

Open access • Posted Content • DOI:10.1101/268953

Rapid proteotyping reveals cancer biology and drug response determinants in the NCI-60 cells — Source link ☑

Tiannan Guo, Tiannan Guo, Augustin Luna, Vinodh N. Rajapakse ...+27 more authors

Institutions: Harvard University, Westlake University, National Institutes of Health, RWTH Aachen University ...+5 more institutions

Published on: 21 Feb 2018 - bioRxiv (Cold Spring Harbor Laboratory)

Topics: Proteome and Systems biology

Related papers:

- Colorectal Cancer Cell Line Proteomes Are Representative of Primary Tumors and Predict Drug Sensitivity.
- Integrating the NCI-60 Data with "Omics" for Drug Discovery
- The NCI Transcriptional Pharmacodynamics Workbench: A Tool to Examine Dynamic Expression Profiling of Therapeutic Response in the NCI-60 Cell Line Panel.
- · Proteomic profiling of breast cancer cell lines and models
- · Data Descriptor: Proteomic profiling across breast cancer cell lines and models

Share this paper: 🚯 🄰 🛅 🗠

1 Rapid proteotyping reveals cancer biology and drug response determinants in the NCI-

- 2 **60 cells**
- 3
- 4 Tiannan Guo^{1,19}*[#], Augustin Luna^{2,3}*, Vinodh N Rajapakse⁴*, Ching Chiek Koh¹*[†],
- 5 Zhicheng Wu¹⁹, Michael P Menden⁵, Yongran Cheng¹⁹, Laurence Calzone⁶, Loredana
- 6 Martignetti⁶, Alessandro Ori⁷, Murat Iskar⁸¹, Ludovic Gillet¹, Qing Zhong⁹, Sudhir Varma
- 7 ¹⁰, Uwe Schmitt ¹¹, Peng Qiu ¹², Yaoting Sun ¹⁹, Yi Zhu ^{1,19}, Peter J Wild ⁹, Mathew J Garnett
- 8 ¹³, Peer Bork ^{8, 14,15,16}, Martin Beck ^{8, 17}, Julio Saez-Rodriguez ⁵, William C. Reinhold ⁴, Chris
- 9 Sander ^{2,3}, Yves Pommier ⁴ [#], Ruedi Aebersold ^{1, 18} [#]
- 10
- 11 * Equal contribution
- 12 # correspondence
- 13

14 Affiliations

- 15 1, Department of Biology, Institute of Molecular Systems Biology, ETH Zurich, Switzerland
- 16 2, Department of Biostatistics and Computational Biology, Dana-Farber Cancer Institute,
- 17 Boston, MA 02115, USA
- 18 3, Department of Cell Biology, Harvard Medical School, Boston, MA 02115, USA
- 19 4, Developmental Therapeutics Branch, Center for Cancer Research, National Cancer
- 20 Institute, National Institutes of Health, Bethesda, MD 20892, United States
- 21 5, RWTH Aachen University, Faculty of Medicine, Joint Research Centre for Computational
- 22 Biomedicine (JRC-COMBINE), Germany
- 6, Institut Curie, PSL Research University, INSERM, U900, Mines Paris Tech, F-75005,
- 24 Paris, France.
- 25 7, Leibniz Institute on Aging, Fritz Lipmann Institute (FLI), Beutenbergstrasse 11, 07745
- 26 Jena, Germany
- 8, Structural and Computational Biology Unit, European Molecular Biology Laboratory,
- 28 69117 Heidelberg, Germany
- 29 9, Institute of Surgical Pathology, University Hospital Zurich, Zurich, Switzerland
- 30 10, HiThru Analytics, Laurel, MD 20707, USA
- 31 11, Scientific IT Services, ETH Zurich, Switzerland
- 32 12, Department of Biomedical Engineering, Georgia Institute of Technology and Emory
- 33 University, 313 Ferst Dr., Atlanta, GA 30332, US
- 13, Sanger Institute, Wellcome Trust Genome Campus, Hinxton, Cambridge CB10 1SA, UK

- 14, Molecular Medicine Partnership Unit, University of Heidelberg and European Molecular
- 36 Biology Laboratory, 69120 Heidelberg, Germany
- 37 15, Max Delbrück Centre for Molecular Medicine, 13125 Berlin, Germany
- 16, Department of Bioinformatics, Biocenter, University of Würzburg, 97074 Würzburg,
- 39 Germany
- 40 17, Cell Biology and Biophysics Unit, European Molecular Biology Laboratory, 69117
- 41 Heidelberg, Germany
- 42 18, Faculty of Science, University of Zurich, Zurich, Switzerland
- 43 19, Westlake Institute for Advanced Study, Westlake University, Hangzhou, Zhejiang, P. R.
- 44 China
- 45
- 46 [†], current address: Sanger Institute, Wellcome Trust Genome Campus, Hinxton, Cambridge
- 47 CB10 1SA, UK
- 1, current address: Division of Molecular Genetics, German Cancer Research Center (DKFZ),
- 49 Im Neuenheimer Feld 280, 69120 Heidelberg, Germany
- 50

51 Summary

- 52
- 53 We describe the rapid and reproducible acquisition of quantitative proteome maps for the
- 54 NCI-60 cancer cell lines and their use to reveal cancer biology and drug response
- determinants. Proteome datasets for the 60 cell lines were acquired in duplicate within 30
- 56 working days using pressure cycling technology and SWATH mass spectrometry. We
- 57 consistently quantified 3,171 SwissProt proteotypic proteins across all cell lines, generating a
- 58 data matrix with 0.1% missing values, allowing analyses of protein complexes and pathway
- 59 activities across all the cancer cells. Systematic and integrative analysis of the genetic
- 60 variation, mRNA expression and proteomic data of the NCI-60 cancer cell lines uncovered
- 61 complementarity between different types of molecular data in the prediction of the response to
- 62 240 drugs. We additionally identified novel proteomic drug response determinants for
- 63 clinically relevant chemotherapeutic and targeted therapies. We anticipate that this study
- 64 represents a landmark effort toward the translational application of proteotypes, which reveal
- biological insights that are easily missed in the absence of proteomic data.

66 Introduction

67

To date, mainly owing to the maturity and availability of high throughput DNA- and 68 RNA- based techniques, forays into the molecular landscape of diseases, in particular cancers, 69 have primarily focused on genomics and transcriptomics ¹⁻³. Protein-level measurements, 70 although showing great potential for providing the granularity and details necessary for 71 personalized therapeutic decisions, are underutilized due to technical hurdles. Advances in 72 data-dependent acquisition (DDA) mass spectrometry (MS) have permitted quantitative 73 proteomic profiling of about 100 tumor samples using multi-dimensional fractionated MS 74 analyses of each sample ⁴⁻⁶, demonstrating the added value of protein measurement in 75 classifying tumor samples. Nevertheless, such DDA workflows suffer from relatively lower 76 77 sample-throughput, relatively higher sample consumption and technical complexity, precluding their routine use in clinically relevant applications (e.g. drug response prediction) 78 on the speed and scale achieved by genomic and transcriptomic approaches ^{2,3}. 79 80 81 To achieve reproducible and high throughput proteomic profiling, we have developed a workflow ^{7,8} integrating pressure cycling technology (PCT), an emerging sample 82

83 preparation method that accelerates and standardizes sample preparation for proteomic profiling⁹, together with SWATH-MS, an MS-based proteomic technique that consists of 84 85 data independent acquisition (DIA) and a targeted data analysis strategy with unique advantages over other MS-based proteomic methods ^{10, 11}. With this technique all MS-86 measurable peptides of a sample are fragmented and recorded in a recursive fashion, thus 87 generating digital proteome maps that can be used to reproducibly detect and quantify 88 proteins across high numbers of samples without the need of isotope labeling. The PCT-89 SWATH technique thus significantly increases the sample throughput and data reproducibility 90 providing excellent quantitative accuracy, and in the meantime reduces sample consumption 91 to ca. 1 microgram of total peptide mass per sample ^{7, 8}. 92

93

In this study, we describe the acquisition of proteome maps of the NCI-60 cell lines in duplicate by PCT-SWATH. The 120 proteome maps were acquired within 30 working days on a single instrument and each sample consumed ca. 1 microgram of total peptide mass. We consistently quantified 3,171 SwissProt proteotypic proteins across all cell lines, generating a data matrix (120 proteomes vs. 3171 proteins) with 0.1% missing values. Raw signals of each peptide and protein in each sample were curated with an expert system. The NCI-60 human

cancer cell line panel contains 60 lines from 9 different tissue types ¹². The NCI-60 have been 100 molecularly and pharmacologically characterized with unparalleled depth and coverage, 101 102 offering a prime in vitro model to further our understanding of cancer biology and cellular responses to anti-cancer agents ^{12, 13}. Discoveries enabled by the NCI-60 in recent years 103 include the development of the FDA approved drugs oxaliplatin for the treatment of colon 104 cancers ¹⁴, eribulin for metastatic breast cancers ¹², bortezomib for the treatment of multiple 105 myeloma¹⁵, and rhomidepsin for cutaneous T-cell lymphomas¹⁶. The sensitivity of the NCI-106 60 has been measured for over 100,000 synthetic or natural compounds derived from a wide 107 range of academic and industrial sources ¹², constructing the most comprehensive resource for 108 cancer pharmacological research. The proteomic data complement the existing NCI-60 109 110 molecular landscapes, allowing systematic investigation of the complementarity among genomics, transcriptomics and proteomics in a number of applications. 111 112

The proteome of the NCI-60 cells has been analyzed previously by data dependent 113 analysis (DDA), a commonly used discovery mass spectrometry technique ¹⁷. Whereas the 114 study reported the cumulative identification of 10,350 IPI proteins from about 1,000 115 116 fractionated and kinase-enriched sample runs, only 492 IPI proteins were quantified across the NCI-60 cell lines without missing value. The present study thus extends the number of 117 consistently quantified proteins, in duplicates, to 3,171, with a ca. six-fold increase. The high 118 quality proteomic data were used for pharmacoproteomic analysis of the response of the cell 119 panel to 240 anti-cancer drugs, resulting in the identification of novel proteomic drug 120 response determinants for clinically relevant chemotherapeutic and targeted therapies. 121

122 **Results**

123

124 Acquisition of the NCI-60 proteome maps

125

We applied the PCT-SWATH workflow ⁷ to generate quantitative proteome maps of 126 the NCI-60 cell lines in technical replicates, resulting in the generation of 120 SWATH maps 127 with high reproducibility at the raw data level (Supplementary Fig. 1). The PCT-assisted 128 sample preparation took about 18 working days and the SWATH-MS analyses consumed 129 130 about 12 working days. Thus, the entire process, from sample preparation to data acquisition, was accomplished within 30 working days, at an unprecedented sample-throughput compared 131 to other cancer proteomic research of comparable scale ^{4-6, 17}, which is due to the elimination 132 of multiple dimensional fractionation, using one barocycler and one mass spectrometer 133 134 (Supplementary Fig. 1, Supplementary Table 1). 135 136 SWATH proteome maps contain fragment ion chromatograms from all MSmeasurable peptides, albeit in a highly convoluted form. To interpret the SWATH maps, we 137 138 built a human cancer cell line spectral library containing 86,209 proteotypic peptides, *i.e.* peptides that uniquely identify a specific protein, from 8,056 SwissProt proteins 139 (Supplementary Table 1). Using this library and the OpenSWATH software ¹¹, we identified 140

6,556 protein groups, covering 81% of the library (Supplementary Fig. 2). To avoid
ambiguity of peptide/protein quantification, we limited our analyses to canonical and

143 proteotypic peptides and proteins by excluding protein isoforms, un-reviewed protein

sequences, and peptide/protein sequence variants.

145

We evaluated the technical variation of each measurement through manual inspection
of the OpenSWATH results based on the replicated measurement for each cell line and
observed in multiple cases substantial technical variation. This is probably due to the fact that
cell type-specific interfering signals leads to invalid SWATH assays, and the presence of
irregular liquid chromatography (LC) and MS behavior of certain peptides in the highly
variable proteomic context of the NCI-60 cells. These phenomena have also been observed
previously in selected reaction monitoring (SRM)-based targeted proteomics studies ¹⁸.

To obtain high accuracy quantitative data for the cell lines, we further developed an expert system, *i.e.* DIA-expert (see Methods), to refine the peptide identification and

quantification provided by automated analysis tools like OpenSWATH (Fig. 1A). We thus 156 157 excluded proteins and peptides that were not reproducibly quantified in technical replicates 158 and focused our analyses on a shorter list of 22,554 proteotypic peptides from 3,171 proteins, 159 with 8% missing values at the peptide level and 0.1% missing values at the protein level across all MS runs (Supplementary Table 1). On average, 7 peptide precursors and 6 unique 160 peptide sequences were identified for each protein (Fig. 1B). Several proteins were identified 161 with over 200 peptides (Fig. 1C). The proteins excluded by DIA-expert may not be incorrect 162 identifications, but rather proteins that could not achieve reproducible quantification by the 163 164 existing algorithm across all cell lines due to either technical issues, for instance the signal-tonoise ratio, or biological issues such as post-translational modifications and splicing variants. 165 166 Improved computational methods will likely rescue some of them in the future. 167 Most peptides for the 3,171 proteins were consistently quantified in all cell lines at 168 both MS1 and MS2 levels. Two representative peptides are shown in Fig. 1A. The coefficient 169 of determination (R^2) between technical replicates, for overall expression of peptides (Fig. 170 1D) and proteins (Fig. 1E), were 0.974 and 0.978, respectively, with a dynamic range over 5 171 172 orders of magnitude (Fig. 1F). We provide the raw MS signals for each quantitative value in Supplementary File 1, allowing visual inspection of the MS signal for every peptide in each 173 sample. When we limited the minimal peptide number per protein to 2, 3 and 4, fewer 174 proteins were quantified however the quantitative accuracy did not substantially improve 175 (Supplementary Figure 3). 176





Figure 1. The acquisition of NCI-60 proteotype. (A) Representative peptide signals as curated and visualized
by the DIA-expert software. (B) The cumulative number of peptide and peptide precursors identified for each
protein. (C) The distribution of peptide precursors and peptides per protein. The overall Pearson correlation
between technical replicates at the peptide level (D) and the protein level (E). Here, the log10 transformed
intensity of each peptide/protein in each cell line technical replicate is plotted in the heatmap. (F) Dynamic range
of the MS signals for 22,968 proteotypic peptides.

185

186 Characterization of the NCI-60 quantitative proteomes

187

The landscape of the 120 thus measured proteotypes is displayed in Fig. 2A. All 188 technical replicates were clustered together using an unsupervised method based on the 189 quantified proteotypes, confirming high quantitative accuracy. In most cases, the proteotypes 190 are not strikingly different across different cancer cell lines, in sharp contrast with the distinct 191 proteomes of tumor versus non-tumor kidney tissues ⁷. The median coefficient of variation 192 (CV) of the protein intensity in different cells was 48%. The CV demonstrated a low 193 dependence on protein abundance, as evident from the distribution of its values for different 194 expression level quantile groups of the measured proteins (Fig. 2B). We then compared our 195 data with the previously reported proteome of the NCI-60 cells using DDA-MS¹⁷. While the 196 DDA data reported comparable number of IPI protein groups to the SwissProt proteotypic 197 protein number from this SWATH data set, the SWATH data exhibited much higher degree of 198

- 199 consistency (Supplementary Table 2 and Supplementary Fig. 5) and better quantitative
- 200 accuracy (Supplementary Fig. 6-7).
- 201



- 202
- 203

204 Figure 2. Characterizing NCI-60 quantitative proteomes. (A) Heatmap overview of NCI-60 proteotype data 205 matrix. 3,171 Swiss-Prot proteins were quantified in 120 SWATH runs. (B) Variation of protein expression, for 206 all proteins (All) and proteins in each abundance quantile group (from low abundance to high abundance). (C) 207 Density plot of correlation of determination between pairs of random proteins versus pairs of proteins within a 208 complex. (D) Stoichiometry variation of protein complexes in the NCI-60 cells. The x-axis shows the average 209 Pearson correlation of each protein complex across the NCI-60. The y-axis shows the average abundance of proteins in a complex. Stable complexes tend to show higher values of average Pearson correlation. (E) Protein 210 211 and mRNA expression of XRCC6/Ku70 and XRCC5/Ku80. (F) Visualization of pathway activity in NCI-60 212 proteotypes. More detailed pathway annotations for this Google map are provided in Supplementary File 2. 213 **Quantification of drug-responsiveness related proteins** 214

215

216 The proteotypes covered 105 protein targets for FDA-approved anti-cancer

compounds, 661 protein drug targets annotated in DrugBank¹⁹ (including 68 drug

metabolizing enzymes, 5 drug carriers, and 15 drug transporters), 694 proteins known to 218 participate in human diseases ^{19, 20}, and 58 human protein kinases, in addition to proteins 219 involved in various biological functions (Supplementary Table 3). Some kinases were found 220 to be broadly expressed in most cells with high abundance, including MST4 and WNK1 221 (Supplementary Fig. 4), consistent with previous reports ^{21, 22}. Other kinases were highly 222 expressed in specific cell lines, for example, EGFR in the breast cancer cell line 223 MDAMB468, ERBB2 in SKOV3 cells, and CDK6 in MOLT4 cells, in agreement with 224 previous studies using antibody-based methods ^{20, 23}. 225 226 One unique benefit of our proteomic data set, compared to genomic and transcriptomic 227 228 data, is its capacity to reveal more accurate information about the abundance of protein complexes and their stoichiometry ²⁴. Our measurements included 101 protein complexes 229 comprising 1,045 proteins (Supplementary Table 4) from a curated resource ²⁴. Significantly 230 higher Pearson correlation coefficients for pairs of proteins that are part of a complex further 231 232 supported the quantitative accuracy of our data matrix (Fig. 2C). We applied our computational pipeline for analyzing co-expression of protein complex numbers ²⁴ to the 233 234 NCI-60 proteotype data and confirmed conserved stoichiometry of protein complexes such as the prefoldin and MCM complexes in various cell lines (Fig. 2D). In a specific case, we 235

observed a high correlation between the protein expression of XRCC6/Ku70 and

237 XRCC5/Ku80, a critical heterodimer involved in DNA repair and responsible for resistance to

radiotherapy and chemotherapy. Ku80 is degraded when not bound to Ku70 $^{25, 26}$.

239 Remarkably, this correlation is not detectable using mRNA measurements (Fig. 2E),

240 indicating that expression of Ku80 is tightly regulated by protein degradation mechanisms

241 independent of cancer types. Indeed, a recent report has shown that RNF8, an E3 ubiquitin

ligase, regulates the expression of Ku80 via its removal from DNA double strand break sites

- 243
- 244

245 Google-map-based visualization of cancer signaling pathways

and its degradation through ubiquitination ²⁷.

246

The NCI-60 proteotypes cover 648 proteins in the Atlas of Cancer Signaling Networks (ACSN), a manually curated pathway database presenting published facts about biochemical reactions involved in cancer using a Google-Maps-style visualization (**Supplementary Fig. 8**) ²⁸. When mapping the mean protein expression per cancer type, we found that multiple pathways in different cell types, including apoptosis, cell survival, motility and DNA repair

among others, displayed a similar pattern (Supplementary File 2), consistent with the fact 252 that the immortal cells retain cancer hallmarks after artificial culturing ²⁹. An example of a 253 clear proteotypic pattern is the delta isoenzyme of protein kinase C, *i.e.* PRKCD, involved in 254 DNA repair and a drug target that has been tested in various cancers 30 . It was reported to be 255 absent in four renal clear cell carcinoma lines ³¹. In agreement, this protein stood out in our 256 visualization, with significantly lower protein expression in renal carcinoma, relative to the 257 average expression in the NCI-60 panel. We provided detailed instructions on how to navigate 258 through the atlas and explore protein abundance in each cancer cell line (see Supplementary 259 260 **File 2**).

261

262 We next compared the activity of cellular pathways using ROMA (Representation and quantification Of Module Activities)³² (Fig. 2F), a gene-set-based quantification algorithm. 263 264 This approach revealed substantial diversity of pathway activity between different proteotypes as evidenced by two-tailed *t*-tests of activity scores (*P*-value < 0.05). When mapping activity 265 266 scores onto ACSN, some tissue specificities were revealed, with particular cell line proteotypes displaying distinct patterns of pathway activity. For instance, the activity of 267 268 apoptosis (with both Caspases and Apoptosis Genes modules) was found to be significantly higher in ovarian cell lines (see Supplementary Table 5). Although there are only two 269 prostate cancer cell lines in the panel, our analysis was able to highlight modules including 270 "AKT-mTOR" and "Apoptosis", whose differential activity can be attributed to HSP90AA1 271 and PRDX. The latter protein has been independently reported to be overexpressed in prostate 272 tumors ³³. 273

274

275 Accessibility of the NCI-60 proteotypes

276

To enable easy data access, visualization, and comparison with other NCI-60 data sets, 277 we have incorporated the SWATH data into the CellMiner database ^{13, 34}. CellMiner allows 278 279 the direct download of the data, as well as comparative and integrative analyses with other molecular data and pharmacological data, e.g. sensitivity of each cell line to over 20,000 280 281 compounds, and the manual inspection of specific genes, up to 150 per query. The detailed 282 instructions for using this resource are provided on the project website (https://discover.nci.nih.gov/cellminer/) and in Supplementary Fig. 9. We have also 283 deposited raw data and processed data matrices of the NCI-60 proteotype in public databases, 284 including PRIDE ³⁵ and ExpressionArray ³⁶. 285

286

287 Predicting drug responsiveness

288

289 The robust, quantitative proteomic data, with almost no missing values, permitted systematic investigation of whether integration of the SWATH-based proteotype with existing 290 genomic and transcriptomic features improves the prediction of drug responsiveness 291 (Supplementary Table 6). We generated various combinations of molecular features, and 292 evaluated their predictive power using the Pearson correlation between predicted and 293 observed drug response values for 240 FDA-approved or investigational compounds in 294 CellMiner^{13, 34, 37}. Each compound is assigned a NSC (National Service Center) identifier 295 upon submission to the National Cancer Institute for evaluation in the NCI-60 panel. The 296 297 largest groups of drugs with target annotations are those that interfere with DNA synthesis 298 and the DNA damage response, including topoisomerase inhibitors. The drug set also contains dozens of targeted agents, including 18 serine and threonine kinase inhibitors and 18 tyrosine 299 300 kinase inhibitors (Fig. 3A).





Using the elastic net algorithm, we then developed multivariate linear models to predict the NCI-60 response for each compound based on genomic, transcriptomic and proteomic features. The Pearson's correlation between observed drug response values and leave-one-out cross validation-predicted response values was applied to evaluate the

- 316 performance of each predictive model.
- 317

As different numbers of features were measured for each omics data set, two strategies 318 were adopted in the modeling analyses. First, we used all omics features (2,282 DNA 319 320 mutations, 14,969 mRNAs and 3,171 proteins), separately and combinatorically, as inputs to 321 evaluate the general performance. Second, we selected 1,566 features that were available for 322 all three molecular data types (denoted as common features). In both cases, we obtained valid 323 models for 224 (93%) of the drugs. The predictive power achieved with all features was 324 slightly higher than that obtained using the common features for all three data types (Fig. 3A); a likely reason for this is that the latter excluded some genomic and transcriptomic features 325 326 not detected at the protein level. We accordingly derived our main analysis results from data including all available molecular features. Our modeling led to the discovery of valid 327 328 biomarkers for drug responsiveness prediction. For instance, we found that the mRNA expression of SLFN11, strikingly responsible for the sensitivity of 45 compounds, out of 329 which 39 were FDA-approved drugs including topoisomerase inhibitors, alkylating agents, 330 and DNA synthesis inhibitors, was the most dominant indicator, in agreement with our 331 previous report ³⁸ (**Supplementary Table 7**). Fourteen ATP-binding cassette family 332 transporters, detected as mutation, transcript or protein levels, were found responsible for 333 sensitivity prediction of 51 compounds including chemotherapeutic agents and protein-334 targeting agents such as HDAC inhibitor Depsipeptide, HSP90 inhibitor Alvespimycin, 335 mTOR inhibitor Temsirolimus and BCR-ABL inhibitor Nilotinib (Supplementary Table 7). 336 337 For ease of reproducibility of data analysis, we developed a Docker container 338

339 (described in **Methods**) that includes our code and other essential dependencies, allowing all340 analyses to be replicated and extended for this and other omics data sets.

341

- 343 Synergies among mutations, transcripts and proteins
- 344

Our pipeline led to the identification of valid models for 224 compounds 345 346 (Supplementary Table 8). Given the relatively small sample size, it was not surprising that accurate predictive models could not be found for every drug, particularly those with limited 347 numbers of responsive lines. We found that the SWATH-MS derived proteotypes displayed 348 higher percentage of predictive features than mutations and transcripts. 1,090 (34%) out of 349 350 3,171 SWATH features are predictive, while 284 (12%) out of 2,282 features for mutations and 1,976 (13%) out of 14,969 transcripts were selected in the models. In general, the 351 SWATH data outperformed the mutation data, however, the mRNA expression data set has 352 353 about a five to six-fold higher number of features than the protein and mutation data sets (Fig. **3A**) and exhibited better overall performance (**Fig. 3C**). 354

355

Our analyses revealed notable synergies among the different molecular measurements. 356 357 Each type of molecular data set demonstrated indispensable benefits in predicting the response to certain drugs/compounds. The responsiveness of 35 compounds (16%) out of 224 358 359 was best predicted with SWATH data, whereas 107 compounds (48%) were best predicted by 360 SWATH data or by combining SWATH data with transcripts and/or DNA data. The most 361 accurate models for over half of the compounds required at least two different types of molecular features. We then computed accumulative sum of Pearson correlation coefficient 362 based on drug responsiveness prediction and observed significant contribution of SWATH 363 data (Fig. 3D). We also compared the predictive power of the DDA data to the SWATH data. 364 While the DDA data were able to generate elastic net models for comparable number of drugs 365 (Fig. 3E), the number of protein predictors is much lower than SWATH data over some 366 367 overlap (Fig. 3F).

368

369 Drug responsiveness prediction

370

Based on the integration of various data sets, global drug response patterns were predicted for the 158 well-modeled drugs (**Fig. 4**, see Methods), with predictive molecular features for individual compounds provided in **Supplementary Table 8**. The data generated from this computational pipeline were validated by the recovery of established pharmacogenomic knowledge. For instance, the mutational status of BRAF was the top predictive molecular feature for sensitivity to BRAF inhibitors, *e.g.* vemurafenib (NSC 761431) and dabrafenib (NSC 764134), and this association was particularly evident in

- 378 melanomas. Activated BRAF mutational status also sensitized cells to the MEK inhibitor
- hypothemycin (NSC: 354462), as has been previously described ³⁹.
- 380
- 381



Figure 4. Predictive power for 224 compounds using different types of omics data. We applied elastic net and
 cross validation to evaluate the drug response predictive accuracy for each omics data set and combinations of
 data sets for 224 drugs which could be effectively modeled. Drug response prediction accuracies across input
 data types are clustered without supervision. MoA of compounds and clinical status of the compounds are

- 387 colored. Each column indicates an input data type or combination of types; each row represents a compound.
- 388 The color indicates the predictive power measured by Pearson correlation of cross-validation predicted versus
- 389 observed drug response values. Black indicates that a valid elastic net model could not be obtained.
- 390

Sensitivity to the antimetabolite 6-thioguanine (6-TG, NSC: 752) (Fig. 5A) was 391 392 predicted by protein expression of NUDT5 and MAT2B within an elastic net model composed of 5 proteomic features: NUDT5, MAT2B, CD47, STX12 and GFAP. The cross-393 394 validation accuracy with this compound and the SWATH-MS data was relatively low (r =0.27), probably due to instability in the selected predictive features with limited sample size. 395 396 Still, we find that for the two strongest predictors in the model, NUDT5 and MAT2B, the expression data were significantly correlated with the activity of 6-TG (Fig. 5B and 5C). 397 398 Additionally, we were able to relate the inter-connected activities of these two proteins to the mechanism of action for 6-TG. In the purine salvage pathway, HPRT1 catalyzes synthesis of 399 inosine monophosphate from hypoxanthine and phosphoribosyl pyrophosphate (PRPP), with 400 production of the latter stimulated by NUDT5. 6-TG can substitute for hypoxanthine, 401 ultimately yielding altered nucleotides that are toxic upon incorporation into DNA ⁴⁰. PRPP is 402 still required, so low NUDT5 expression could possibly induce 6-TG resistance. This is 403 consistent with our NCI-60 data and recent experimental work showing that depletion of 404 NUDT5 confers resistance to 6-TG⁴¹. As noted in Fig. 5A, a metabolite of 6-TG, 405 406 thioguanosine monophosphate (TGMP) can be inactivated by methylation. Production of the 407 methyl group donor, S-adenosylmethionine (SAMe), is catalyzed by the methionine adenosyltransferase IIa (MAT2A) enzyme. The MAT2B protein, exhibiting high correlation 408 with MAT2A (Fig. 5D), is a regulatory component of MAT which may enhance feedback 409 inhibition by SAMe ⁴². Increased MAT inhibition and diminished TGMP methylation may 410 shunt more TGMP toward DNA incorporation, enhancing the 6-TG response. In spite of its 411 relatively low cross-validation accuracy, the presented model may provide a starting point for 412 further exploration, in light of the supporting prior research. 413



415

Figure 5. Drug responsiveness predicted by SWATH data. (A) molecular mechanisms of 6TG. (B)
correlation between NUDT5 protein expression and 6-TG activity. (C) correlation between MAT2B protein
expression and 6-TG activity. (D) correlation between MAT2B and MAT2A protein expression. (E)
LAMTOR3 facilitates MEK/ERK pathway activation by binding MEK and ERK. (F) correlation between
LAMTOR3 protein expression and Vemurafenib activity. (G) Association of BRAF mutation and LAMTOR3
protein expression with Vemurafenib activity.

422

Analysis of the protein kinase inhibitor vemurafenib (NSC 761431) yielded a 423 multivariate model based on BRAF V600E activating mutation status ⁴³ and the protein 424 expression level of LAMTOR3. LAMTOR3 (MP1) is part of an endosomal scaffolding 425 complex that interacts with components of the RAF/MEK/ERK mitogenic signaling pathway 426 (Fig. 5E). In particular, LAMTOR3 binds MEK1 and ERK1, facilitating activation of the 427 latter protein ⁴⁴. Elevated LAMTOR3 protein expression was correlated with vemurafenib 428 429 resistance (r= 0.44, Fig. 5F), consistent with the hypothesis that LAMTOR3 has the capacity to enhance RAF/MEK/ERK pathway signaling downstream from RAF. In particular, 430 increased protein expression of LAMTOR3 was observed in two BRAF mutant cell lines, 431 ME:SK-MEL-5 and ME:LOXIMVI, which are relatively resistant to Vemurafenib (Fig. 5G). 432 Due to the limited number of BRAF mutant cell lines exhibiting relative drug resistance (*i.e.* 2 433 cell lines), additional statistical analyses with sufficient power are not possible. Robust 434 statistical validation of this model may possible when larger cell line databases (e.g. the 435

436 Sanger and Broad resources) expand to include proteomic coverage of LAMTOR3. Still, this

- 437 finding remains relevant in light of the recent research into the activity of LAMTOR3,
- 438 including the observation that reduced LAMTOR3 protein levels decreased the activation of

439 MEK1/2 and ERK1/2 $^{44, 45}$. Additionally, LAMTOR3 has been shown to affect proliferation

- of pancreatic and breast cancers $^{46, 47}$, and has been patented as a diagnostic biomarker for
- 441 breast cancer 47 .
- 442

Our elastic net analysis also produced multiple recurrent predictors with plausible drug 443 response associations. ABCC4 was a negatively weighted predictor for several alkylating 444 agents, including chlorambucil (NSC: 3088), uracil mustard (NSC: 34462), nitrogen mustard 445 (NSC: 762), consistent with its established role as a drug efflux pump ⁴⁸. Another recurrent, 446 negatively-weighted predictor was CTNND1, which was identified for several compounds, 447 448 including bendamustine (NSC: 138783), etoposide (NSC: 141540), valrubicin (NSC: 246131), and carmustine (NSC: 409962). CTNND1 encodes delta-catenin, whose 449 overexpression promotes cell survival through activation of Wnt pathway signaling ⁴⁹. The 450 resulting inhibition of apoptosis ⁵⁰ could plausibly confer resistance to the mentioned DNA-451 452 damage inducing drugs.

453

454 **Discussion**

455

Due to the complementarity of protein and transcript data ^{4-6, 51}, it can be expected that 456 the rapid and consistent quantification of thousands of proteins across a large sample cohort 457 will revel new biological information that is not apparent from the commonly used transcript 458 profiles. However, due to technical limitations, such proteomic cohort datasets have been 459 challenging to acquire. Here, using the NCI-60 cell line compendium, we demonstrate the 460 ability of the PCT-SWATH proteomic technique to consistently quantify in excess of 3000 461 proteins across the 60 cell lines measured in duplicate. The data were acquired in 30 working 462 463 days on a single mass spectrometer and for each sample measurement ca. 1 microgram of total peptide mass was consumed. This has been enabled by the pressure cycling technology 464 which minimizes samples consumption and the data-independent MS data acquisition using 465 SWATH-MS⁷. The data generated and their use to reveal cancer biology and drug response 466 determinants represent a significant advance in the field. 467

The proteome of the NCI-60 cells has been previously measured by extensive sample 469 fractionation and DDA-MS analysis of over 1,000 fractionated samples ¹⁷. In this study, data 470 acquisition for each cell line required an average of about 29.16 hours MS instrument time. 471 472 That shotgun proteomics study reported the cumulative identification of 10,350 IPI proteins over the NCI-60 cell lines. However, only 492 proteins were quantified in all cell lines 473 without missing value. The PCT-SWATH methodology adopted in this study offers an over 474 10-fold increase in sample-throughput, which has allowed us to acquire the proteotype for 475 each cell in the NCI-60 panel in duplicate, with standardized sample preparation, within 30 476 working days. In addition, our data have 0.1% amount of missing values at protein level 477 owing to the data acquisition strategy and improvements in bioinformatics analysis. This 478 study demonstrates that the human proteotype can be obtained with a throughput comparable 479 to genomic and transcriptomic analyses, though still at relatively lower coverage. 480

481

Two aspects of our workflow ensure robust and quantitatively accurate protein 482 483 expression measurements. First, we obtained technical duplicates for the entire set of NCI-60 proteotypes, which was feasible due to the unparalleled high sample-throughput of the PCT-484 485 SWATH methodology which is now gaining popularity in proteomic profiling of clinical specimens. In addition, we developed an expert system software (manuscript in preparation) 486 to further curate peptide and protein identification and quantification. Applying stringent 487 criteria, 3,171 proteins were included for further analyses. The raw MS signal for each of 488 these quantified proteins, in each cell line, was inspected by the expert system, simulating 489 manual inspection, and is available for visual inspection in the supplementary data. We further 490 compared the expression of a few proteins with known expression in certain cell lines, 491 obtaining good agreement. Nevertheless, we cannot conclude that the peptides and proteins 492 that failed to pass curation by the expert system are not biological signals, due to the 493 unpredictable degree of biological heterogeneity, and the fact that we did not analyze non-494 canonical peptide variants and post-translational modification. The latter can be potentially 495 496 dissected and quantitated by future in silico analyses of our SWATH maps. Since the NCI-60 cell lines are widely used in cell biology, we anticipate broad utility of this highly curated 497 proteomic data. Additionally, our rapid proteotype acquisition pipeline using PCT-SWATH 498 requires little biological material, making it suitable for clinical settings and in precision 499 medicine efforts 7, 8, 52. 500

Compared to other omics data, the proteotypes obtained here offered unique insights 502 into the coordinated expression of protein complexes. Interactions amongst their component 503 subunits contribute to our understanding of protein function, as well as human diseases ^{24, 53-} 504 ⁵⁵. Several protein complexes have been identified as biomarkers of disease, including cancer 505 progression ⁵⁶. Our high quality proteomic data allowed systematic investigation of the 506 composition of 101 protein complexes in 60 cell lines. We expect that this represents a proof-507 of-principle for a generic, high-throughput approach, applicable to clinical specimens⁷, for 508 exploring the association between protein complexes and biological/disease phenotypes. 509

510

511 The NCI-60 continues to enable important contributions that have come and continue to come from this resource, and often emerging technologies are first tested on this cell line 512 panel due to its diversity and depth of surrounding knowledge ^{3, 12, 57-59}. Each cancer cell line 513 514 in the NCI-60 has been tested against tens of thousands of compounds, including the 240 FDA-approved and investigational drugs featured in our analyses. With the addition of the 515 516 SWATH proteomic data, the NCI-60 remains positioned as one of most comprehensive models for cancer research and drug discovery ^{12, 15}. It uniquely enabled our thorough, 517 518 integrative analysis of different molecular profiles (genomic, transcriptomic, and proteomic) in predicting drug responsiveness. Our findings strengthen the body of work highlighting the 519 importance of integrative omic approaches in understanding drug mechanisms and establish 520 the benefit of large-scale proteomic measurements. Therefore, we expect this work to become 521 a seminal work in the area of pharmacoproteomics, the benefit of which will grow with 522 anticipated expansion of sample size, proteomic coverage, including extension to 523 phosphoproteomic expression, as well as extension to mouse models ⁶⁰ and human specimens 524 7 525

526

The existing SWATH data specifically enabled the use of advanced analysis 527 techniques to produce multivariate models of drug response. Great effort was put into making 528 529 our work accessible to a large audience through data submission to the NCI-60 CellMiner database and availability through an accompanying R package, rcellminer. We expect this 530 531 pipeline based on the widely used elastic net method will continue to evolve and enable future 532 studies on additional data sets and phenotypes. And while the strengths of the elastic net method over other related methods have been previously described ^{61, 62}, the resulting models 533 still require careful scrutiny by individual researchers. The interpretation of the models 534 535 developed here, and by others using our pipeline, should be guided by understanding of the

biological activities of the associated predictors in the context of the mechanisms of action for 536 the input drugs. From the models generated by the current analyses, we identified several 537 potential determinants of drug responses, including NUDT5 and MAT2B protein levels for 538 the antimetabolite 6-TG, as well as complementary markers, such as LAMTOR3 protein 539 levels in conjunction with BRAF mutational status for Vemurafenib and other BRAF 540 inhibitors. These determinants may provide clinically relevant insights toward understanding 541 mechanisms of resistance to these and other agents. Together, these results invite further 542 investigation of this unique proteomic data resource. For example, in the current study's 543 544 analysis of protein complexes, we identified discrepancies between data at the transcriptomic and proteomic levels. This observation has been similarly made in tumor samples, with 545 additional variation across tissue types ⁶³. These differences can be used in future studies to 546 develop drug response models with non-redundant predictor sets including both data types. 547 548 However, due to the tissue diversity of the NCI-60 cells and the limited number of cell lines, data from more cancer cell lines of specific tissue type and extension to clinical specimens are 549 550 required to advance our findings to clinical applications.

551

552 Acknowledgements

We thank Margot Sunshine who developed CellMiner and the NCI-DTP team (Dr. 553 Jerry Collins and Dr. James H. Doroshow) for the drug data and support for data acquisition, 554 Emanuel Goncalves for comments to the manuscript. The work was supported by the 555 SystemsX.ch project PhosphoNetX PPM (to R.A.), the Swiss National Science Foundation 556 (grant no. 3100A0-688 107679 to R.A.), the European Research Council (grants no. ERC-557 558 2008-AdG 233226 and ERC-20140AdG 670821 to R.A.), European Union's Horizon 2020 research and innovation programme under grant agreement No 668858 (to R.A., J.S.-R., L.C., 559 P.W.), the Ruth L. Kirschstein National Research Service Award (grant no. F32 CA192901 to 560 A.L.), the National Resource for Network Biology (NRNB) from the National Institute of 561 562 General Medical Sciences (NIGMS) (grant no. P41 GM103504 to C.S.), and the Center for 563 Cancer Research, Intramural Program of the National Cancer Institute (grant no. Z01 BC006150 to Y.P), and the Wellcome Trust Award (102696) to M.J.G.. We thank An Guo for 564 565 help in preparing the graphics.

566

567 Author contributions

T.G. designed and coordinated the project with supervision from R.A. C.C.K.
processed the samples. L.G., C.C.K. and T.G. acquired the SWATH data. T.G. performed the

- 570 SWATH data interpretation and benchmarking with help from C.C.K., and the expert system
- analysis with help from U.S. A.L., V.N.R. and Z.W. performed the drug response prediction
- analysis, and developed the reproducible research infrastructure, with critical inputs from
- 573 M.P.M., J.S.R., M.J.G., S.V., W.C.R., C.S, and Y.P., L.C. and L.M. performed the pathway
- analysis. A.L., V.N.R., W.C.R. and S.V. integrated the SWATH data into rcellminer and
- 575 CellMiner. A.O., M.I. and R.C. performed the protein complex analysis, with help from A.L.,
- 576 Z.W., Y.C., V.N.R, C.S., Y.S., Y.Z., Y.P. P.Q. and Q.Z. contributed to the data analysis.
- 577 T.G., A.L. and V.N.R. wrote the manuscript with inputs from all co-authors. P.J.W., P.B.,
- 578 M.R., J.S.R., W.C.R., C.S., Y.P. and R.A. supervised the project.
- 579

580 **Competing financial interests**

- 581 R.A. holds shares of Biognosys AG, which operates in the field covered by the article.
- 582 The research group of R.A. is supported by SCIEX, which provides access to prototype
- 583 instrumentation, and Pressure Biosciences, which provides access to advanced sample
- 584 preparation instrumentation.
- 585
- 586

587 Materials and Methods

588

589 PCT-assisted sample preparation for MS analyses

590

The NCI-60 cells were obtained as frozen, non-viable cell pellets from the 591 Developmental Therapeutics Program (DTP), National Cancer Institute (NCI-NIH) and 592 processed using Barocycler® NEP2320 (PressureBioSciences Inc, South Easton, MA). The 593 IDs of the NCI-60 cells in our study matching to the IDs in Cellminer and a previous 594 proteomic study by the Kuster group are provided in Supplementary Table 1. Briefly, cell 595 pellets were lysed in a buffer containing 8M urea, 0.1M ammonium bicarbonate, and 596 CompleteTM protease inhibitor using barocycler program (20 seconds 45 kpsi, 10 seconds 0 597 kpsi, 120 cycles) at 35°C⁷. Whole cell lysates were sonicated for 25 seconds with 1 min 598 interval on ice for 3 times. Cellular debris was removed by centrifugation and sample protein 599 600 concentration was determined by BCA assay prior to protein reduction with 10 mM TCEP for 20 min at 35°C, and alkylation with 40 mM iodoacetamide in the dark for 30 min at room 601 602 temperature. Lys-C digestion (1/50, w/w) was performed in 6 M urea using PCT program (25 seconds 25 kpsi, 10 seconds 0 kpsi 75 cycles) at 35°C; whereas trypsin digestion (1/30, w/w) 603 604 was performed in further diluted urea (1.6M) using PCT program (25 seconds 25 kpsi, 10 seconds 0 kpsi, 160 cycles) at 35°C. Digestion was stopped by acidification with 605 606 trifluoroacetic acid to a final pH of around 2 before C18 column desalting using SEP-PAK 607 C18 cartridges (Waters Corp., Milford, MA, USA).

608

609 Off-gel electrophoresis

610

To create a comprehensive spectral library for SWATH-MS analysis, we pooled 20-611 40% of desalted peptide solutions from each NCI-60 sample and performed off-gel 612 fractionation. Briefly, pooled peptides were resolubilised in OGE buffer containing 5% (v/v)613 glycerol, 0.7% (v/v) acetonitrile (ACN) and 1% (v/v) carrier ampholytes mixture (IPG buffer 614 pH 3.0-10.0, GE Healthcare). Fractionation was performed on a 3100 OFFGEL (OGE) 615 Fractionator (Agilent Technologies) using a 24 cm pH3-10 IPG strip (Immobilised pH 616 617 Gradient strip from GE Healthcare) according to manufacturer's instructions using a program of 1 h rehydration at a maximum of 500 V, 50 µA and 200 mW followed by separation at a 618 maximum of 8000 V, 100 µA and 300 mW until 50 kVh were reached. Each of 24 fraction 619 was recovered and cleaned up by C18 reversed-phase MicroSpin columns (The Nest Group 620

Inc.). Based on the sample complexity (based on Nanodrop, A280 measurement), for each

- strip, the following fractions were pooled into 12 samples for MS injections: pool 1 (fraction
- 623 1-2), pool 2 (fraction 3), pool 3 (fraction 4), pool 4 (fraction 5), pool 5 (fraction 6-7), pool 6
- 624 (fraction 8-9), pool 7 (fraction 10-11), pool 8 (fraction 12-15), pool 9 (fraction 16-19), pool 10
- 625 (fraction 20-21), pool 11 (fraction 22), pool 12 (fraction 23-24). Those were injected in
- 626 quadruplicate, resulting in 48 DDA injections of fractionated samples.
- 627

628 DDA MS for spectral library generation

629

For spectral library generation, a SCIEX TripleTOF 5600 System mass spectrometer 630 was operated essentially as described before ⁶⁴: all samples were analyzed on an Eksigent 631 nanoLC (AS-2/1Dplus or AS-2/2Dplus) system coupled with a SWATH-MS-enabled AB 632 633 SCIEX TripleTOF 5600 System. The HPLC solvent system consisted of buffer A (2% ACN and 0.1% formic acid, v/v) and buffer B (95% ACN with 0.1% formic acid, v/v). Samples 634 635 were separated in a 75 µm diameter PicoTip emitter (New Objective) packed with 20 cm of Magic 3 µm, 200A C18 AQ material (Bischoff Chromatography). The loaded material was 636 eluted from the column at a flow rate of 300 nL min⁻¹ with the following gradient: linear 2 -637 35% B over 120 min, linear 35 - 90% B for 1 min, isocratic 90% B for 4 min, linear 90 - 2% 638 B for 1 min and isocratic 2% solvent B for 9 min. The mass spectrometer was operated in 639 DDA mode using a top20 method, with 500 ms and 150 ms acquisition time for the MS1 and 640 MS2 scans respectively, and 20 s dynamic exclusion for the fragmented precursors. Rolling 641 collision energy using the following equation $(0.0625 \times m/z - 3.5)$ with a collision energy 642 spread of 15 eV was used for fragmentation regardless of the charge state of the precursors, to 643 mimic as close as possible the fragmentation conditions of the precursors in SWATH-MS 644 mode. Altogether, we had 66 DDA-MS injections, including the 48 OGE samples and another 645 18 pooled peptide samples from the unfractionated cell lysate of the NCI-60 cells. 646

647

648 Spectral and assay library generation

649

All raw instrument data were centroided using Proteowizard msconvert (version 2.0).

651 The assay library was generated using an established protocol ^{<math>64}. In short, the shotgun data

sets were searched individually using X!Tandem 65 (2011.12.01.1) with k-score plugin 66 ,

653 Myrimatch ⁶⁷ (2.1.138), OMSSA ⁶⁸ (2.1.8) and Comet ⁶⁹ (2013.02r2) against the reviewed

654 UniProtKB/Swiss-Prot (2014_02) protein sequence database containing 20,270 proteins

appended with 11 iRT peptides and decoy sequences. Carbamidomethyl was used as a fixed 655 modification and oxidation as the variable modification. Maximally two missed cleavages 656 were allowed. Peptide mass tolerance was set to 50 ppm, fragment mass error to 0.1 Da. The 657 search identifications were combined and statistically scored using PeptideProphet⁷⁰ and 658 iProphet⁷¹ available within the Trans-Proteomics Pipeline (TPP) toolset (version 4.7.0)⁷². 659 MAYU⁷³ (v. 1.07) was used to determine the iProphet cutoff (0.999354) corresponding to a 660 protein FDR of 1.03%. SpectraST was used in library generation mode with CID-QTOF 661 settings and iRT normal-isation at import against the iRT Kit ⁷⁴ peptide sequences (-662 c_IRTirt.txt -c_IRR) and a consensus library was consecutively generated. An in-house 663 python script, spec-trast2tsv.py31 (msproteomicstools 0.2.2) was then used to generate the 664 assay library with the following settings: -1 350,2000 -s b,y -x 1,2 -o 6 -n 6 -p 0.05 -d -e -w 665 swath32.txt -k openswath (fragment ions between 350 and 2000 m/z, b and y ions authorized, 666 667 fragment charges 1+ and 2+, 6 most intense transitions, precision of fragment ion retrieved 0.05 Da, exact fragment ion mass calculated, exclude fragments in the swath window). The 668 669 OpenSWATH tool, ConvertTSVToTraML converted the TSV file to TraML format; Open-SwathDecoyGenerator generated the decoy assays in shuffle mode and appended them to the 670 671 TraML assay library. In this study, we built a SWATH assay library containing 86,209 proteotypic peptide precursors in 8,056 proteotypic SwissProt proteins. This library is 672 supplied in PRIDE project PXD003539. 673

674

675 SWATH-MS

676

The SWATH-MS data acquisition in a Sciex TripleTOF 5600 mass spectrometer was performed as described before ¹⁰, using 32 windows of 25 Da effective isolation width (with an additional 1 Da overlap on the left side of the window) and with a dwell time of 100 ms to cover the mass range of 400 - 1200 m/z in 3.3 s. The collision energy for each window was set using the collision energy of a 2+ ion centered in the middle of the window (equation: 0.0625 x m/z - 3.5) with a spread of 15 eV. The sequential precursor isolation window setup was as follows: [400-425], [424-450], [449-475], ..., [1174-1200].

684

685 Protein identification using OpenSWATH

686

We analyzed the SWATH data using OpenSWATH software ¹¹ using parameters as
 described previously ²⁴. We identified 48,374 peptides from 6,556 protein groups from the

689 NCI-60 panel with < 1% false discovery rate at both peptide and protein level evaluated by 690 OpenSWATH ¹¹and Mayu ⁷⁵ (supplied in PRIDE project PXD003539).

691

692 **DIA-expert analyses**

693

The DIA-expert software read OpenSWATH output result file which contains 694 statistical scores (i.e. mProphet score or mScore) indicating the confidence of identification 695 for each peptide precursor in each sample, and from there selected the sample in which a 696 697 peptide precursor was identified with highest confidence. It then obtained extracted ion chromatograms (XICs) for the target peptide precursor and all associated annotated b and y 698 fragments in the reference sample, and refined fragments based on the peak shape of each 699 700 fragment and its peak boundary. The refined fragments and precursor XIC traces from each of 701 the rest samples were subsequently compared with the reference peak group using empirical expert rules, based on which the best matched peak group in each sample was picked and 702 703 visualized. Duplicated measurements were used to evaluate the accuracy of peptide and protein quantification. The protein quantity was normalized based on total ion 704 705 chromatography of the MS1 spectra from each raw SWATH file. All codes are provided in 706 Github https://github.com/tiannanguo/dia-expert.

707

708 Protein complexes analysis

709

For this analysis, technical replicates were averaged to generate the NCI-60 710 proteotypes. To assess the coverage of protein complexes by NCI-60 proteotypes, we 711 retrieved a large resource of mammalian protein complexes assembled from CORUM ⁷⁶, 712 COMPLEAT ⁷⁷ and literature-curated complexes ^{24, 78}. This resource contains 2,041 proteins 713 as members of 279 distinct complexes and it is available at http://variablecomplexes.embl.de/. 714 101 complexes were represented in the NCI-60 proteotypes with at least 5 members 715 716 quantified. These complexes, in total, contain 1,045 distinct proteins quantified in the NCI-60 proteotypes. Pearson's correlation coefficient was calculated for all the pairwise comparisons 717 718 of 3,171 proteins across the NCI-60 cell lines. All pairwise comparisons were classified into two categories: either two proteins were members of the same complex or not. Average 719 720 abundance, standard deviation and average Pearson correlation of each complex were calculated based on the abundance of complex members in the NCI-60 proteotypes. 721 722

For this analysis, technical replicates were averaged to generate the NCI-60 723 724 proteotypes. To assess the coverage of protein complexes by NCI-60 proteotypes, we retrieved a large resource of mammalian protein complexes assembled from CORUM ⁷⁶, 725 COMPLEAT ⁷⁷ and literature-curated complexes ^{24, 78}. This resource contains 2041 proteins 726 as members of 279 distinct complexes and it is available at http://variablecomplexes.embl.de/. 727 728 158 complexes were represented in the NCI-60 proteotypes with at least 5 members quantified. These complexes, in total, contain 1,045 distinct proteins quantified in the NCI-60 729 proteotypes. Pearson's correlation coefficient was calculated for all the pairwise comparisons 730 731 of 3,171 proteins across the NCI-60 cell lines. All pairwise comparisons were classified into 732 two categories: either two proteins were members of the same complex or not. Average 733 abundance and standard deviation of each complex were calculated based on the mean 734 abundance of complex members in the NCI-60 proteotypes. 735

736 737

The activity of pathways, as they are described in ACSN, has been computed using 738 739 ROMA ³². Among all the modules defined in ACSN, only 11 show a significant dispersion 740 over the data set: AKT_MTOR, HR (Homologous Recombination), NER (nucleotide Excision Repair), TNF response, Death Receptors regulators, Apoptosis, caspases, E2F3 and 741 E2F4 targets, HIF1 and cytoskeleton polarity. For these modules, the mean activity score for 742 each type of cancer cell lines was computed and mapped onto the atlas (from bright green for 743 low values to bright red for high values). To assess module differential activity between 744 745 proteotypes, we computed a *t*-test on the activity scores in cell lines of a cancer type versus 746 the activity of all other cancer cell lines. The definition of genes composing each module can 747 be found in http://acsn.curie.fr

- 748
- 749

750 Drug sensitivity prediction using elastic net

Pathway activity analysis

751

The elastic net regularized regression algorithm was applied to predict drug response for 240 FDA-approved or investigational NSC-designated compounds. Some widely studied drugs are represented by more than one NSC identifier, with each identifier associated with a distinct compound sample and series of NCI-60 drug activity assays. For each compound, 7 combinations of input data were evaluated. These included NCI-60 mRNA expression, gene-

level mutation, and SWATH-MS protein expression, both alone and in all possible 757 combinations. mRNA expression data was available for 14,969 genes, and derived from 758 759 CellMiner, with missing values imputed using the impute.knn function (with default parameters) of the Bioconductor impute package. Gene-level mutation profiles were 760 available for 2,282 genes, and were obtained from CellMiner exome sequencing data, with 761 762 values indicating the percent conversion to a variant form for the case of expected functionimpacting alterations (frameshift, nonsense, splice-sense, missense mutations by 763 SIFT/PolyPhen2 analysis). SWATH-MS based protein expression data was available for 764 765 3,171 proteins.

766

Elastic net analysis was done using the glmnet R package ⁷⁹. The elastic net analysis was conducted using a multi-step pipeline involving cross-validations performed in a nested manner. The "outer" cross-validation is a leave-one-out cross validation that is conducted over all computational steps present in the "inner" pipeline, and it is used to validate model performance. The "inner" cross-validation are conducted to select elastic net hyperparameters (alpha and lambda) and for predictor set trimming, using data from a set of ~59 cell lines.

774 The elastic net parameters alpha and lambda were selected by minimizing the crossvalidation error (average of 10 replicates of 10-fold cross-validation) within the "inner" 775 776 pipeline. The selected alpha and lambda parameters were then applied to 200 runs of the 777 elastic net algorithm, each using a random data subset derived from 90% of the available cell lines. The 200 resulting coefficient vectors were then averaged, and predictors were ranked by 778 the magnitude of their average coefficient weight. To select a limited number of predictors 779 with potential to generalize to new data, top k-element predictor sets (by average coefficient 780 weight magnitude) were evaluated using standard linear regression and 10-fold cross-781 validation. The appropriate k was set to the smallest value yielding a cross-validation error 782 within one standard deviation of the minimum cross-validation error. 783

784

To obtain a robust estimate of performance on unseen data, leave-one-out crossvalidation was applied to the overall procedure as part of the "outer" pipeline. Specifically, drug response for each cell line was predicted using an elastic net model derived using the remaining held out data (and the steps outlined above). The vector of predicted response values was then correlated with the actual response values, with the Pearson's correlation

- coefficient providing an estimate of the predictive value of the applied input data
- combination. More details of the elastic net algorithm are provided in File S3.
- 792

793 Elastic net analysis was done using the rcellminerElasticNet R package

794 (<u>https://bitbucket.org/cbio_mskcc/rcellminerelasticnet</u>), which facilitates the application of the

glmnet R package (which provides the elastic net algorithm code) to data from the rcellminer

- and rcellminerData packages ⁸⁰. rcellminerElasticNet also provides utility functions for
- summarizing and visualizing elastic net results.
- 798

799

Results for the elastic net analysis are available from this URL:

800 https://discover.nci.nih.gov/cellminerreviewdata/swath_analysis/swathOutput_062316_all.tar.

gz. This compressed file contains results for the analysis run with all features and selected

802 common features. Each drug compound has three files for each combination of molecular

features used in a particular run of the elastic net algorithm: 1) a knitr report R Markdown

804 (.Rmd) file containing the code that was run, 2) an RData (.Rdata) file containing the results

of each elastic net run (see elasticNet() documentation in the rcellminerElasticNet package),

- 3) the rendered knitr report as a webpage (.html).
- 807

Beyond the knitr report containing code, the elastic net pipeline is made reproducible using a Docker image. Docker (www.docker.com) is an emerging platform for conducting reproducible research in the biomedical research community. All necessary software and dependencies to run the described analysis have been embedded in the available Docker container to provide readers an environment that runs on all major operating systems (including Windows, OSX, and Linux), making Docker containers self-contained, portable, and capable of performing at levels similar to the host system.

815

The Docker container is available at the Docker Hub repository: cannin/swath (https://hub.docker.com/r/cannin/swath/). Key dependencies installed, include: RStudio Server (https://www.rstudio.com/), rcellminer/rcellminerData ⁸⁰, and rcellminerElasticNet. With these installed dependencies, readers have the opportunity to 1) re-run analysis for specific drug compounds and modify the code in order to extend the analysis using RStudio Server, a web-based version of the RStudio R editor, and 2) use an R Shiny app web-based data explorer to further understand described results. Instructions on the usage of the Docker

823	container are located at the rcellminerElasticNet project page
824	(https://bitbucket.org/cbio_mskcc/rcellminerelasticnet).
825	
826	Data deposition
827	
828	The NCI-60 SWATH data sets and SWATH assay library has been deposited in
829	PRIDE. Project Name: NCI60 proteome by PCT-SWATH; Project accession: PXD003539.
830	Reviewer account details:
831	Username: reviewer15254@ebi.ac.uk
832	Password: dWdyptzf
833	The protein data matrix has also been deposited in ArrayExpress. Project accession: E-
834	PROT-2. Project title: Proteomic profiling of NCI60 cell lines from Cancer Cell Line
835	Encyclopedia.
836	Reviewer account details:
837	Username: Reviewer_E-PROT-2
838	Password: gdgywGco
839	The protein data matrix is also accessible in CellMiner website ¹³ and R package
840	rcellminer ³⁷ .

Cancer Genome Atlas Research, N. et al. The Cancer Genome Atlas Pan-Cancer analysis

842 **References**

1.

845		project. <i>Nat Genet</i> 45 , 1113-1120 (2013).
846	2.	Barretina, J. et al. The Cancer Cell Line Encyclopedia enables predictive modelling of
847		anticancer drug sensitivity. Nature 483, 603-607 (2012).
848	3.	Garnett, M.J. et al. Systematic identification of genomic markers of drug sensitivity in cancer
849		cells. Nature 483 , 570-575 (2012).
850	4.	Zhang, B. et al. Proteogenomic characterization of human colon and rectal cancer. Nature
851		513 , 382-387 (2014).
852	5.	Mertins, P. et al. Proteogenomics connects somatic mutations to signalling in breast cancer.
853		Nature 534 , 55-62 (2016).
854	6.	Zhang, H. et al. Integrated Proteogenomic Characterization of Human High-Grade Serous
855		Ovarian Cancer. Cell (2016).
856	7.	Guo, T. et al. Rapid mass spectrometric conversion of tissue biopsy samples into permanent
857		quantitative digital proteome maps. <i>Nat Med</i> (2015).
858	8.	Shao, S. et al. Minimal sample requirement for highly multiplexed protein quantification in
859		cell lines and tissues by PCT-SWATH mass spectrometry. Proteomics (2015).
860	9.	Powell, B.S., Lazarev, A.V., Carlson, G., Ivanov, A.R. & Rozak, D.A. Pressure cycling technology
861		in systems biology. Methods Mol Biol 881, 27-62 (2012).
862	10.	Gillet, L.C. et al. Targeted data extraction of the MS/MS spectra generated by data-
863		independent acquisition: a new concept for consistent and accurate proteome analysis.
864		Molecular & cellular proteomics : MCP 11, 0111 016717 (2012).
865	11.	Rost, H.L. et al. OpenSWATH enables automated, targeted analysis of data-independent
866		acquisition MS data. Nat Biotechnol 32, 219-223 (2014).
867	12.	Shoemaker, R.H. The NCI60 human tumour cell line anticancer drug screen. Nat Rev Cancer
868		6 , 813-823 (2006).
869	13.	Reinhold, W.C. et al. CellMiner: A Web-Based Suite of Genomic and Pharmacologic Tools to
870		Explore Transcript and Drug Patterns in the NCI-60 Cell Line Set. Cancer Res 72, 3499-3511
871		(2012).
872	14.	Fojo, T. et al. Identification of non-cross-resistant platinum compounds with novel
873		cytotoxicity profiles using the NCI anticancer drug screen and clustered image map
874		visualizations. Crit Rev Oncol Hematol 53, 25-34 (2005).
875	15.	Holbeck, S.L., Collins, J.M. & Doroshow, J.H. Analysis of Food and Drug Administration-
876		approved anticancer agents in the NCI60 panel of human tumor cell lines. <i>Mol Cancer Ther</i> 9,
877		1451-1460 (2010).
878	16.	Bates, S.E. et al. Romidepsin in peripheral and cutaneous T-cell lymphoma: mechanistic
879		implications from clinical and correlative data. <i>Br J Haematol</i> 170 , 96-109 (2015).
880	17.	Gholami, A.M. et al. Global Proteome Analysis of the NCI-60 Cell Line Panel. <i>Cell Rep</i> 4 , 609-
881		620 (2013).
882	18.	Picotti, P. & Aebersold, R. Selected reaction monitoring-based proteomics: workflows,
883		potential, pitfalls and future directions. <i>Nat Methods</i> 9 , 555-566 (2012).
884	19.	Law, V. et al. DrugBank 4.0: shedding new light on drug metabolism. <i>Nucleic Acids Res</i> 42 ,
885		D1091-1097 (2014).
886	20.	Uhlen, M. et al. Proteomics. Tissue-based map of the human proteome. Science 347 ,
887	24	1260419 (2015).
888	21.	Lin, J.L. et al. MST4, a new Ste20-related kinase that mediates cell growth and transformation
889	22	via modulating ERK pathway. <i>Uncogene</i> 20 , 6559-6569 (2001).
890	22.	Huang, C.L., Cha, S.K., Wang, H.K., Xie, J. & Cobb, M.H. WNKs: protein kinases with a unique
891		kinase domain. <i>Exp Mol Med</i> 39 , 565-573 (2007).

892 893 894	23.	Xu, H. et al. Epidermal growth factor receptor (EGFR)-related protein inhibits multiple members of the EGFR family in colon and breast cancer cells. <i>Mol Cancer Ther</i> 4 , 435-442 (2005).
895 896	24.	Ori, A. et al. Spatiotemporal variation of mammalian protein complex stoichiometries. <i>Genome Biol</i> 17 , 47 (2016).
897 898	25.	Kanungo, J. Exogenously expressed human Ku70 stabilizes Ku80 in Xenopus oocytes and induces heterologous DNA-PK catalytic activity. <i>Mol Cell Biochem</i> 338 , 291-298 (2010).
899 900 901	26.	Chang, H.W. et al. Effect of beta-catenin silencing in overcoming radioresistance of head and neck cancer cells by antagonizing the effects of AMPK on Ku70/Ku80. <i>Head Neck</i> 38 Suppl 1 , E1909-1917 (2016)
902 903	27.	Feng, L. & Chen, J. The E3 ligase RNF8 regulates KU80 removal and NHEJ repair. <i>Nat Struct</i> Mol Biol 19 , 201-206 (2012).
904 005	28.	Kuperstein, I. et al. Atlas of Cancer Signalling Network: a systems biology resource for
905 906 907	29.	Hanahan, D. & Weinberg, R.A. Hallmarks of cancer: the next generation. <i>Cell</i> 144 , 646-674 (2011).
908 909	30.	Mackay, H.J. & Twelves, C.J. Targeting the protein kinase C family: are we there yet? <i>Nat Rev Cancer</i> 7 , 554-562 (2007).
910 911	31.	Engers, R. et al. Protein kinase C in human renal cell carcinomas: role in invasion and differential isoenzyme expression. <i>Br J Cancer</i> 82 , 1063-1069 (2000).
912 913	32.	Martignetti, L., Calzone, L., Bonnet, E., Barillot, E. & Zinovyev, A. ROMA: Representation and Quantification of Module Activity from Target Expression Data. <i>Front Genet</i> 7 , 18 (2016)
914	33.	Ummanni, R. et al. Peroxiredoxins 3 and 4 are overexpressed in prostate cancer tissue and
915 916 917	34.	affect the proliferation of prostate cancer cells in vitro. <i>J Proteome Res</i> 11 , 2452-2466 (2012). Shankavaram, U.T. et al. CellMiner: a relational database and query tool for the NCI-60 cancer cell lines. <i>BMC genomics</i> 10 , 277 (2009)
918 919	35.	Jones, P. et al. PRIDE: a public repository of protein and peptide identifications for the proteomics community. <i>Nucleic Acids Res</i> 34 , D659-663 (2006).
920 921	36.	Brazma, A. et al. ArrayExpressa public repository for microarray gene expression data at the EBL Nucleic Acids Res 31 , 68-71 (2003)
922	37.	Luna, A. et al. rcellminer: exploring molecular profiles and drug response of the NCI-60 cell
923 924	38.	Zoppoli, G. et al. Putative DNA/RNA helicase Schlafen-11 (SLFN11) sensitizes cancer cells to
925 926	39.	DNA-damaging agents. <i>Proc Natl Acad Sci U S A</i> 109 , 15030-15035 (2012). Solit, D.B. et al. BRAF mutation predicts sensitivity to MEK inhibition. <i>Nature</i> 439 , 358-362
927 928	40	(2006). de Boer NK, van Bodegraven AA, Ibaran B, de Graaf P, & Mulder C I, Drug Insight:
929 930	40.	pharmacology and toxicity of thiopurine therapy in patients with IBD. <i>Nat Clin Pract</i> <i>Gastroenterol Hepatol</i> 4 , 686-694 (2007).
931 932	41.	Doench, J.G. et al. Optimized sgRNA design to maximize activity and minimize off-target effects of CRISPR-Cas9. <i>Nat Biotechnol</i> 34 , 184-191 (2016).
933 934 935	42.	Halim, A.B., LeGros, L., Geller, A. & Kotb, M. Expression and functional interaction of the catalytic and regulatory subunits of human methionine adenosyltransferase in mammalian cells. <i>The Journal of biological chemistry</i> 274 , 29720-29725 (1999).
936 937	43.	Flaherty, K.T. et al. Inhibition of mutated, activated BRAF in metastatic melanoma. <i>The New England journal of medicine</i> 363 , 809-819 (2010).
938 939	44.	Schaeffer, H.J. et al. MP1: a MEK binding partner that enhances enzymatic activation of the MAP kinase cascade. <i>Science</i> 281 , 1668-1671 (1998).
940 941	45.	Teis, D., Wunderlich, W. & Huber, L.A. Localization of the MP1-MAPK scaffold complex to endosomes is mediated by p14 and required for signal transduction. <i>Dev Cell</i> 3 , 803-814
942 943 944	46.	(2002). Jun, S. et al. PAF-mediated MAPK signaling hyperactivation via LAMTOR3 induces pancreatic tumorigenesis. <i>Cell Rep</i> 5 , 314-322 (2013).

945 946 947	47.	De Araujo, M.E. et al. Polymorphisms in the gene regions of the adaptor complex LAMTOR2/LAMTOR3 and their association with breast cancer risk. <i>PLoS One</i> 8 , e53768 (2013).
948	48.	Borst, P. & Elferink, R.O. Mammalian ABC transporters in health and disease. <i>Annu Rev</i>
949	40	Biochem 71, 557-552 (2002).
950 951 052	49.	characters through activation of Wnt/beta-catenin signaling. <i>J Exp Clin Cancer Res</i> 35 , 82
953 953	50.	Chen, S. et al. Wnt-1 signaling inhibits apoptosis by activating beta-catenin/T cell factor-
954		mediated transcription. J Cell Biol 152, 87-96 (2001).
955	51.	Liu, Y., Beyer, A. & Aebersold, R. On the Dependency of Cellular Protein Levels on mRNA
956		Abundance. <i>Cell</i> 165 , 535-550 (2016).
957	52.	Shao, S. et al. Reproducible Tissue Homogenization and Protein Extraction for Quantitative
958		Proteomics Using MicroPestle-Assisted Pressure-Cycling Technology. J Proteome Res 15,
959		1821-1829 (2016).
960	53.	Dudley, A.M., Janse, D.M., Tanay, A., Shamir, R. & Church, G.M. A global view of pleiotropy
961		and phenotypically derived gene function in yeast. <i>Mol Syst Biol</i> 1 , 2005 0001 (2005).
962	54.	Wang, Q. et al. Community of protein complexes impacts disease association. Eur J Hum
963		Genet 20 , 1162-1167 (2012).
964	55.	Fraser, H.B. & Plotkin, J.B. Using protein complexes to predict phenotypic effects of gene
965		mutation. Genome Biol 8, R252 (2007).
966	56.	Le, D.H. A novel method for identifying disease associated protein complexes based on
967		functional similarity protein complex networks. <i>Algorithms Mol Biol</i> 10 , 14 (2015).
968	57.	Barretina, J. et al. The Cancer Cell Line Encyclopedia enables predictive modelling of
969		anticancer drug sensitivity. <i>Nature</i> 483 , 603-607 (2012).
970	58.	Weinstein, J.N. Drug discovery: Cell lines battle cancer. <i>Nature</i> 483 . 544-545 (2012).
971	59.	Abaan, O.D. et al. The exomes of the NCI-60 panel: a genomic resource for cancer biology
972		and systems pharmacology. Cancer Res 73, 4372-4382 (2013).
973	60.	Gao, H. et al. High-throughput screening using patient-derived tumor xenografts to predict
974		clinical trial drug response. Nat Med 21 , 1318-1325 (2015).
975	61.	Papillon-Cavanagh, S. et al. Comparison and validation of genomic predictors for anticancer
976		drug sensitivity. J Am Med Inform Assoc 20, 597-602 (2013).
977	62.	Jang, I.S., Neto, E.C., Guinney, J., Friend, S.H. & Margolin, A.A. Systematic assessment of
978		analytical methods for drug sensitivity prediction from cancer cell line data. Pacific
979		Symposium on Biocomputing. Pacific Symposium on Biocomputing, 63-74 (2014).
980	63.	Kosti, I., Jain, N., Aran, D., Butte, A.J. & Sirota, M. Cross-tissue Analysis of Gene and Protein
981		Expression in Normal and Cancer Tissues. Sci Rep 6, 24799 (2016).
982	64.	Schubert, O.T. et al. Building high-quality assay libraries for targeted analysis of SWATH MS
983		data. Nat Protoc 10, 426-441 (2015).
984	65.	Craig, R. & Beavis, R.C. A method for reducing the time required to match protein sequences
985		with tandem mass spectra. Rapid Commun Mass Spectrom 17, 2310-2316 (2003).
986	66.	MacLean, B., Eng, J.K., Beavis, R.C. & McIntosh, M. General framework for developing and
987		evaluating database scoring algorithms using the TANDEM search engine. <i>Bioinformatics</i> 22 ,
988		2830-2832 (2006).
989	67.	Tabb, D.L., Fernando, C.G. & Chambers, M.C. MyriMatch: highly accurate tandem mass
990		spectral peptide identification by multivariate hypergeometric analysis. J Proteome Res 6,
991		654-661 (2007).
992	68.	Geer, L.Y. et al. Open mass spectrometry search algorithm. <i>J Proteome Res</i> 3 , 958-964 (2004).
993	69.	Eng, J.K., Jahan, T.A. & Hoopmann, M.R. Comet: an open-source MS/MS sequence database
994		search tool. Proteomics 13, 22-24 (2013).
995	70.	Keller, A., Nesvizhskii, A.I., Kolker, E. & Aebersold, R. Empirical statistical model to estimate
996		the accuracy of peptide identifications made by MS/MS and database search. Anal Chem 74 .
997		5383-5392 (2002).

998	71.	Shteynberg, D. et al. iProphet: multi-level integrative analysis of shotgun proteomic data
999		improves peptide and protein identification rates and error estimates. <i>Molecular & cellular</i>
1000		proteomics: mCP 10, M111 007690 (2011).
1001	72.	Keller, A., Eng, J., Zhang, N., Li, X.J. & Aebersold, R. A uniform proteomics MS/MS analysis
1002		platform utilizing open XML file formats. <i>Mol Syst Biol</i> 1 , 2005 0017 (2005).
1003	73.	Reiter, L. et al. Protein identification false discovery rates for very large proteomics data sets
1004		generated by tandem mass spectrometry. Molecular & cellular proteomics : MCP 8, 2405-
1005		2417 (2009).
1006	74.	Escher, C. et al. Using iRT, a normalized retention time for more targeted measurement of
1007		peptides. Proteomics 12, 1111-1121 (2012).
1008	75.	Reiter, L. et al. Protein Identification False Discovery Rates for Very Large Proteomics Data
1009		Sets Generated by Tandem Mass Spectrometry. Molecular & Cellular Proteomics 8, 2405-
1010		2417 (2009).
1011	76.	Ruepp, A. et al. CORUM: the comprehensive resource of mammalian protein complexes
1012		2009. Nucleic Acids Res 38 , D497-501 (2010).
1013	77.	Vinayagam, A. et al. Protein complex-based analysis framework for high-throughput data
1014		sets. <i>Sci Signal</i> 6 , rs5 (2013).
1015	78.	Ori, A. et al. Cell type-specific nuclear pores: a case in point for context-dependent
1016		stoichiometry of molecular machines. <i>Mol Syst Biol</i> 9 , 648 (2013).
1017	79.	Friedman, J., Hastie, T. & Tibshirani, R. Regularization Paths for Generalized Linear Models via
1018		Coordinate Descent. <i>J Stat Softw</i> 33 , 1-22 (2010).
1019	80.	Luna, A. et al. rcellminer: exploring molecular profiles and drug response of the NCI-60 cell
1020		lines in R. <i>Bioinformatics</i> 32 , 1272-1274 (2016).