including isoforms, 31,562 sequences) were used as the target sequence databases for mouse

and rat datasets, respectively. The built-in contaminant database within MaxQuant was used

and a decoy database was generated by MaxQuant by reversing the input database sequences

576 after the respective enzymatic digestion. The datasets were run separately in multi-threaded

577 mode.

578

579 **4.3. Post-processing**

The post-processing of results from MaxQuant is explained in detail in [14]. In brief, the protein groups labelled as potential contaminants, decoys and those with fewer than 2 PSMs were removed. Protein intensities in each sample were normalised by scaling the iBAQ intensity values to the total amount of signal in each MS run and converted to parts per billion (ppb).

$$ppb_{i}BAQ_{i} = \left(\frac{iBAQ_{i}}{\sum_{i=1}^{n} iBAQ_{i}}\right) x 1,000,000,000$$

586 The 'majority protein identifiers' within each protein group were mapped to their Ensembl 587 gene identifiers/annotations using the Bioconductor package 'mygene'. For downstream 588 analysis only protein groups whose isoforms mapped to a single unique Ensembl gene ID 589 were considered. Protein groups that mapped to more than one Ensembl gene ID are provided 590 in Supplementary File 1. The protein intensity values from different protein groups with the 591 same Ensembl gene ID were aggregated as median values. The parent genes to which the 592 different protein groups were mapped to are equivalent to 'canonical proteins' in UniProt 593 (https://www.uniprot.org/help/canonical_and_isoforms) and therefore the term protein 594 abundance is used to describe the protein abundance of the canonical protein throughout the 595 manuscript.

596

597 4.4. Integration into Expression Atlas

598 The calculated canonical protein abundances (mapped to genes), together with the validated599 SDRF files, summary files detailing the quality of post-processing and the input MaxQuant

- 600 parameter files (mqpar.xml) were integrated into Expression Atlas
- 601 (<u>https://www.ebi.ac.uk/gxa/home</u>) as proteomics baseline experiments (E-PROT identifiers
- 602 are available in Table 1).
- 603

604 **4.5. Protein abundance comparison across datasets**

605 To compare protein abundances, the normalised protein abundances (in ppb) from each group 606 of tissues in a dataset were converted into ranked bins. In this study, 'tissue' is defined as a 607 distinct functional or structural region within an 'organ'. For example, hippocampus, 608 cerebellum and cortex are defined as 'tissues' that are part of the brain (organ) and similarly 609 sinus node, left atria, left ventricle, right atria, right ventricle are defined as 'tissues' in heart 610 (organ). Protein abundances were transformed into bins by first grouping MS runs from each 611 tissue within a dataset as a batch. The normalised protein abundances (ppb) for each MS run 612 within a batch were sorted from lowest to highest abundance and ranked into 5 bins. Proteins 613 whose ppb abundances are ranked in the lowest bin (bin 1) represent lowest abundance and 614 correspondingly proteins within bin 5 are of highest abundance in their respective tissue.

615 When merging tissues into organs, median bin values were used.

616 Proteins that were detected in at least 50% of the samples were selected for PCA (Principal 617 Component Analysis) and was performed using R (The R Stats package) [54] using binned 618 abundance values. For generating heatmaps, a Pearson correlation coefficient for all samples 619 was calculated on pairwise complete observations of bin transformed values. Missing values were marked as NA (not available). For each organ a median R^2 was calculated from all 620 621 pairwise R² values of their respective samples. Samples were hierarchically clustered on 622 columns and rows using Euclidean distances. To compare the correlation in protein 623 expression of shared organs between datasets, the FOT normalised protein abundances (ppb) 624 were aggregated by calculating the median over samples. The regression line was computed 625 using the 'linear model' (lm) method in R.

4.6 Comparison of protein abundances using iBAQ and spectral counting data availablein PaxDB

628 To compare protein abundances generated from iBAQ in this study and spectral counting 629 methods, protein abundance data from different mouse organs was obtained from PaxDB 630 (https://www.pax-db.org/) [16]. FOT normalised iBAQ abundances, as described above, were 631 compared with the spectral counting abundances for the matching mouse organs. Organs 632 from mouse labelled as 'integrated' in PaxDB were selected. It was not possible to perform 633 this comparison for rat organs since data in PaxDB for rat are available for either the 'whole 634 organism' or for "cell types" only. Abundances were compared across mouse adipose tissue, 635 brain, heart, kidney, liver, lung, pancreas and spleen. The Ensembl ENSG gene ids were 636 mapped to ENSP protein ids in PaxDB using the 'mygene' bioconductor package in R.

637 4.7. UMAP analysis

638 To generate the UMAP visualisations we used the binned protein abundance values generated 639 in this study from rat and mouse, as well as the binned human protein abundance values from 640 [14]. First, we reduced this data to only contain the orthologs found in all three species. For 641 the purpose of only the initial visualisation layout, we filtered the data to include those 642 proteins present in 90% of samples. Once the initial layout was generated, we then used the 643 full protein abundance values to generate protein-specific visualisations. We use R v4.1.0 644 with the package 'umap' (Uniform Manifold Approximation and Projection in R) [55] 645 v0.2.7.0 to generate the UMAP visualisations.

646 **4.8. Organ-specific expression profile analysis**

647 For comparison across organs, the tissues were aggregated into organs and their median bin

values were considered. As described previously [14] the classification scheme done by

649 Uhlén *et al.* [17] was modified to classify the proteins into one of the three categories: (1)

650 "Organ-enriched": present in one unique organ with bin values 2-fold higher than the mean

bin value across all organs; (2) "Group enriched": present in at least 7 organs in mouse or in
at least 4 organs in rat, with bin values 2-fold higher than the mean bin value across all
organs; and (3) "Mixed": the remaining canonical proteins that are not part of the above two
categories.

655

Enriched gene ontology (GO) terms analysis was carried out through over-representation test
described previously [14], it was combined with "Organ-enriched" and "Group enriched"
mapped gene lists for each organ. In addition, Reactome [56] pathway analysis was
performed using mapped gene lists and running pathway-topology and over-representation
analysis, as reported previously [14].

661 **4.9. Comparison of protein expression across species**

The g:Orth Orthology search function in the g:Profiler suite of programs [57] was used for translating gene identifiers between organisms. Since a custom list of gene identifiers could not be used as the background search set, the mouse and rat genes were first mapped against the background Ensembl database. The resulting list of mouse and rat genes mapped to human orthologs were then filtered so that they only included parent gene identifiers of the protein groups from mouse and rat organs identified in this study and the parent genes of human organs described in our previous study (Supplementary File 2 in [14]), respectively.

The orthologs were grouped into various categories denoting the resulting mapping between identifiers: "one-to-one", "one-to-many", "many-to-one", "many-to-many", and "no mappings" between gene identifiers. Only "one-to-one" mapped ortholog identifiers were used to compare protein intensities between mouse, rat and human organs. The normalised ppb protein abundances of the one-to-one mapped orthologues in 6 organs (brain, heart,

675	kidney, liver, lung and testis), that were studied across all three organisms were used to assess
676	the pairwise correlation of protein abundances. The linear regression was calculated using the
677	linear fit 'lm' method in R.

678

Data availability 679

Expression Atlas E-PROT identifiers and PRIDE original dataset identifiers are included in 680 681 Table 1.

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- 683 First of all, we would like to thank all data submitters who made their datasets available via
- 684 PRIDE and ProteomeXchange. We would also like to thank Andrew Leach and the rest of the
- team involved in the Open Targets "Target Safety" project, for helpful discussions. 685

Authors' contributions 686

- 687 SW, DGS, AP, DJK selected and curated the datasets. SW, AP, DGS performed analyses. AC
- and ARJ helped in the interpretation of results and designed approach for data normalisation. 688
- 689 NG, SF, PM, and IP helped in the integration of the results into Expression Atlas. SW, AP,
- 690 DGS, JAV wrote the manuscript. JAV, ARJ and IP obtained the funding for performing the
- 691 study. All authors have read and approved the manuscript.
- 692

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- and analysis, decision to publish, or preparation of the manuscript.
- 701
- 702 Supporting information
- 703 Supplementary file 1: Protein groups from all datasets that are mapped to more than one
- Ensembl Gene ID.
- 705 Supplementary file 2: Median protein abundances (in ppb) for each protein group across
- various tissue samples in each organ.
- 707 Supplementary file 3: Median binned protein abundances across various tissue samples in

- 709 Supplementary file 4: Supplementary figures (S1) illustrating correlation of protein
- abundances in organs represented in different datasets. (S2) Correlation of protein
- abundances generated using iBAQ and spectral counting methods in various mouse organs.
- 712 (S3) Correlation of protein expression between organs within human, mouse and rat.
- 713 **Supplementary file 5:** Organ distribution of canonical proteins in mouse and rat.

714 Supplementary file 6: Gene Ontology enrichment analysis of 'organ-enriched' and 'group-

- 715 enriched' proteins.
- Supplementary file 7: Binned protein abundances of one-to-one mapped orthologs across all
 organs studied.
- 718 Supplementary file 8: Figure illustrating binned protein abundances of all one-to-one
- 719 mapped orthologs across six common organs in mouse, rat and human.
- 720 Supplementary file 9: UMAP co-ordinates and source data of UMAP analysis.

721

722 Abbreviations

- 723 DDA: Data Dependent Acquisition
- 724 DIA: Data Independent Acquisition
- 725 FOT: Fraction of Total
- 726 GO: Gene Ontology
- 727 iBAQ: intensity-based absolute quantification
- 728 iTRAQ: Isobaric tag for relative and absolute quantitation
- 729 IDF: Investigation Description Format
- 730 MS: Mass Spectrometry
- 731 ppb: Parts per billion
- 732 PCA: Principal Component Analysis
- 733 SDRF: Sample and Data Relationship Format
- 734 TMT: Tandem Mass Tagging
- 735 UMAP: Uniform Manifold Approximation and Projection

Aebersold R, Mann M. Mass-spectrometric exploration of proteome structure and

function. Nature. 2016;537(7620):347-55. Epub 2016/09/16. doi: 10.1038/nature19949.

736 **References**

PubMed PMID: 27629641.

1.

737 738

739

740

741 Perez-Riverol Y, Csordas A, Bai J, Bernal-Llinares M, Hewapathirana S, Kundu DJ, 2. 742 et al. The PRIDE database and related tools and resources in 2019: improving support for 743 quantification data. Nucleic Acids Res. 2019;47(D1):D442-D50. Epub 2018/11/06. doi: 744 10.1093/nar/gky1106. PubMed PMID: 30395289; PubMed Central PMCID: 745 PMCPMC6323896. 746 Vizcaino JA, Deutsch EW, Wang R, Csordas A, Reisinger F, Rios D, et al. 3. 747 ProteomeXchange provides globally coordinated proteomics data submission and 748 dissemination. Nat Biotechnol. 2014;32(3):223-6. Epub 2014/04/15. doi: 10.1038/nbt.2839. 749 PubMed PMID: 24727771; PubMed Central PMCID: PMCPMC3986813. 750 Martens L, Vizcaino JA. A Golden Age for Working with Public Proteomics Data. 4. 751 Trends Biochem Sci. 2017;42(5):333-41. Epub 2017/01/26. doi: 10.1016/j.tibs.2017.01.001. 752 PubMed PMID: 28118949; PubMed Central PMCID: PMCPMC5414595. 753 Romanov N, Kuhn M, Aebersold R, Ori A, Beck M, Bork P. Disentangling Genetic 5. 754 and Environmental Effects on the Proteotypes of Individuals. Cell. 2019;177(5):1308-18 e10. 755 Epub 2019/04/30. doi: 10.1016/j.cell.2019.03.015. PubMed PMID: 31031010; PubMed 756 Central PMCID: PMCPMC6988111. 757 Skinnider MA, Foster LJ. Meta-analysis defines principles for the design and analysis 6. 758 of co-fractionation mass spectrometry experiments. Nat Methods. 2021;18(7):806-15. Epub 759 2021/07/03. doi: 10.1038/s41592-021-01194-4. PubMed PMID: 34211188. Ochoa D, Jarnuczak AF, Vieitez C, Gehre M, Soucheray M, Mateus A, et al. The 760 7. 761 functional landscape of the human phosphoproteome. Nat Biotechnol. 2020;38(3):365-73. 762 Epub 2019/12/11. doi: 10.1038/s41587-019-0344-3. PubMed PMID: 31819260; PubMed 763 Central PMCID: PMCPMC7100915. 764 8. Vaudel M, Verheggen K, Csordas A, Raeder H, Berven FS, Martens L, et al. 765 Exploring the potential of public proteomics data. Proteomics. 2016;16(2):214-25. Epub 766 2015/10/10. doi: 10.1002/pmic.201500295. PubMed PMID: 26449181; PubMed Central 767 PMCID: PMCPMC4738454. 768 9. Kumar D, Yadav AK, Jia X, Mulvenna J, Dash D. Integrated Transcriptomic-769 Proteomic Analysis Using a Proteogenomic Workflow Refines Rat Genome Annotation. Mol 770 Cell Proteomics. 2016;15(1):329-39. Epub 2015/11/13. doi: 10.1074/mcp.M114.047126. 771 PubMed PMID: 26560066; PubMed Central PMCID: PMCPMC4762527. 772 Brunet MA, Brunelle M, Lucier JF, Delcourt V, Levesque M, Grenier F, et al. 10. 773 OpenProt: a more comprehensive guide to explore eukaryotic coding potential and 774 proteomes. Nucleic Acids Res. 2019;47(D1):D403-D10. Epub 2018/10/10. doi: 775 10.1093/nar/gky936. PubMed PMID: 30299502; PubMed Central PMCID: 776 PMCPMC6323990. 777 11. Levitsky LI, Kliuchnikova AA, Kuznetsova KG, Karpov DS, Ivanov MV, Pyatnitskiy 778 MA, et al. Adenosine-to-Inosine RNA Editing in Mouse and Human Brain Proteomes. 779 Proteomics. 2019;19(23):e1900195. Epub 2019/10/03. doi: 10.1002/pmic.201900195. 780 PubMed PMID: 31576663. Li H, Zhou R, Xu S, Chen X, Hong Y, Lu Q, et al. Improving Gene Annotation of the 781 12. 782 Peanut Genome by Integrated Proteogenomics Workflow. J Proteome Res. 2020;19(6):2226-783 35. Epub 2020/05/06. doi: 10.1021/acs.jproteome.9b00723. PubMed PMID: 32367721. 784 13. Jarnuczak AF, Najgebauer H, Barzine M, Kundu DJ, Ghavidel F, Perez-Riverol Y, et al. An integrated landscape of protein expression in human cancer. Sci Data. 2021;8(1):115. 785 43

- 786 Epub 2021/04/25. doi: 10.1038/s41597-021-00890-2. PubMed PMID: 33893311; PubMed
 787 Central PMCID: PMCPMC8065022.
- Prakash A, García-Seisdedos D, Wang S, Kundu DJ, Collins A, George N, et al. An
 integrated view of baseline protein expression in human tissues. bioRxiv.
- 790 2021:2021.09.10.459811. doi: 10.1101/2021.09.10.459811.
- 15. Samaras P, Schmidt T, Frejno M, Gessulat S, Reinecke M, Jarzab A, et al.

792 ProteomicsDB: a multi-omics and multi-organism resource for life science research. Nucleic

793 Acids Res. 2020;48(D1):D1153-D63. Epub 2019/10/31. doi: 10.1093/nar/gkz974. PubMed

794 PMID: 31665479; PubMed Central PMCID: PMCPMC7145565.

- 795 16. Wang M, Herrmann CJ, Simonovic M, Szklarczyk D, von Mering C. Version 4.0 of
- 796 PaxDb: Protein abundance data, integrated across model organisms, tissues, and cell-lines.
- Proteomics. 2015;15(18):3163-8. Epub 2015/02/07. doi: 10.1002/pmic.201400441. PubMed
 PMID: 25656970; PubMed Central PMCID: PMCPMC6680238.
- 17. Uhlen M, Fagerberg L, Hallstrom BM, Lindskog C, Oksvold P, Mardinoglu A, et al.
- 800 Proteomics. Tissue-based map of the human proteome. Science. 2015;347(6220):1260419.
- 801 Epub 2015/01/24. doi: 10.1126/science.1260419. PubMed PMID: 25613900.
- 802 18. Azimifar SB, Nagaraj N, Cox J, Mann M. Cell-type-resolved quantitative proteomics
- 803 of murine liver. Cell Metab. 2014;20(6):1076-87. Epub 2014/12/04. doi:
- 804 10.1016/j.cmet.2014.11.002. PubMed PMID: 25470552.
- 805 19. Deshmukh AS, Murgia M, Nagaraj N, Treebak JT, Cox J, Mann M. Deep proteomics
- 806 of mouse skeletal muscle enables quantitation of protein isoforms, metabolic pathways, and
- transcription factors. Mol Cell Proteomics. 2015;14(4):841-53. Epub 2015/01/27. doi:

808 10.1074/mcp.M114.044222. PubMed PMID: 25616865; PubMed Central PMCID:

- 809 PMCPMC4390264.
- 810 20. Meierhofer D, Halbach M, Sen NE, Gispert S, Auburger G. Ataxin-2 (Atxn2)-Knock-
- 811 Out Mice Show Branched Chain Amino Acids and Fatty Acids Pathway Alterations. Mol
- 812 Cell Proteomics. 2016;15(5):1728-39. Epub 2016/02/07. doi: 10.1074/mcp.M115.056770.
- 813 PubMed PMID: 26850065; PubMed Central PMCID: PMCPMC4858951.
- 814 21. Sarver DC, Kharaz YA, Sugg KB, Gumucio JP, Comerford E, Mendias CL. Sex
 815 differences in tendon structure and function. J Orthop Res. 2017;35(10):2117-26. Epub
 816 2017/01/11. doi: 10.1002/jor.23516. PubMed PMID: 28071813; PubMed Central PMCID:
 817 PMCPMC5503813.
- 818 22. Duda P, Wojcicka O, Wisniewski JR, Rakus D. Global quantitative TPA-based
- 819 proteomics of mouse brain structures reveals significant alterations in expression of proteins
- 820 involved in neuronal plasticity during aging. Aging (Albany NY). 2018;10(7):1682-97. Epub
- 821 2018/07/22. doi: 10.18632/aging.101501. PubMed PMID: 30026405; PubMed Central
- 822 PMCID: PMCPMC6075443.
- 823 23. Harman JC, Guidry JJ, Gidday JM. Comprehensive characterization of the adult ND4
 824 Swiss Webster mouse retina: Using discovery-based mass spectrometry to decipher the total
 825 proteome and phosphoproteome. Mol Vis. 2018;24:875-89. Epub 2019/02/05. PubMed
 826 PMID: 30713425; PubMed Central PMCID: PMCPMC6334985.
- 827 24. Angelidis I, Simon LM, Fernandez IE, Strunz M, Mayr CH, Greiffo FR, et al. An
- atlas of the aging lung mapped by single cell transcriptomics and deep tissue proteomics. Nat
- 829 Commun. 2019;10(1):963. Epub 2019/03/01. doi: 10.1038/s41467-019-08831-9. PubMed
 830 PMID: 30814501; PubMed Central PMCID: PMCPMC6393476.
- 831 25. Zhao Y, Wilmarth PA, Cheng C, Limi S, Fowler VM, Zheng D, et al. Proteome-
- transcriptome analysis and proteome remodeling in mouse lens epithelium and fibers. Exp
- 833 Eye Res. 2019;179:32-46. Epub 2018/10/26. doi: 10.1016/j.exer.2018.10.011. PubMed
- 834 PMID: 30359574; PubMed Central PMCID: PMCPMC6360118.

- 835 26. Huttlin EL, Jedrychowski MP, Elias JE, Goswami T, Rad R, Beausoleil SA, et al. A
 836 tissue-specific atlas of mouse protein phosphorylation and expression. Cell.
- 837 2010;143(7):1174-89. Epub 2010/12/25. doi: 10.1016/j.cell.2010.12.001. PubMed PMID:
 838 21183079.
- 839 27. Linscheid N, Santos A, Poulsen PC, Mills RW, Calloe K, Leurs U, et al. Quantitative
 840 proteome comparison of human hearts with those of model organisms. PLoS Biol.
- 841 2021;19(4):e3001144. Epub 2021/04/20. doi: 10.1371/journal.pbio.3001144. PubMed PMID:
- 842 33872299; PubMed Central PMCID: PMCPMC8084454.
- 843 28. Dudek M, Angelucci C, Pathiranage D, Wang P, Mallikarjun V, Lawless C, et al.
- 844 Circadian time series proteomics reveals daily dynamics in cartilage physiology.
- 845 Osteoarthritis Cartilage. 2021;29(5):739-49. Epub 2021/02/22. doi:
- 846 10.1016/j.joca.2021.02.008. PubMed PMID: 33610821; PubMed Central PMCID:
 847 PMCPMC8113022.
- 848 29. Schroeder S, Hofer SJ, Zimmermann A, Pechlaner R, Dammbrueck C, Pendl T, et al.
- 849 Dietary spermidine improves cognitive function. Cell Rep. 2021;35(2):108985. Epub
- 850 2021/04/15. doi: 10.1016/j.celrep.2021.108985. PubMed PMID: 33852843.
- 851 30. Bundy JL, Vied C, Nowakowski RS. Sex differences in the molecular signature of the
- developing mouse hippocampus. BMC Genomics. 2017;18(1):237. Epub 2017/03/18. doi:
- 853 10.1186/s12864-017-3608-7. PubMed PMID: 28302071; PubMed Central PMCID:
 854 PMCPMC5356301.
- 855 31. Linscheid N, Logantha S, Poulsen PC, Zhang S, Schrolkamp M, Egerod KL, et al.
 856 Quantitative proteomics and single-nucleus transcriptomics of the sinus node elucidates the
- foundation of cardiac pacemaking. Nat Commun. 2019;10(1):2889. Epub 2019/06/30. doi:
- 858 10.1038/s41467-019-10709-9. PubMed PMID: 31253831; PubMed Central PMCID:
 859 PMCPMC6599035.
- Alugubelly N, Mohammed AN, Edelmann MJ, Nanduri B, Sayed M, Park JW, et al.
 Adolescent rat social play: Amygdalar proteomic and transcriptomic data. Data Brief.
- 862 2019;27:104589. Epub 2019/11/02. doi: 10.1016/j.dib.2019.104589. PubMed PMID:
- 863 31673590; PubMed Central PMCID: PMCPMC6817652.
- 864 33. Roffia V, De Palma A, Lonati C, Di Silvestre D, Rossi R, Mantero M, et al. Proteome
 865 Investigation of Rat Lungs subjected to Ex Vivo Perfusion (EVLP). Molecules. 2018;23(12).
 866 Epub 2018/11/24. doi: 10.3390/molecules23123061. PubMed PMID: 30467300; PubMed
 867 Central PMCID: PMCPMC6321151.
- 868 34. Bernier M, Harney D, Koay YC, Diaz A, Singh A, Wahl D, et al. Elucidating the 869 mechanisms by which disulfiram protects against obesity and metabolic syndrome. NPJ
- 870 Aging Mech Dis. 2020;6:8. Epub 2020/07/28. doi: 10.1038/s41514-020-0046-6. PubMed
- 871 PMID: 32714562; PubMed Central PMCID: PMCPMC7374720.
- 872 35. Huang Q, Luo L, Alamdar A, Zhang J, Liu L, Tian M, et al. Integrated proteomics
- 873 and metabolomics analysis of rat testis: Mechanism of arsenic-induced male reproductive
- 874 toxicity. Sci Rep. 2016;6:32518. Epub 2016/09/03. doi: 10.1038/srep32518. PubMed PMID: 27585557: PubMed Control PMCD: PMCPMC5000432
- 875 27585557; PubMed Central PMCID: PMCPMC5009432.
- 876 36. Kaushik G, Spenlehauer A, Sessions AO, Trujillo AS, Fuhrmann A, Fu Z, et al.
- Vinculin network-mediated cytoskeletal remodeling regulates contractile function in the
 aging heart. Sci Transl Med. 2015;7(292):292ra99. Epub 2015/06/19. doi:
- 879 10.1126/scitranslmed.aaa5843. PubMed PMID: 26084806; PubMed Central PMCID:
- 880 PMCPMC4507505.
- 881 37. Vileigas DF, Harman VM, Freire PP, Marciano CLC, Sant'Ana PG, de Souza SLB, et
- al. Landscape of heart proteome changes in a diet-induced obesity model. Sci Rep.
- 883 2019;9(1):18050. Epub 2019/12/04. doi: 10.1038/s41598-019-54522-2. PubMed PMID:
- 884 31792287; PubMed Central PMCID: PMCPMC6888820.

885 Limbutara K, Chou CL, Knepper MA. Quantitative Proteomics of All 14 Renal 38. 886 Tubule Segments in Rat. J Am Soc Nephrol. 2020;31(6):1255-66. Epub 2020/05/03. doi: 887 10.1681/ASN.2020010071. PubMed PMID: 32358040; PubMed Central PMCID: 888 PMCPMC7269347. 889 Devaux S, Cizkova D, Quanico J, Franck J, Nataf S, Pays L, et al. Proteomic Analysis 39. 890 of the Spatio-temporal Based Molecular Kinetics of Acute Spinal Cord Injury Identifies a 891 Time- and Segment-specific Window for Effective Tissue Repair. Mol Cell Proteomics. 892 2016;15(8):2641-70. Epub 2016/06/03. doi: 10.1074/mcp.M115.057794. PubMed PMID: 893 27250205; PubMed Central PMCID: PMCPMC4974342. 894 40. Choi H, Simpson D, Wang D, Prescott M, Pitsillides AA, Dudhia J, et al. 895 Heterogeneity of proteome dynamics between connective tissue phases of adult tendon. Elife. 896 2020;9. Epub 2020/05/13. doi: 10.7554/eLife.55262. PubMed PMID: 32393437; PubMed 897 Central PMCID: PMCPMC7217697. 898 Hodge K, Have ST, Hutton L, Lamond AI. Cleaning up the masses: exclusion lists to 41. 899 reduce contamination with HPLC-MS/MS. J Proteomics. 2013;88:92-103. Epub 2013/03/19. 900 doi: 10.1016/j.jprot.2013.02.023. PubMed PMID: 23501838; PubMed Central PMCID: 901 PMCPMC3714598. 902 Dai C, Fullgrabe A, Pfeuffer J, Solovyeva EM, Deng J, Moreno P, et al. A proteomics 42. 903 sample metadata representation for multiomics integration and big data analysis. Nat 904 Commun. 2021;12(1):5854. Epub 2021/10/08. doi: 10.1038/s41467-021-26111-3. PubMed 905 PMID: 34615866; PubMed Central PMCID: PMCPMC8494749. 906 43. Evans R, O'Neill M, Pritzel A, Antropova N, Senior A, Green T, et al. Protein 907 complex prediction with AlphaFold-Multimer. bioRxiv. 2021:2021.10.04.463034. doi: 908 10.1101/2021.10.04.463034. 909 Bregman CL, Adler RR, Morton DG, Regan KS, Yano BL, Society of Toxicologic P. 44. 910 Recommended tissue list for histopathologic examination in repeat-dose toxicity and 911 carcinogenicity studies: a proposal of the Society of Toxicologic Pathology (STP). Toxicol 912 Pathol. 2003;31(2):252-3. Epub 2003/04/17. doi: 10.1080/01926230390183751. PubMed 913 PMID: 12696587. 914 Walzer M, García-Seisdedos D, Prakash A, Brack P, Crowther P, Graham RL, et al. 45. 915 Implementing the re-use of public DIA proteomics datasets: from the PRIDE database to 916 Expression Atlas. bioRxiv. 2021:2021.06.08.447493. doi: 10.1101/2021.06.08.447493. 917 46. Sollner JF, Leparc G, Hildebrandt T, Klein H, Thomas L, Stupka E, et al. An RNA-918 Seq atlas of gene expression in mouse and rat normal tissues. Sci Data. 2017;4:170185. Epub 919 2017/12/13. doi: 10.1038/sdata.2017.185. PubMed PMID: 29231921; PubMed Central 920 PMCID: PMCPMC5726313. 921 Sudmant PH, Alexis MS, Burge CB. Meta-analysis of RNA-seq expression data 47. 922 across species, tissues and studies. Genome Biol. 2015;16:287. Epub 2015/12/24. doi: 923 10.1186/s13059-015-0853-4. PubMed PMID: 26694591; PubMed Central PMCID: 924 PMCPMC4699362. 925 48. Zheng-Bradley X, Rung J, Parkinson H, Brazma A. Large scale comparison of global 926 gene expression patterns in human and mouse. Genome Biol. 2010;11(12):R124. Epub 927 2010/12/25. doi: 10.1186/gb-2010-11-12-r124. PubMed PMID: 21182765; PubMed Central 928 PMCID: PMCPMC3046484. 929 49. Liao BY, Zhang J. Evolutionary conservation of expression profiles between human 930 and mouse orthologous genes. Mol Biol Evol. 2006;23(3):530-40. Epub 2005/11/11. doi: 931 10.1093/molbev/msj054. PubMed PMID: 16280543. 932 50. Prasad A, Kumar SS, Dessimoz C, Bleuler S, Laule O, Hruz T, et al. Global 933 regulatory architecture of human, mouse and rat tissue transcriptomes. BMC Genomics.

- 934 2013;14:716. Epub 2013/10/22. doi: 10.1186/1471-2164-14-716. PubMed PMID: 24138449;
 935 PubMed Central PMCID: PMCPMC4008137.
- 51. Athar A, Fullgrabe A, George N, Iqbal H, Huerta L, Ali A, et al. ArrayExpress update
 from bulk to single-cell expression data. Nucleic Acids Res. 2019;47(D1):D711-D5. Epub
 2018/10/26. doi: 10.1093/nar/gky964. PubMed PMID: 30357387; PubMed Central PMCID:
- 939 PMCPMC6323929.
- 52. Cox J, Mann M. MaxQuant enables high peptide identification rates, individualized
 p.p.b.-range mass accuracies and proteome-wide protein quantification. Nat Biotechnol.
- 942 2008;26(12):1367-72. Epub 2008/11/26. doi: 10.1038/nbt.1511. PubMed PMID: 19029910.
- 943 53. Tyanova S, Temu T, Cox J. The MaxQuant computational platform for mass
- 944 spectrometry-based shotgun proteomics. Nat Protoc. 2016;11(12):2301-19. Epub 2016/11/04.
 945 doi: 10.1038/nprot.2016.136. PubMed PMID: 27809316.
- 946 54. Team RC. R: A language and environment for statistical computing. R
- Foundation for Statistical Computing, Vienna, Austria. ISBN 3-900051-07-0, URL
 http://www.R-project.org/. 2013.
- 949 55. McInnes L HJ, Melville J. UMAP: Uniform Manifold Approximation and Projection 950 for dimension reduction. arXiv. 2018.
- 951 56. Wu G, Haw R. Functional Interaction Network Construction and Analysis for Disease
- 952 Discovery. Methods Mol Biol. 2017;1558:235-53. Epub 2017/02/06. doi: 10.1007/978-1-
- 953 4939-6783-4_11. PubMed PMID: 28150241.
- 954 57. Raudvere U, Kolberg L, Kuzmin I, Arak T, Adler P, Peterson H, et al. g:Profiler: a
- 955 web server for functional enrichment analysis and conversions of gene lists (2019 update).
- 956 Nucleic Acids Res. 2019;47(W1):W191-W8. Epub 2019/05/09. doi: 10.1093/nar/gkz369.
- 957 PubMed PMID: 31066453; PubMed Central PMCID: PMCPMC6602461.