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1 PLMD: an updated data resource of protein lysine modifications

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- 10 **Running title:** A protein lysine modification database

ABSTRACT

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Post-translational modifications (PTMs) occurring at protein lysine residues, or protein lysine modifications (PLMs), play critical roles in regulating biological processes. Due to the explosive expansion of the amount of PLM substrates and the discovery of novel PLM types, here we greatly updated our previous studies, and presented a much more integrative resource of protein lysine modification database (PLMD). In PLMD, we totally collected and integrated 284,780 modification events in 53,501 proteins across 176 eukaryotes and prokaryotes for up to 20 types of PLMs, including ubiquitination, acetylation, sumoylation, methylation, succinylation, malonylation, glutarylation, glycation, formylation, hydroxylation, butyrylation, neddylation, 2-hydroxyisobutyrylation. pupylation, propionylation, crotonylation, phosphoglycerylation, carboxylation, lipoylation and biotinylation. Using the data set, a motif-based analysis was performed for each PLM type, and the results demonstrated that different PLM types preferentially recognize distinct sequence motifs for the modifications. Moreover, various PLMs synergistically orchestrate specific cellular biological processes by mutual crosstalks with each other, and we totally found 65,297 PLM events involved in 90 types of PLM co-occurrences on the same lysine residues. Finally, various options were provided for accessing the data, while original references and other annotations were also present for each PLM substrate. Taken together, we anticipated the PLMD database can serve as a useful resource for further researches of PLMs. PLMD 3.0 was implemented in PHP + MySQL and freely available at http://plmd.biocuckoo.org.

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1. Introduction

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If not all, most of cellular functions are controlled by cell signaling pathways involving the proteins which are frequently modified by reversible post-translational modifications (PTMs) that dynamically coordinate the signaling networks (Mann and Jensen 2003; Mertins et al. 2013; Morris et al. 2015; Strzyz 2016). Among the major types of amino acids that can be modified, positively charged lysine residues play important roles in regulating protein functions, while the neutralization of the charge frequently brings enormous influences on the substrate proteins (Olsen et al. 2004). Accordingly, PTMs occurring at specific lysine residues in proteins, or protein lysine modifications (PLMs), play critical roles in regulating a broad spectrum of biological processes (Shaid et al. 2013; Choudhary et al. 2014; Huang et al. 2015; Hendriks and Vertegaal 2016). In general, specific lysine residues undergo different PLM processes (i) by addition of small molecule functional groups, which occurs in acetylation (Choudhary et al. 2014), methylation (Lanouette et al. 2014), succinylation (Zhang et al. 2011), malonylation (Xie et al. 2012), glutarylation (Tan et al. 2014), butyrylation (Chen et al. 2007), propionylation (Chen et al. 2007), crotonylation (Tan et al. 2011), and biotinylation (Sharp et al. 2006), (ii) by covalent linkage of some protein modifiers, which are critical in ubiquitination (Shaid et al. 2013), sumoylation (Lamoliatte et al. 2014), and pupylation (Poulsen et al. 2010) as well as NEDD8-mediated neddylation (Rabut and Peter 2008), or (iii) by non-enzymatic attachment of sugar molecules such as glycation (Goodman 2013). Recently, rapid progresses of proteomic methods (e.g., high-throughput liquid chromatography-mass spectrometry (LC-MS) techniques) and the introduction of pan-antibodies specific for distinct PLMs (Hendriks et al. 2014; Elia et al. 2015; Svinkina et al. 2015) have greatly promoted the identification of well-characterized PLMs as well as new ones. For example, using a monoclonal anti-Lys-ε-Gly-Gly (anti-diGly) antibody and a polyclonal acetyl-lysine antibody, Elia et al. (2015) identified and quantified 33,500 ubiquitination and 16,740 acetylation sites, respectively. Also, Svinkina et al. (2015) identified over 10,000 acetylation sites in Jurkat cells with a mixture of anti-lysine acetylation antibodies. In addition, applying the combined fractional diagonal chromatography technology, Stes et al. (2014) identified more than 7,500 endogenous ubiquitination sites in over 3,300 proteins. Hendriks et al. (2014) also profiled a global sumovlation in human cells with high-resolution MS in a

site-specific manner and totally identified over 4,300 sumoylation sites in over 1,600 proteins, 62 63 and Weinert et al. (2013) identified almost 8,000 succinylation sites from multiple species. In 64 particular, using the state-of-the-art proteomic techniques in combination with chemical 65 biology or biochemistry as validation tools, Dr. Yingming Zhao's group has identified a considerable number of novel PLMs such as succinylation, malonylation, propionylation 66 67 crotonylation, glutarylation, and 2-hydroxyisobutyrylation, which has greatly advanced the identification and functional investigation of these PLMs (Chen et al. 2007; Xie et al. 2012; 68 69 Dai et al. 2014; Tan et al. 2014). Since a flood of PLM sites have been uncovered, it has 70 emerged to be a great challenge for the collection and integration of bulky PLM substrates and 71 sites from different studies. Although several public databases, such as UniProt (Consortium 72 2014), dbPTM (Lu et al. 2012), PhosphositePlus (Hornbeck et al. 2015), HPRD (Prasad et al. 73 2009) and SysPTM (Li et al. 2014), also compiled PLM information, they mainly focused on a general purpose of the collection of PTMs, and only a limited part of the identified PLM 74 75 substrates and sites were included. Many newly identified PLM events still remain to be 76 integrated. In 2011, we developed a database of the compendium of protein lysine acetylation (CPLA) 77 78 by manually collecting 7151 known acetylation sites in 3311 proteins (Liu et al. 2011). Later, 79 we updated the CPLA 1.0 by extending acetylation to 12 types of PLMs, and renamed the 80 database as the compendium of protein lysine modifications (CPLM 2.0) (Liu et al. 2014). In 81 this study, we greatly improved our previous databases, and developed a much more 82 comprehensive data resource of protein lysine modification database (PLMD). Compared to 83 CPLA 1.0 and CPLM 2.0, PLMD 3.0 database has been greatly expanded in terms of 84 modification types, species numbers, protein numbers and total modification sites. It contained 284,780 modification events in 53,501 proteins from 176 species for 20 types of PLMs (Fig. 1). 85 Also, the detailed annotations of each protein entry together with the information of primary 86 references were provided. Based on the PLMD data set, a motif-based analysis of sequence 87 preferences was performed, and the most significantly over-represented sequence motif was 88 89 discovered around modification sites for 16 types of PLMs. Additionally, we detected 65,297 90 PLM events of 90 types of PLM co-occurrences on the same lysine residues, such as 24,487 91 acetylation-succinylation acetylation-ubiquitination sites, 11,056 5542 sites,

- acetylation-malonylation sites, 4033 ubiquitination-succinylation sites, 3363 ubiquitination-sumoylation sites and 1992 succinylation-malonylation sites, and the results demonstrated that different types of PLMs prefer to crosstalk with each other. Taken together, PLMD 3.0 can service as an informative platform for the community to access PLM information, and we anticipate it could be a useful resource for further experimental or
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2. Construction and content

- 99 We manually curated the searchable literature from PubMed to collect experimentally identified PLM substrates and sites by inputting keywords, including 'ubiquitination', 100 'acetylation', 'sumoylation', 'methylation', 'succinylation', 'malonylation', 'glutarylation', 101 102 'glycation', 'formylation', 'hydroxylation', 'butyrylation', 'propionylation', 'crotonylation', 103 'pupylation', 'neddylation', '2-hydroxyisobutyrylation', 'phosphoglycerylation', 104 'carboxylation', 'lipoylation' and 'biotinylation'. In order to provide a more comprehensive 105 data resource for researchers, additional keywords such as 'ubiquitinated', 'acetyl', 'acetylated', 106 'SUMO', 'succinyl' and other related nomenclatures were adopted for collecting more data. 107 All modified lysine residues were then mapped to the benchmark sequences retrieved from the 108 UniProt database. In addition, more detailed annotations such as protein names, gene names 109 and functional descriptions as well as sequence annotations of modified proteins were retrieved 110 from UniProt and further integrated into the PLMD database for providing rich information. 111 Also, primary references of PLM substrates and sites were offered to guarantee the reliability 112 and quality of the database. 113 Totally, we found 284,780 PLM events occurring at 234,062 lysine residues of 53,501
 - Totally, we found 284,780 PLM events occurring at 234,062 lysine residues of 53,501 proteins for 20 types of PLMs across 176 eukaryotes and prokaryotes (Table S1). Then we classified the PLMs into three categories: 1) four types of ubiquitin and ubiquitin-like modifications (Ub-UbLs), 2) nine types of acylations, and 3) seven types of other PLMs (Fig. 2). Among them, acylations and Ub-UbLs account for the vast majority of PLM events: the former possesses 141,276 (49.61%) acylation sites and the latter contains 130,194 (45.72%) sites. More specifically, ubiquitination (121,742 sites, 42.75%) and acetylation (111,253 sites, 39.07%), two extensively studied PLMs, still occupy a large proportion of all PLM sites with a growing number identified owing to their significant functional roles. However, it is

noteworthy that the amount of newly identified lysine acylations is dramatically increasing. For example, the succinylation and malonylation sites have reached up to 18,593 (6.53%) and 9,584 (3.37%), respectively. The rapid progress in the identification of these new lysine acylations is attributed to the advancement of proteomic techniques along with the emerging evidence that suggests that these new lysine acylations are important in regulating cellular metabolisms in both physiological and pathophysiological states (Hirschey and Zhao 2015; Xu et al. 2015; Sadhukhan et al. 2016). However, only a small number of substrates could be detected regarding other new lysine acylations such as butyrylation, crotonylation, glutarylation and propionylation, which are mainly identified on histones. Moreover, for Ub-UbLs, the number of identified sumoylation sites is also increased rapidly, especially in the last two years.

Although plenty of PLMs were experimentally discovered across 176 species from our data, the amount of identified substrates is generally limited in most organisms. The distribution of PLM substrates and sites in 20 major species with >600 substrates were analyzed. For most PLM types, a considerable number of substrates (Fig. 3A) and sites (Fig. 3B) were identified in mammals, especially in *Homo sapiens*, *Mus musculus* and *Rattus norvegicus*. There are also more than half of the PLM types found in the *Escherichia coli* and *Saccharomyces cerevisiae*. In addition, we observed that several types of Ub-UbLs were only exclusively identified in distinct species. For instance, ubiquitination, sumoylation and neddylation were only available in eukaryotes, while pupylation was only discovered in actinomyces.

3. Usage

Our database was developed in a user-friendly manner, and multiple options were provided for users to access the information of PLMs. Because of the variety of types and species of the modification data, two browse options including 'Browse by types' and 'Browse by species' were supplied in the database (Fig. 4). For convenience, PLMD allowed browsing 19 major species, while all the other species were denoted as 'Others'. Here we used lysine acetylation substrates from *H. sapiens* as an example to illustrate the usage of the browse options in PLMD. In the option of 'Browse by types', there were 20 simplified diagrams showing molecular structures of ligands conjugated to lysine residues, which were employed to

represent the 20 types of PLMs and were divided into three categories including Ub-UbLs, acylation and other PLMs (Fig. 4A). First, by clicking on the 'Acetylation' button, a brief description of protein lysine acetylation and the distribution of acetylated proteins of different species were displayed. Then the acetylation substrates in *H. sapiens* could be listed through clicking on the 'Homo sapiens' link (Fig. 4B). In the option of 'Browse by species', after clicking on the species diagram of *H. sapiens* (Fig. 4C), the distribution of lysine modified proteins of 20 types of PLMs in *H. sapiens* will be displayed, and then users can click on the link of 'Acetylation' to view the list of acetylated substrates in *H. sapiens*. The detailed information for any specified PLM protein could be accessed through the links in the list (Fig. 4D).

Besides above two options, PLMD also provided up to four implemented search options for users to query the database with one or multiple keywords including 'Substrate Search'

for users to query the database with one or multiple keywords including 'Substrate Search' (Fig. 5A), 'Advanced search' (Fig. 5B), 'Multiple Search' (Fig. 5C) and 'BLAST Search' (Fig. 5D). For the 'Substrate Search', users can input one or multiple keywords, e.g., using 'TP53' and selecting the 'Gene Name', the results will be shown in a tabular format with 'PLMD ID', 'UniProt Accession', 'Species' and 'Protein Name' (Fig. 5A). Furthermore, users can put up to three terms together by means of three operators of 'and', 'or' and 'exclude' to receive a more exhaustive acquirement via 'Advanced Search' (Fig. 5B). Also, 'Multiple Search' was present for retrieving multiple PLM proteins with a list of keywords (Fig. 5C). At last, with 'BLAST Search' option, users can find identical or homologous proteins with a protein sequence in the FASTA format by the application of NCBI BLAST packages (Fig. 5D) (Boratyn et al. 2013).

4. Discussion

The property of electron-rich and nucleophilic nature of the lysine chain makes it ideal for undergoing PTM reactions within multiple substrates (Shannon and Weerapana 2015), which could affect the protein stability and activity and further regulate various physiological and biological functions. Moreover, an increasing number of studies also suggested that aberrances of PLMs are highly related to a considerable number of diseases (Greer and Shi 2012; Flotho and Melchior 2013; Morris et al. 2015; Sadoul and Khochbin 2016). Recent advancement of proteomic techniques explosively expanded the number of PLM substrates and the discovery of novel PLMs. In view of the data accumulation, there is an urgent need to integrate these

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In this work, we updated the database of CPLM 2.0 into PLMD 3.0. The modification types, species numbers, protein numbers and total modification sites in PLMD 3.0 have been greatly expanded. Totally, it contained 284,780 PLM events in 53,501 substrates from 176 species for up to 20 types of PLMs, which then were classified into three categories including four types of Ub-UbLs, nine types of acylations and seven types of other PLMs.

In addition, PLM events occurring at specific positions of specific amino acids or peptides are catalyzed by a variety of corresponding enzymes (Xiong and Guan 2012; Deng et al. 2016; Xu et al. 2016), so the flanking sequences around the modification sites of the same PLM types may possess a strong sequence conservation due to the enzyme specificity. Therefore, using the data set in PLMD combined with Motif-X (Schwartz and Gygi 2005), we carried out a motif-based analysis of sequence preferences around modification sites for each PLM type. Sixteen types of PLMs were found to process the most significantly over-represented sequence motifs, including nine types of acylations, three types of Ub-UbLs and four types of other PLMs (Fig. 6A). Among them, the 2-hydroxyisobutyrylation was analyzed with P < 0.001 and glutarylation and phosphoglycerylation were analyzed with P < 0.0001 due to the data limitation, while other 13 types of PLMs were explored with P < 0.00001. For acylation motifs, the L and E residues were significantly enriched at -2 and -1 positions of the acetylation sequences, whereas N, K and R residues at +3, -4, and -1 positions are over-represented for lysine propionylation, butyrylation and 2-hydroxyisobutyrylation, respectively. Particularly, it was found that G residues were significantly enriched at -1 position for both succinylation and malonylation, suggesting that succinylation and malonylation incline to co-occur at the same position. For Ub-UbLs, LXXXKXL was discovered as the most predominant motif for ubiquitination. Meanwhile, KXE was found to be the over-represented motif for sumoylation, which was accordant with many reports that proteins conjugated by SUMO contain the basal KXE type motif (Geiss-Friedlander and Melchior 2007). We also found the EEXXK is the most remarkable motif for lysine methylation. Due to the data limitation, we didn't find any over-represented motifs for the remaining four PLM types, including neddylation, carboxylation, lipoylation and biotinylation. In addition, some of the modification sites in the PLMD were reported by only one literature, while some were reported by multiple evidences.

We checked modification events reported by multiple evidences and performed a motif analysis upon them (Fig. S1). From the results, for some PLMs, although the most significantly over-represented sequence motifs of identified sites reported multiple times are not fully consistent with the ones of all identified sites in the PLMD, those could be matched with the top five significantly over-represented sequence motifs of all identified PLM sites, which also indicated the reliability of our data sets.

Moreover, by competitively occurring at the same residue, multiple PLMs appear to act in combinatorial ways. From the collected data, we totally found 65,297 PLM events involved in 90 types of PLM co-occurrences. The distribution of co-occurrences of two different PLM types were pairwisely analyzed (Fig. 6B), and the detailed results were shown in Table S2. As the representatives of mostly abundant **PLM** crosstalks, there 11,056 are acetylation-succinylation, 5542 acetylation-malonylation and 1992 succinylation-malonylation sites in our results. Since lysine acylations depend on the similar acyl-CoA metabolic intermediates, such as acetyl-CoA (Ac-CoA), succinyl-CoA and malony-CoA (Hirschey and Zhao 2015), it can be expected that they are more likely to occur in the same locations. Furthermore, a comprehensive analysis of the crosstalks among different PLMs in same tissues or cell lines was performed (Fig.S2). The result showed that the co-occurrences of multiple PLMs on the same lysine residue significantly occurred in majority of tissues or cell lines. Moreover, the in situ crosstalks between different PLMs in same tissues or cell lines are consistent with that in all identified PLM sites. Since multiple PLMs are significantly co-occurred, the preference of the crosstalk among different PLMs and the properties on the PLM crosstalk need further explorations.

In the future, PLMD will be continuously maintained to expand with the increasing availability of data in various resources as well as enhancements on the text mining algorithm which will enable PubMed to pinpoint and select more PLM substrates and sites from research articles. We anticipated this updated database can provide a more useful resource for further computational or experimental studies.

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- 243 2014DFB30020).
- 244 References
- Boratyn, G.M., Camacho, C., Cooper, P.S., Coulouris, G., Fong, A., Ma, N., Madden, T.L.,
- 246 Matten, W.T., McGinnis, S.D., Merezhuk, Y., 2013. BLAST: a more efficient report with
- usability improvements. Nucleic Acids Res. 41, W29-W33.
- Chen, Y., Sprung, R., Tang, Y., Ball, H., Sangras, B., Kim, S.C., Falck, J.R., Peng, J., Gu, W.,
- 249 Zhao, Y., 2007. Lysine propionylation and butyrylation are novel post-translational
- 250 modifications in histones. Mol. Cell. Proteomics 6, 812-819.
- 251 Choudhary, C., Weinert, B.T., Nishida, Y., Verdin, E., Mann, M., 2014. The growing landscape
- of lysine acetylation links metabolism and cell signalling. Nat. Rev. Mol. Cell Bio. 15,
- 253 536-550.
- UniProt Consortium, 2015. UniProt: a hub for protein information. Nucleic Acids Res. 43,
- 255 D204-D212.
- Dai, L., Peng, C., Montellier, E., Lu, Z., Chen, Y., Ishii, H., Debernardi, A., Buchou, T.,
- Rousseaux, S., Jin, F., 2014. Lysine 2-hydroxyisobutyrylation is a widely distributed active
- 258 histone mark. Nat. Chem. Biol. 10, 365-370.
- Deng, W., Wang, Y., Ma, L., Zhang, Y., Ullah, S., Xue, Y., 2016. Computational prediction of
- 260 methylation types of covalently modified lysine and arginine residues in proteins. Brief.
- Bioinform. DOI: https://doi.org/10.1093/bib/bbw041.
- Elia, A.E., Boardman, A.P., Wang, D.C., Huttlin, E.L., Everley, R.A., Dephoure, N., Zhou, C.,
- Koren, I., Gygi, S.P., Elledge, S.J., 2015. Quantitative proteomic atlas of ubiquitination and
- acetylation in the DNA damage response. Mol. Cell 59, 867-881.
- 265 Flotho, A., Melchior, F., 2013. Sumoylation: a regulatory protein modification in health and
- 266 disease. Annu. Rev. Biochem. 82, 357-385.
- Geiss-Friedlander R, Melchior F., 2007. Concepts in sumoylation: a decade on. Nat. Rev. Mol.
- 268 Cell Bio. 8, 947-956.
- Goodman, C., 2013. Post-translational modifications: Considering conditions. Nat. Chem. Biol.

- 270 9, 601-601.
- 271 Greer, E.L., Shi, Y., 2012. Histone methylation: a dynamic mark in health, disease and
- 272 inheritance. Nat. Rev. Genet. 13, 343-357.
- Hendriks, I.A., D'Souza, R.C., Yang, B., Verlaan-de Vries, M., Mann, M., Vertegaal, A.C., 2014.
- 274 Uncovering global SUMOylation signaling networks in a site-specific manner. Nat. Struct. Mol.
- 275 Biol. 21, 927-936.
- Hendriks, I.A., Vertegaal, A.C., 2016. A comprehensive compilation of SUMO proteomics. Nat.
- 277 Rev. Mol. Cell Bio. 17, 581-595.
- Hirschey, M.D., Zhao, Y., 2015. Metabolic regulation by lysine malonylation, succinylation,
- and glutarylation. Mol. Cell. Proteomics 14, 2308-2315.
- Hornbeck, P.V., Zhang, B., Murray, B., Kornhauser, J.M., Latham, V., Skrzypek, E., 2015.
- 281 PhosphoSitePlus, 2014: mutations, PTMs and recalibrations. Nucleic Acids Res. 43,
- 282 D512-D520.
- Huang, H., Lin, S., Garcia, B.A., Zhao, Y., 2015. Quantitative proteomic analysis of histone
- 284 modifications. Chem. Rev. 115, 2376-2418.
- Lamoliatte, F., Caron, D., Durette, C., Mahrouche, L., Maroui, M.A., Caron-Lizotte, O.,
- Bonneil, E., Chelbi-Alix, M.K., Thibault, P., 2014. Large-scale analysis of lysine SUMOylation
- by SUMO remnant immunoaffinity profiling. Nat. Commun. 5, 5409.
- Lanouette, S., Mongeon, V., Figeys, D., Couture, J.F., 2014. The functional diversity of protein
- 289 lysine methylation. Mol. Syst. Biol. 10, 724.
- 290 Li, J., Jia, J., Li, H., Yu, J., Sun, H., He, Y., Lv, D., Yang, X., Glocker, M.O., Ma, L., 2014.
- 291 SysPTM 2.0: an updated systematic resource for post-translational modification. Database
- 292 2014, bau025.
- 293 Liu, Z., Cao, J., Gao, X., Zhou, Y., Wen, L., Yang, X., Yao, X., Ren, J., Xue, Y., 2011. CPLA
- 294 1.0: an integrated database of protein lysine acetylation. Nucleic Acids Res. 39(suppl 1),
- 295 D1029-D1034.
- 296 Liu, Z., Wang, Y., Gao, T., Pan, Z., Cheng, H., Yang, Q., Cheng, Z., Guo, A., Ren, J., Xue, Y.,
- 297 2014. CPLM: a database of protein lysine modifications. Nucleic Acids Res. 42, D531-D536.
- 298 Lu, C.-T., Huang, K.-Y., Su, M.-G., Lee, T.-Y., Bretaña, N.A., Chang, W.-C., Chen, Y.-J., Chen,
- 299 Y.-J., Huang, H.-D., 2012. DbPTM 3.0: an informative resource for investigating substrate site

- 300 specificity and functional association of protein post-translational modifications. Nucleic Acids
- 301 Res., gks1229.
- Mann, M., Jensen, O.N., 2003. Proteomic analysis of post-translational modifications. Nat.
- 303 Biotechnol. 21, 255-261.
- Mertins, P., Qiao, J.W., Patel, J., Udeshi, N.D., Clauser, K.R., Mani, D., Burgess, M.W.,
- Gillette, M.A., Jaffe, J.D., Carr, S.A., 2013. Integrated proteomic analysis of post-translational
- modifications by serial enrichment. Nat. Methods 10, 634-637.
- 307 Morris, M., Knudsen, G.M., Maeda, S., Trinidad, J.C., Ioanoviciu, A., Burlingame, A.L.,
- 308 Mucke, L., 2015. Tau post-translational modifications in wild-type and human amyloid
- precursor protein transgenic mice. Nat. Neurosci. 18, 1183-1189.
- 310 Olsen, J.V., Ong, S.-E., Mann, M., 2004. Trypsin cleaves exclusively C-terminal to arginine
- and lysine residues. Mol. Cell. Proteomics 3, 608-614.
- Poulsen, C., Akhter, Y., Jeon, A.H.W., Schmitt Ulms, G., Meyer, H.E., Stefanski, A., Stühler,
- 313 K., Wilmanns, M., Song, Y.H., 2010. Proteome-wide identification of mycobacterial
- 314 pupylation targets. Mol. Syst. Biol. 6, 386.
- Prasad, T.K., Goel, R., Kandasamy, K., Keerthikumar, S., Kumar, S., Mathivanan, S.,
- 316 Telikicherla, D., Raju, R., Shafreen, B., Venugopal, A., 2009. Human protein reference
- database–2009 update. Nucleic Acids Res. 37(suppl 1), D767-D772.
- Rabut, G., Peter, M., 2008. Function and regulation of protein neddylation. Embo. Rep. 9,
- 319 969-976.
- 320 Sadhukhan, S., Liu, X., Ryu, D., Nelson, O.D., Stupinski, J.A., Li, Z., Chen, W., Zhang, S.,
- Weiss, R.S., Locasale, J.W., 2016. Metabolomics-assisted proteomics identifies succinylation
- and SIRT5 as important regulators of cardiac function. Proc. Natl. Acad. Sci. U. S. A. 113,
- 323 4320-4325.
- 324 Sadoul, K., Khochbin, S., 2016. The growing landscape of tubulin acetylation: lysine 40 and
- 325 many more. Biochem. J. 473, 1859-1868.
- 326 Schwartz, D., Gygi, S.P., 2005. An iterative statistical approach to the identification of protein
- 327 phosphorylation motifs from large-scale data sets. Nat. Biotechnol. 23, 1391-1398.
- 328 Shaid, S., Brandts, C., Serve, H., Dikic, I., 2013. Ubiquitination and selective autophagy. Cell
- 329 Death Differ. 20, 21-30.

- 330 Shannon, D.A., Weerapana, E., 2015. Covalent protein modification: the current landscape of
- residue-specific electrophiles. Curr. Opin. Chem. Biol. 24, 18-26.
- 332 Sharp, J.S., Nelson, S., Brown, D., Tomer, K.B., 2006. Structural characterization of the E2
- 333 glycoprotein from Sindbis by lysine biotinylation and LC-MS/MS. Virology 348, 216-223.
- 334 Stes, E., Laga, M., Walton, A., Samyn, N., Timmerman, E., De Smet, I., Goormachtig, S.,
- Gevaert, K., 2014. A COFRADIC protocol to study protein ubiquitination. J. Proteome Res. 13,
- 336 3107-3113.
- 337 Strzyz, P., 2016. Post-translational modifications: Extension of the tubulin code. Nat. Rev. Mol.
- 338 Cell Bio. 17, 609.
- 339 Svinkina, T., Gu, H., Silva, J.C., Mertins, P., Qiao, J., Fereshetian, S., Jaffe, J.D., Kuhn, E.,
- 340 Udeshi, N.D., Carr, S.A., 2015. Deep, quantitative coverage of the lysine acetylome using
- 341 novel anti-acetyl-lysine antibodies and an optimized proteomic workflow. Mol. Cell.
- 342 Proteomics 14, 2429-2440.
- Tan, M., Luo, H., Lee, S., Jin, F., Yang, J.S., Montellier, E., Buchou, T., Cheng, Z., Rousseaux,
- 344 S., Rajagopal, N., 2011. Identification of 67 histone marks and histone lysine crotonylation as a
- new type of histone modification. Cell 146, 1016-1028.
- Tan, M., Peng, C., Anderson, K.A., Chhoy, P., Xie, Z., Dai, L., Park, J., Chen, Y., Huang, H.,
- 347 Zhang, Y., 2014. Lysine glutarylation is a protein posttranslational modification regulated by
- 348 SIRT5. Cell. Metab. 19, 605-617.
- Weinert, B.T., Schölz, C., Wagner, S.A., Iesmantavicius, V., Su, D., Daniel, J.A., Choudhary, C.,
- 350 2013. Lysine succinylation is a frequently occurring modification in prokaryotes and
- eukaryotes and extensively overlaps with acetylation. Cell Rep. 4, 842-851.
- 352 Xie, Z., Dai, J., Dai, L., Tan, M., Cheng, Z., Wu, Y., Boeke, J.D., Zhao, Y., 2012. Lysine
- succinylation and lysine malonylation in histones. Mol. Cell. Proteomics 11, 100-107.
- 354 Xiong, Y., Guan, K.-L., 2012. Mechanistic insights into the regulation of metabolic enzymes by
- 355 acetylation. J. Cell Biol. 198, 155-164.
- 356 Xu, H.-D., Shi, S.-P., Wen, P.-P., Qiu, J.-D., 2015. SuccFind: a novel succinylation sites online
- prediction tool via enhanced characteristic strategy. Bioinformatics 31, 3748-3750.
- 358 Xu, Y., Zhang, S., Lin, S., Guo, Y., Deng, W., Zhang, Y., Xue, Y., 2016. WERAM: a database of
- writers, erasers and readers of histone acetylation and methylation in eukaryotes. Nucleic Acids

360	D	gkw1011	
3011	RAS	OKWILLI	

361 Zhang, Z., Tan, M., Xie, Z., Dai, L., Chen, Y., Zhao, Y., 2011. Identification of lysine

succinylation as a new post-translational modification. Nat. Chem. Biol. 7, 58-63.

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364	Figures	
365	Fig. 1. Comparison of CPLA 1.0, CPLM 2.0 and PLMD 3.0.	
366	CPLA, Compendium of Protein Lysine Acetylation; CPLM, Compendium of Protein Lysin	
367	Modifications; PLMD, Protein Lysine Modification Database; Types, total lysine modification	
368	types; Species, total species lysine modification involved; Proteins, total lysine modificatio	
369	protein; Total sites, total lysine modification sites.	
370		
371	Fig. 2. Simplified diagram of 20 types of PLMs with molecular structures of ligands	
372	conjugated to lysine residues.	
373	A-C: Twenty types of PLMs classified into three categories: Ub-UbLs (A), acylation (B) and	
374	other PLMs (C). Ub-UbLs, ubiquitin and ubiquitin-like modifications.	
375		
376	Fig. 3. The heatmaps for the distribution of protein or site number of different PLM types	
377	among different species.	
378	A: The heatmap for the distribution of substrate numbers. B: The heatmap for the distribution	
379	of modified lysine residue numbers.	
380		
381	Fig. 4. Two browse options of PLMD.	
382	A and B : Browse by PLM types. C and D : Browse by species.	
383		
384	Fig. 5. The search options of PLMD.	
385	A: 'Substrate Search' with one or multiple keywords. B: The 'Advanced Search' permits users	
386	to input up to three terms for query. C: The 'Multiple Search' permits users to enter multiple	
387	protein entries with a list of terms. D: The 'BLAST Search' allows users to find identical of	
388	homologous proteins with a protein sequence in FASTA format.	
389		
390	Fig. 6. The over-represented sequence motifs and potential crosstalks of different PLM types	
391	A: Motif-based analyses of sequence preferences around PLM sites. Totally, 16 types of PLM	
392	were found to process the most significantly over-represented sequence motifs. Due to the dat	
393	limitation, the <i>P</i> -value of 2-hydroxyisobutyrylation was set as 0.001, and that of glutarylatio	

394 and phosphoglycerylation was set as 0.0001. The other 13 types of PLMs were analyzed with a 395

P-value < 0.00001. **B**: The heatmap for the distribution of co-occurrences of two different PLM

types. The detailed numbers are provided in Table S2.

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