

Research on the Human Proteome Reaches a Major Milestone: >90% of Predicted Human Proteins Now Credibly Detected, According to the HUPO Human Proteome Project

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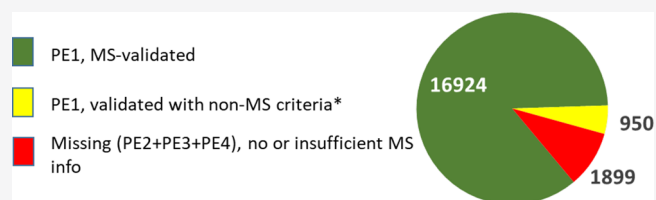
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ABSTRACT: According to the 2020 Metrics of the HUPO Human Proteome Project (HPP), expression has now been detected at the protein level for >90% of the 19 773 predicted proteins coded in the human genome. The HPP annually reports on progress made throughout the world toward credibly identifying and characterizing the complete human protein parts list and promoting proteomics as an integral part of multiomics studies in medicine and the life sciences. NeXtProt release 2020–01 classified 17 874 proteins as PE1, having strong protein-level evidence, up 180 from 17 694 one year earlier. These represent 90.4% of the 19 773 predicted coding genes (all PE1,2,3,4 proteins in neXtProt). Conversely, the number of neXtProt PE2,3,4 proteins, termed the “missing proteins” (MPs), was reduced by 230 from 2129 to 1899 since the neXtProt 2019–01 release. PeptideAtlas is the primary source of uniform reanalysis of raw mass spectrometry data for neXtProt, supplemented this year with extensive data from MassIVE. PeptideAtlas 2020–01 added 362 canonical proteins between 2019 and 2020 and MassIVE contributed 84 more, many of which converted PE1 entries based on non-MS evidence to the MS-based subgroup. The 19 Biology and Disease-driven B/D-HPP teams continue to pursue the identification of driver proteins that underlie disease states, the characterization of regulatory mechanisms controlling the functions of these proteins, their proteoforms, and their interactions, and the progression of transitions from correlation to coexpression to causal networks after system perturbations. And the Human Protein Atlas published Blood, Brain, and Metabolic Atlases.

KEYWORDS: Human Proteome Project (HPP), neXtProt protein existence (PE) metrics, missing proteins (MPs), non-MS PE1 proteins, uncharacterized protein existence 1 (uPE1), Chromosome-Centric HPP (C-HPP), Biology and Disease-HPP (B/D-HPP), PeptideAtlas, Mass Spectrometry Interactive Virtual Environment (MassIVE), Human Protein Atlas



INTRODUCTION

The Human Proteome Project (HPP) of the Human Proteome Organization (HUPO) has provided a useful framework for international collaboration, data sharing and reanalysis, quality assurance, guidelines, communication, and acceleration of progress in building and utilizing proteomic knowledge since its launch in 2010.¹ Remarkable progress has been documented in neXtProt, PeptideAtlas, and MassIVE and the annual Metrics paper in the HPP special issues of the *Journal of Proteome Research* (JPR).^{2–9} Our complementary decadal “A High-Stringency Blueprint of the Human Proteome” draws a striking parallel to the celebration of the completion of 90% of the Human Genome sequence in 2001.¹⁰

To celebrate the announcement of the completion of the first draft of the human proteome by HUPO and the HPP, JPR has published a **Virtual Issue** of the most significant papers in the HPP that have appeared in the past 10 years in the Journal.

PROGRESS ON THE HUMAN PROTEOME PARTS LIST: ACHIEVING THE MILESTONE OF DETECTING 90% OF PREDICTED PROTEIN-CODING GENES

Table 1 shows the increase from 13 975 PE1 proteins in neXtProt release 2012–02 to 17 874 in release 2020–01. This PE1 total now represents 90.4% of the 19 773 PE1,2,3,4 proteins predicted from the Human Genome version GRCh38. Conversely, the number of PE2,3,4 “Missing Proteins” has

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Table 1. Numbers of Proteins Classified in neXtProt Protein Existence Evidence Levels PE1,2,3,4 from 2012 to 2020, Showing Remarkable Progress in Confidently Identifying Predicted Human Proteins as PE1^a

PE level/neXtProt release	2012–02	2013–09	2014–10	2016–01	2017–01	2018–01	2019–01	2020–01
1: Evidence at protein level	13 975	15 646	16 491	16 518	17 008	17 470	17 694	17 874
2: Evidence at transcript level	5205	3570	2647	2290	1939	1660	1548	1596
3: Inferred from homology	218	187	214	565	563	452	510	253
4: Predicted	88	87	87	94	77	74	71	50
MP = PE2 + PE3 + PE4	5511	3844	2948	2949	2579	2186	2129	1899
Human PeptideAtlas canonical proteins	12 509	13 377	14 928	14 569	15 173	15 798	16 293	16 655

^aAs of 2020–01, PE1/PE1 + 2 + 3 + 4 = 17 874/19 773 = 90.4% now PE1. The PE2,3,4 proteins are still “missing proteins” (MP). Dates represent the neXtProt releases used as baseline for the annual HPP metrics papers.^{2–9} More stringent Guidelines 2.1 were imposed in time for the 2016–01 release.¹¹ Since 2016, the January release of neXtProt has served as the annual baseline. The final row shows the critical contribution of canonical proteins resulting from PeptideAtlas reanalyses of available MS datasets (see text and Figure 1).

declined from 5511 in 2012 to 1899 in 2020, including a reduction of 230 from 2019.

The components of the HPP are the Chromosome-centric C-HPP, with 25 teams addressing the 24 chromosomes and mitochondria; the Biology and Disease-driven B/D-HPP, with 19 teams investigating organs, diseases, and biological functions; and the four Resource Pillars for Mass Spectrometry, Antibody-profiling, Knowledgebases, and Pathology. The C-HPP held its 21st Workshop on Future Strategies for the Human Proteome Project at Saint-Malo, France, in May 2019. The HPP Mass Spectrometry Data Interpretation Guidelines 3.0 were finalized, and integration of the MassIVE data resource was agreed upon for the neXtProt 2020–01 release. On the basis of the Guidelines 3.0,¹² neXtProt has 16 836 entries that were validated by PeptideAtlas, 15 214 entries validated by MassIVE, 16 920 validated by one resource or the other, 15 130 by both resources, 84 only by MassIVE, and 1706 only by PeptideAtlas. There are four additional entries identified and curated by neXtProt from PTM proteomics papers. [Supplementary Table S1](#) contains Excel files with all of these lists from neXtProt. In order to call a protein validated by MS, neXtProt requires that the minimum MS-evidence requirements described in the HPP Guidelines 3.0 are satisfied by either PeptideAtlas or MassIVE (or both) independently; i.e., insufficient evidence in both PeptideAtlas and MassIVE, such as one peptide in each, may not be combined to form sufficient evidence.

A major adjustment had occurred in 2015 as the HPP Guidelines 1.0 based on false-discovery rates ≤ 0.01 at the protein level were superseded by Guidelines 2.1, requiring two proteotypic (uniquely mapping) non-nested peptides of at least 9 amino acids in length.¹¹ These requirements removed many proteins that were “one-hit wonders”, were based on peptides as short as 6 or 7 amino acids, or matched better to well-established proteins. That caused 485 proteins that would previously have been classified as PE1 to be put in the PE2,3,4 missing proteins set (438 PE2, 40 PE3, 7 PE4); as shown in [Table 1](#), that substantially slowed the identification of PE1 proteins from 2014 to 2016. Meanwhile, in 2016 there was a large increase in PE3 (214 to 565) due to a decision of UniProt/SwissProt to remove upgrades to PE2 that relied on inclusion in the ArrayExpress or CleanEx transcriptome repositories, while greatly increasing the number of PE3 entries based only on homology (expression detected in nonhuman species).⁵ For the 2020–01 release neXtProt incorporated the merged RNA_seq data sets from Human Protein Atlas, GTEx, and FANTOM5¹³ to upgrade PE3 and PE4 to PE2 if the expression values were ≥ 1 FPKM (fragments per kilobase of exon model per million reads mapped).

[Figure 1](#) shows the major categories of experimental evidence upon which neXtProt has classified the human proteome. Of the

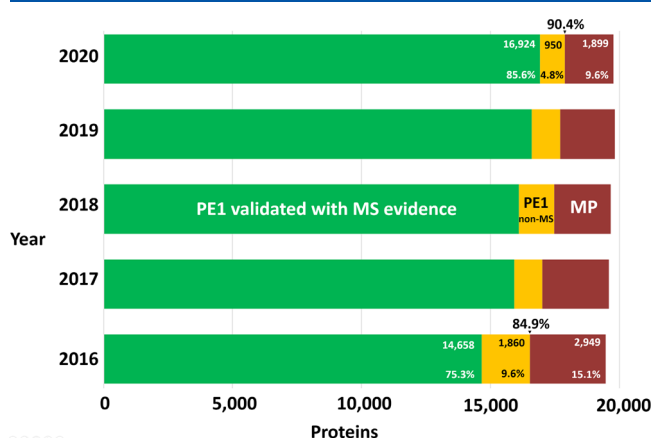


Figure 1. Bar chart showing the progression of the state of protein validation for the past five years, since the HPP Guidelines 2.1 have been in effect. The PE1 proteins validated with mass spectrometry (MS) evidence are depicted in green, PE1 proteins based on other types of protein evidence in yellow, and the PE2,3,4 missing proteins (MPs) that have no or insufficient protein-level evidence in human specimens and cell lines in red. The green bars for 2016–2019 are from PeptideAtlas, while for 2020 it is a combination with MassIVE. Data for the green + yellow (PE1) and red (PE2,3,4) bars are provided in [Table 1](#).

17 874 PE1, 16 924 are now based on validated mass spectrometry results (green bar). The yellow bar in [Figure 1](#) represents 950 PE1 proteins identified by non-MS methods and curated by SwissProt/neXtProt based on the published literature: 73 from Edman sequencing, 122 from disease mutations, 35 from 3D structures, 342 from protein–protein interactions, 49 from antibody-based techniques, 127 from PTMs or proteolytic processing, and 202 from biochemical studies; because many proteins have multiple lines of evidence, these numbers represent the first evidence reported for each protein. [Supplementary Table S2](#) provides a comprehensive list of the kinds of evidence used by SwissProt curators to classify 906 of these 950 as PE1, while [Supplementary Table S3](#) further annotates 820 entries of this list with information from the Human Protein Atlas (see below). There are no HPP Guidelines for Data Interpretation for these different studies grouped in the non-MS category; SwissProt describes their PE process at https://www.uniprot.org/help/protein_existence. The yellow bar has declined quite dramatically from 1860 in 2016 to 950 in 2020, as more and more proteins have been confirmed with

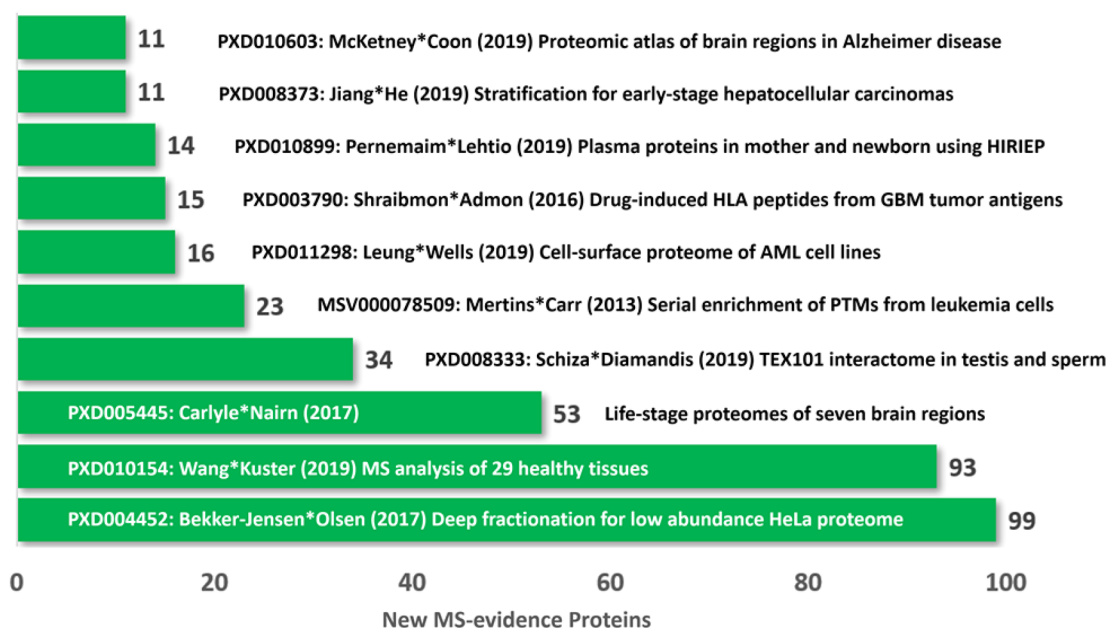


Figure 2. Top 10 data sets that contributed to the increment in canonical proteins in between 2019 and 2020 builds of PeptideAtlas. These new MS-based canonical proteins represent both new PE1 proteins that previously were PE2,3,4 (red in Figure 1) and previous non-MS-based PE1 proteins (yellow in Figure 1) that are now validated with MS evidence in PeptideAtlas. Each data set is labeled with the ProteomeXchange identifier (PXD), first*last authors, publication year, number of new canonical proteins, and method highlighted.^{14–23}

independent identification by increasingly sensitive and accurate mass spectrometry analyses. The missing proteins PE2,3,4 (Table 1) now number 1899 (red bar in Figure 1), hence 9.6% of the total predicted proteins.

From its beginning in 2012 neXtProt has relied on the standardized reanalysis of available data sets by PeptideAtlas. From these, there are now 16 655 canonical proteins in the 2020–01 Human PeptideAtlas build (Table 1). To the 2019–01 build, PeptideAtlas added 257 samples from 24 ProteomeXchange data sets comprising 101 million new spectrum identifications that passed stringent thresholds. This subset of data sets submitted to ProteomeXchange was selected to contribute new identifications reflecting new instrumentation, methods, or sample types. Note that the more data that are added, the more stringent the threshold must be for all data sets in order to maintain a constant FDR; thus, including data sets that do not contribute new identifications is counterproductive.

Figure 2 shows the top 10 newly analyzed data sets located through ProteomeXchange contributing to the increased number of canonical proteins in PeptideAtlas during 2019; they account for 369 of the 413 added canonical proteins. Note that, while 413 canonical proteins were added in PeptideAtlas, 51 previously canonical proteins were removed from the reference proteome due to Swiss-Prot entry merging as discussed below, yielding the net increase of 362 in Table 1. While the HPP and JPR encourage investigators to validate their findings of newly detected missing proteins with synthetic peptides, neither neXtProt nor PeptideAtlas nor MassIVE requires confirmatory synthetic peptide MS matching results for their annual updates of community MS data for the human proteome. Efforts to incorporate synthetic peptide MS matching are underway, but PeptideAtlas does not have an automated method to incorporate SRM or PRM results. Nevertheless, several PRM data sets are now included in PeptideAtlas, applying the HUPO/PSI Universal Spectrum Identifier in spectrum-based guidelines, as PRM acquires full MS2 spectra.

The largest gain came from PeptideAtlas reprocessing of PXD004452, a deep analysis of the extensively studied HeLa cell,^{24–26} using shotgun proteomics, three proteases, high peptide load, and 2D high-resolution reversed-phase peptide fractionation to saturate the sequencing speed with short gradients, as well as modern MS instruments.²³ The original paper reported 584 000 unique peptide sequences and 14 200 protein isoforms from 12 200 protein-coding genes, 7000 N-acetylation sites, and 10 000 phosphorylation sites. This depth is comparable to the depth of next-generation RNA sequencing. Enhancing the sensitivity of the MS protocol reaches less-abundant proteins.

The second largest gain was from PeptideAtlas reprocessing of PXD010154, with combined RNA-seq and MS analyses of adjacent cryosections from 29 tissues.²² They reported detection of 13 640 proteins, including 37 PE2,3,4 MPs, based on “at least 1 unique peptide” confirmed with a synthetic peptide; only 8 met the HPP Guidelines 2.1 of two uniquely mapping peptides of at least 9 aa. Correlation of mRNA and protein abundance was generally low; 478 proteins were not detected in testis and 1408 across all tissues even when mRNA FPKM values were above the mean values. Moreover, 300 of the 478 not detected in testis by MS were not detected in HPA with antibodies. Among transcripts below the mean FPKM, GPCR, ion channels, and cytokine-related proteins were notable missing proteins. An extensive analysis showed challenges in finding protein-level evidence of sequence variants or splice isoforms; not a single lncRNA peptide was substantiated. The 93 new canonical proteins found by PeptideAtlas predominantly represent proteins previously classified as PE1 by non-MS methods.

The third largest gain with 53 new canonical proteins came from PeptideAtlas reanalysis of PXD005445, data from the BrainSpan Project, which examined 7 regions of the brain in individuals in 6 age intervals from infancy to age 42.²¹ After testis, brain is the most promising organ for deeper analysis to

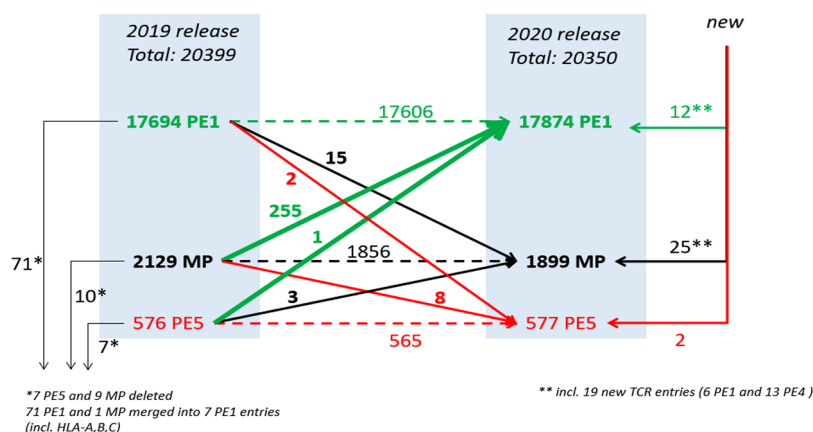


Figure 3. Flowchart depicts the detailed changes in neXtProt PE1–4 categories (plus PE5) from neXtProt release 2019–01 to neXtProt release 2020–01 (see text).

Table 2. Chromosome-by-Chromosome Status of Predicted Proteins in neXtProt 2020–01^a

Chr	PE1	PE2	PE3	PE4	PE2 + 3 + 4	PE1/PE1–4	% PE1/PE1–4	net decrease PE2,3,4 (2016–2020)	uPE1 proteins
1	1819	171	27	3	201	1819/2020	90.0%	108 (309/201)	145
2	1193	68	9	2	79	1193/1272	93.8%	55 (134/79)	87
3	981	72	9	2	83	981/1064	92.2%	82 (141/59)	55
4	688	44	15	0	59	688/747	92.1%	36 (95/59)	50
5	817	43	2	1	46	817/863	94.7%	76 (122/46)	50
6	930	72	7	4	83	930/1013	91.8%	53 (136/83)	59
7	828	81	37	16	134	828/962	86.1%	3 (137/134)	53
8	612	42	7	2	51	612/663	92.3%	44 (95/51)	35
9	683	80	8	3	91	683/774	88.2%	38 (129/91)	59
10	670	61	2	1	64	670/734	91.3%	51 (115/64)	52
11	1029	205	58	1	264	1029/1293	79.6%	55 (319/264)	71
12	942	66	3	0	69	942/1011	93.2%	50 (119/69)	52
13	304	17	1	0	18	304/322	94.4%	25 (43/18)	25
14	614	50	43	5	98	614/712	86.2%	–5 (93/98)	32
15	524	47	2	0	49	524/573	91.4%	24 (73/49)	38
16	756	60	2	0	62	756/818	92.4%	37 (99/62)	47
17	1061	83	2	2	87	1061/1148	92.4%	61 (148/87)	66
18	252	12	1	0	13	252/265	95.1%	11 (24/13)	10
19	1281	112	10	1	123	1281/1404	91.2%	138 (261/123)	80
20	490	45	0	3	48	490/538	91.1%	34 (82/48)	42
21	191	36	5	1	42	191/233	82.0%	7 (49/42)	11
22	439	33	2	1	36	439/475	92.4%	28 (64/36)	33
X	731	82	1	2	85	731/816	89.6%	60 (145/85)	101
Y	27	13	0	0	13	27/40	67.5%	3 (16/13)	1
Mito	15	0	0	0	0	15/15	100.0%	0 (0/0)	0
Unk	2	2	0	0	2	2/4	50.0%	0 (2/2)	0
ALL	17 874	1596	253	50	1899	17 879/19 779	90.4%	1050 (2949/1899)	1254
Sums	17 879	1597	253	50	1900	17 699/19 829	90.2%	1050 (2950/1900)	1254

^aNote: There are discrepancies between the true total numbers of proteins in each PE category (**ALL**) and the Sums because six proteins are derived from two genes on two different chromosomes, and thus appear twice under the per-chromosome table values.

find MPs.²² Their analysis revealed that 8980 proteins were detected in at least one region and 6529 in all 7 regions. Differential analysis of protein expression by region showed prominence for nuclear functions, including RNA processing, especially in the cerebellum. Our [Supplementary Table S3](#) provides information from the Human Protein Atlas of transcript expression and immunohistochemical localization of 820 of the 950 non-MS-based PE1 proteins, ordered alphabetically by tissue type.

■ DYNAMIC CHANGES IN neXtProt CLASSIFICATION OF PROTEINS

The year-to-year changes in neXtProt categories, including their inputs from UniProt/SwissProt, are quite dynamic and quite challenging to track, as shown in [Figure 3](#). Note that 255 PE2,3,4 MPs were promoted by new evidence to PE1, while 10 MPs were deleted, 8 were demoted to PE5, 15 were gained from PE1, and 25 were newly added to neXtProt. PE1 had an unusual reason for a smaller increase than what came from PE2,3,4 MPs (a net of 180): 71 PE1 entries were merged into 7, mostly due to a major reclassification of HLA-A, HLA-B, and HLA-C genes

and their protein products. In 2013, we noted that there were 21 genes for HLA-A, 35 for HLA-B, 14 for HLA-C, and 13 for HLA-DRB1 as neXtProt entries for Chromosome 6.² Also, 15 previous PE1 proteins were demoted to PE2,3,4. Further details show 12 PE1 entries new to neXtProt, of which 6 involved the family of T Cell Receptor proteins, from which 13 more were added to the MPs according to predictive models (PE4). Previous annual metrics papers have reported corresponding dynamic year-to-year changes.^{6–8}

There were multiple minor changes in the category of PE5. In 2013, the HPP excluded category PE5 from the denominator of all predicted proteins, due to the status of these sequences as “dubious or uncertain” genes with a high proportion of pseudogenes.³ When C-HPP announced the MP50 Missing Proteins Challenge, PE5 entries were excluded.⁶

■ UPDATE FROM THE CHROMOSOME-CENTRIC HPP (C-HPP)

The 25 teams of the C-HPP have mounted two initiatives to characterize “the Dark Proteome”: the 2016 neXt-MP50 Challenge to find or document the finding of at least 50 PE2,3,4 missing proteins per chromosome⁵ and the 2018 neXt-CP50 Challenge to use experimental and informatics approaches to annotate functions for 1260 unannotated PE1 proteins (uPE1).⁷ Both challenges reflect the overall goal of the HPP to both detect and characterize the predicted human proteins. Among the advances on missing proteins that came from the entire community, there were 32 MPs identified in the 2019 HPP special issue of *JPR* and proposed to be reviewed for PE1: 1 from rarely studied bone (plus 5 non-MS PE1),²⁷ 3 from placenta,²⁸ 5 from testis,²⁹ which has been a rich source of MPs, and 23 from matching peptides to their counterparts in SRM Atlas.³⁰ Of the 9 PE1 candidates from the first three papers, Q6NT89/TMF-regulated nuclear protein (chr 1) from Lin et al., and Q8TAA1/RNase11 (chr 14) and Q8WW27/ABOBEC-4 (chr 1) from Sun et al. became PE1 in neXtProt 2020–01, as did Q5T5N4/C6orf118, which initially did not quite meet Guidelines criteria; the other 6 of the 9 remained PE2. None of these 3 data sets was released publicly in time to be included in the PeptideAtlas update release of 2020–01. Of the 23 from Elguoshy et al.,³⁰ 19 became PE1.

As of release 2019–01,⁸ chromosomes 19, 1, 5, and 17 had each noted a reduction in their number of PE2,3,4 MPs by 50 or more; as of release 2020–01, MPs had been reduced by at least 50 for Chromosomes 2, 3, 6, 10, 11, 12, 22, and X, as well (www.nextprot.org/about/protein-existence), as shown in Table 2, Chromosome-by-Chromosome Status of Predicted Proteins in neXtProt 2020–01.

The C-HPP also monitors progress on demonstrating and predicting functions for the 1260 uPE1 proteins unannotated for function as of 2018, when C-HPP initiated the next-CP50 Challenge seeking to stimulate discovery and confirmation of function. Table 2 (column 10) gives the uPE1 numbers for each chromosome. The column sum of 1254 is identical with that for 2019. While many chromosome uPE1 numbers in 2020 are very close to 2019,⁸ chromosomes 1, 6, and 19 stand out with increases from 138 to 145, 54 to 59, and 77 to 80, respectively, as the PE1 total grew larger.

Another significant development was the application of the I-TASSER/COFACTOR algorithms to predict Gene Ontology terms for uPE1 proteins.^{31,32} neXtProt added a link for individual protein entries to facilitate submission of uPE1 proteins for a report of predicted functions. As of May 15,

2020, documented requests for C-I-TASSER function predictions totaled 561 proteins from 181 users from 35 countries, including 201 neXtProt proteins [<https://zhanglab.ccmb.med.umich.edu/C-I-TASSER/bin/stat.cgi>].

■ HIGHLIGHTS FROM PEPTIDE ATLAS AND MASSIVE

The PeptideAtlas processing pipeline at a high level includes conversion of vendor files to mzML³³ with the ProteoWizard³⁴ msconvert tool. The mzML files are processed through Comet^{35,36} using the most recent THISP³⁷ Level 3 database and SpectraST³⁸ using the most relevant NIST spectral library. The search results are validated with PeptideProphet³⁹ and iProphet⁴⁰ tools within the Trans-Proteomic Pipeline^{41–43} environment. Data sets encompass all the most common instrument types, fragmentation modes, and labeling methods, including iTRAQ and TMT.⁴⁴ Most data sets are available in ProteomeXchange,⁴⁵ although there remain 114 older samples that predated ProteomeXchange and are not available there. All 1804 samples are thresholded at a constant iProphet-model estimated 0.0002 PSM-level FDR in order to achieve a 0.01 protein-level FDR. Canonical proteins require two non-nested uniquely mapping peptides of length 9 or more residues, as described in the HPP Guidelines 3.0.¹² However, all peptides are reported to neXtProt and undergo remapping and filtering there. All spectra supporting the Human PeptideAtlas 2020–01 Build are available interactively via the web interface. All spectra that belong to a data set deposited in ProteomeXchange are referenced by their HUPO Protein Standards Initiative Universal Spectrum Identifier (USI).¹²

The MassIVE processing pipeline (Mass Spectrometry Interactive Virtual Environment)⁴⁶ also starts with ProteoWizard conversion of raw files into mzML using msConvert, and follows it with MS-GF+ database search against the UniProt human reference proteome with isoforms as well as common contaminants. To ensure strict FDR controls at multiple levels, the MassIVE pipeline follows the MassIVE-KB process for construction of spectral libraries, with 0.1% spectrum-level FDR, <1% length-specific peptide level FDR, and 1% FDR for all proteins matched by at least one unique peptide, which corresponds to 0.013% protein-level FDR for proteins matched by 2+ peptides. In total, 227 unlabeled data sets from 27 404 LC-MS runs encompassing 658 million MS/MS HCD spectra were reprocessed. All spectra were mapped to exons in ENSEMBL; we noted whether the peptide maps uniquely to an exon, covers a splice junction, or is mapped to an exon at all. Canonical proteins require two non-nested uniquely mapping peptides of length 9 or more residues, per HPP Guidelines 3.0. However, all peptides are reported to neXtProt and undergo remapping and filtering there. All spectra are viewable online using MassIVE's ProteinExplorer,⁴⁷ including the complete provenance from each sequence to spectra, searches, and data sets that support it. Additionally, all matching synthetic spectra can be viewed for each sequence.

A major asset of the HPP is ProteomeXchange.⁴⁵ All MS data sets submitted to the *JPR* annual issues are required to have a PXD identifier for manuscript reviewers and readers to access the primary data sets and metadata in one of the participating repositories: PRIDE,^{48,49} PeptideAtlas,^{50,51} MassIVE,⁴⁷ jPOSTdb,⁵² iProX,⁵³ or Panorama Public.⁵⁴ As of October 3, 2020, ProteomeXchange listed 12 801 all-species data sets and 5296 human data sets [www.proteomexchange.org].

■ HIGHLIGHTS FROM THE BIOLOGY AND DISEASE-DRIVEN HPP (B/D-HPP)

The B/D-HPP comprises 19 teams (<https://hupo.org/B/D-HPP>) that focus on distinct biomedical research areas relevant to understanding basic biology and the development of human diseases. The B/D teams are collaborative communities that are open to the participation of scientists from around the globe and welcome productive interactions that disseminate the use of proteomics across the research community. The B/D-HPP has been active in developing the SRMAtlas, the use of DIA-SWATH-MS, and the search for organ-specific priority proteins.

Themes at the core of all B/D-HPP teams include (1) the identification of driver proteins that underlie health and disease states, and (2) the characterization of regulatory mechanisms that control the functions of these proteins, including posttranslational modifications, protein interactions, physicochemical properties, and their dynamic organization within a cell or tissue.

The identification and characterization of driver proteins provide critical insights into the molecular underpinning of health and disease, then guide the development of assays for early detection of disease and innovative therapeutic interventions. Two examples in 2019 of collaboration between B/D-HPP teams included the Plasma and Diabetes teams applying a robust workflow to monitor 1508 plasma proteins and cellular pathways associated with weight loss and weight maintenance,⁵⁵ and the Rheumatic and Autoimmune Disorders and Cardiovascular teams defining a panel of priority proteins that reveal both relevant mechanistic insights and promising biomarker candidates.⁵⁶

The ability to profile proteomes with high sensitivity and accuracy has uncovered disease- and age-associated signatures.^{52–54} Another study identified changes in alternative splicing and autophagy connected to a decline in skeletal muscle function.⁵⁷ As members of the B/D-Cancer team, Jiang et al. identified novel therapeutic targets of early stage hepatocellular carcinoma,¹⁵ and the NCI Clinical Proteomics Tumor Analysis Consortium has provided vast publicly accessible data sets and major resource papers in 2019 on proteogenomic and immune characterization of colon cancers⁵⁸ and clear cell renal cell carcinomas,⁵⁹ with others in the pipeline. Phosphoproteome analyses of kinases and kinase substrates informed prediction of promising targeted therapies. The Infectious Diseases team identified secreted proteins as microbial biomarkers for early detection of infection with the pathogen causing Lyme disease, *Borrelia burgdorferi*.⁶⁰

Several elegant studies demonstrate the fundamental role of proteomics in biological discoveries, such as characterizing phenotypes associated with distinct protein aggregation states: pathological TDP-43 conformers that have differential toxicities in neurons and correlate with frontotemporal lobar neurodegeneration;⁶¹ cullin-RING ubiquitin ligase and diacyl glycerol biosynthesis associated with protein aggregation events in neurodegenerative diseases;^{62,63} and the necessity of nuclear aggregation of a DNA sensing protein, IFI16, for human cells to elicit immune signaling and suppress viral gene expression during a herpesvirus infection.⁶⁴

Protein posttranslational modifications (PTMs) have received special attention: glycosylated immunoglobulin light chain is a biomarker for early diagnosis of amyloidosis;⁶⁵ tyrosine-nitration of cystathionine β -synthase blocks the trans-sulfuration pathway in acute pancreatitis, promoting homo-

cysteine accumulation upon S-adenosylmethionine treatment;⁶⁶ a top-down approach revealed deamidation, methylation, acetylation, trimethylation, phosphorylation, and S-glutathionylation of sarcomeric proteins of nonhuman primate skeletal muscle;⁶⁷ and protein lysine hyperacetylation contributes to development of type 2 diabetes when mitochondrial activity of beta cells or insulin target tissues is perturbed.⁶⁸

These accomplishments were aided by the methods development components of many B/D-HPP teams to promote enhanced proteomic and multiomic experimental workflows, as well as computational platforms for data analysis and integration. The cancer team reported a data integration pipeline for predicting kinase activities from phosphoproteome data sets,⁶⁹ while the cardiovascular team published a method for automated contaminant removal from peptide samples.⁷⁰ A broadly applicable strategy from the Aebersold laboratory explores the transitions from correlation to coexpression to causal networks,⁷¹ using an experimental design with at least two systems perturbations to identify causal relationships of proteins in complex processes. This approach could identify new proteins involved in biological processes and their functions, thereby breaking the trend of a high percentage of omics-era investigations focused on the already-most-studied proteins.

To promote the dissemination and application of proteomics methods within diverse biological and clinical studies, members of B/D-HPP teams have organized educational activities, conferences, and symposia, including during 2019 the Brain Proteome Project Workshop in Amsterdam, a FASEB conference on protein acetylation in Lisbon, the Human Immunopeptidome satellite meeting at HUPO-Adelaide, a virtual EMBL Workshop on Proteomics in Cell Biology and Disease Mechanisms, the Applied Public-Private Research enabling OsteoArthritis Clinical Headway (APPROACH) project, and a journal special issue on Understanding Proteome Organization in Space and Time.⁷²

■ HIGHLIGHTS FROM HUMAN PROTEIN ATLAS

The prolific output from the Human Protein Atlas (HPA), which serves as the Antibody-profiling Resource Pillar of the HPP, continues. In 2019 the Tissue, Cell, and Pathology Atlas series was complemented by releases of the Blood, Brain, and Metabolic Atlases.^{73–75} The Blood Atlas focuses on expression of all protein-coding genes in different blood cell types, as well as a separate analysis on The Human Secretome,⁷⁶ which provides a comprehensive annotation of 2600 proteins predicted to be secreted by human cells. This represents a large fraction of the targets for pharmaceutical drugs, current clinical chemistry, and future diagnostics, and categorizes the different secreted proteins as secreted locally from different tissues or secreted into blood. For the 729 proteins that were found to be secreted into blood, concentration levels based on mass spectrometry or antibody-based immune assays are provided.⁷⁶ Besides classical plasma proteins, cytokines, interleukins, hormones and receptors, about 100 have no known function (presumably uPE1 or uPE2,3,4; candidates for I-TASSER analysis). Many of these circulating proteins are not currently detected with mass spectrometry-based proteomics or antibody-based immunoassays. The analysis revealed that assays are lacking for a large fraction of the secreted blood proteins.

HPA applied its vast array of antibody reagents to COVID-19 studies. A separate portal has been created with direct links to antibody-based protein data for >300 proteins suggested to interact with SARS-CoV-2. For a full understanding of the

susceptibility for SARS-CoV-2 infection and aid in development of effective treatments, it is necessary to study the cell type-specific expression of COVID-19 related proteins on the tissue, cellular and subcellular levels. Surprisingly, an in-depth characterization in >150 different cell types of the angiotensin I converting enzyme 2 (ACE2), the widely presumed target of the SARS-CoV-2 spike protein, revealed very limited expression in human lung and respiratory epithelia.⁷⁷ Instead, highest ACE2 expression was observed in enterocytes, renal tubules, gallbladder, cardiomyocytes, male reproductive cells, placental trophoblasts, ductal cells, eye and vasculature. These other organs show prominent damage in some COVID-19 patients. The study highlights the need for further studies on samples from COVID-19 patients to confirm the histological colocalization of the virus with ACE2 and other SARS-CoV-2 related proteins, including the host serum protease TMPRSS2, as these proteins constitute targets for ongoing drug development efforts. There is evidence that ACE2 expression in airway epithelial cells is stimulated by interferon.⁷⁸

While neXtProt includes information from HPA for protein characterization, and the HPA utilizes neXtProt information, the antibody-based data generated by HPA are currently not sufficient for PE1 classification in the absence of other compelling evidence. From version 19 of the HPA, 56 proteins defined as missing proteins and 171 uPE1 proteins were targeted by antibodies of high reliability meeting the criteria for enhanced validation as defined by the International Working Group for Antibody Validation (IWGAV).^{79,80} Further special efforts are being developed to link antibody-based protein detection and localization with MS-based identification of proteins, especially missing proteins, led by Cecilia Lindskog and Charles Pineau. The initial focus will be on testis, known to harbor a large number of missing proteins.⁸¹ This represents a special version of the general strategy of enrichment of low abundance proteins the HPP has been encouraging for several years.

We have created [Supplementary Tables S2](#) and [S3](#) listing the 950 non-MS-based PE1 proteins (see above). For 906 it was feasible to retrieve results from up to 10 different types of studies contributing to the PE1 assignment. Of the 950, 820 proteins matched entries in HPA version 19 based on Ensembl version 92. Only 10 entries were solely based on antibody information, but 127 have available information on high confidence (reliability score = Enhanced or Supported) in the HPA. These could be priority targets for further studies. The table also includes information on tissue specificity based on mRNA expression data across 37 different organs based on HPA, GTEx and FANTOM5 data sets; 707 had elevated expression in 1–5 tissues, which could guide tissue-specific efforts to target these proteins, while 96 had low tissue specificity, presumably representing widely expressed housekeeping proteins.

DISCUSSION

The neXtProt and PeptideAtlas releases of 2020–01 mark a major milestone in identifying and characterizing the human proteome, with stringent protein-level PE1 detections exceeding 90% of the predicted proteins according to neXtProt. For historical perspective, the Human Genome Project had a major public celebration hosted by President Clinton on June 26, 2000 when the sequencing “approached 90%” of the goal. Of course, much more research is underway to capture and characterize splice variants, PTMs, proteolytic products, and protein–protein, protein–nucleic acid, and protein–lipid interactions and their roles in biological networks in health and disease. For

nearly a decade we have called this phase “building the parts list”.³

Simultaneously, the proteomics community strives to make proteomics a regular and major part of multiomics research. The critical parameters of dynamics, localization, and function of proteins and especially their splice variants, sequence variants, and various PTMs can be clarified only by direct study of proteins individually and as part of the cellular proteome; they cannot be predicted from DNA and RNA studies. Thus, the emergence of proteogenomics as a strategic approach to molecular characterization of cancers, cardiovascular diseases, brain circuitry, reproductive processes, and infectious agent/host interactions is a harbinger of a productive future.

Proteomics is poised to play an increasing role in “precision medicine” and “precision health”, including use of proteomic results to predict clinically useful targets for small molecule and protein or antibody therapeutics in many disease categories. Proteins likewise are critical to development of companion individual cells, thereby revealing the heterogeneity of tissue and organ function. As always, advances in instrumentation and methods are critical for asking new questions about the biology and creating new diagnostics and therapeutics. diagnostic tests to guide use of chemo- and immunotherapies. At the cellular level, there is a major impetus to conduct RNA-seq, protein, and proteoform analyses in

A specific initiative of the HUPO Human Proteome Project since 2016 has been the neXt-MP50 Challenge to each HPP chromosome team to identify or document the identification by others of at least 50 neXtProt PE2,3,4 “missing proteins”, proteins not yet detected in any human specimen with credible high-quality protein evidence, either by mass spectrometry meeting HPP Guidelines or by expert curation of a variety of non-MS protein methods (see [Figure 1](#) and [Supplementary Table S2](#)). This Challenge now has been met for 12 of the 24 chromosomes ([Table 2](#)). Over the past three years, as some investigators have reported finding only small numbers of missing proteins, we have wondered whether the ability to detect more MPs is becoming saturated. However, we see in [Table 1](#) and [Figures 1](#) and [3](#) that the number and proportion of PE2,3,4 missing proteins continue to decline and the number of PE1 proteins continues to grow. In the most recent year, PE1 proteins grew by a net of 180 proteins and 255 PE2,3,4 proteins were promoted to PE1 ([Figure 3](#)). It is noteworthy that detection by MS has also reduced the number of PE1 proteins identified only or primarily by non-MS methods from 1860 to 950 in the past four years.

There are several major reasons for Missing Proteins: transcription and translation may fail to generate sufficient protein to be detected with current MS or other methods; proteins’ sequences may not yield two non-nested, uniquely mapping tryptic peptides of at least 9 amino acids, especially among highly homologous families of proteins; and/or membrane proteins may not be solubilized. There are large differences in gene expression across tissue types and the life span. A review of HPA, GTEx, and FANTOM5 transcript databases led to an estimate of 800 to 1000 PE2,3,4 predicted proteins lacking detectable transcript expression across all tissues.¹³ There are 399 olfactory receptor genes for which no proteins have been found, even when long-sought supra-nasal olfactory epithelium specimens were analyzed,⁸² though that study did find 5 other MPs. Some distinctive tissue types may not yet have been studied, illustrated by the report of 15 MP candidates in fallopian tube.²² For proteins expressed at levels

below the sensitivity of current methods, there is progress as enrichment strategies, fractionation methods, and MS instruments continue to increase sensitivity, illustrated in Figure 2 by the HeLa cell proteome.²³ Data on transcript expression levels can guide researchers to the cell or organ types most likely to be expressing the protein, of which testis and brain have been most productive. Several clever methods have been applied by HPP investigators to enrich low-abundance proteins, including adsorption to ProteoMiner hexapeptide-covered beads,⁸³ and the plans to utilize HPA antibodies together with MS to capture and thereby enrich still-missing proteins in testis, guided by transcript expression data (see HPA, above).

Meanwhile, some proteins whose sequences lack two predicted proteotypic tryptic peptides of at least 9 aa have been detected by using additional proteases with different cleavage specificity⁸⁴ or finding N- and C-terminal peptides and peptides with missed cleavages.⁷ Low MW proteins also are under-represented in PE1; specific efforts to enrich for these proteins have not yet yielded many PE1 candidates, nor have top-down methods revealed a new PE1 protein. Detection based on protein–protein interactions must avoid claiming recombinant bait proteins as expressed. An analysis of homologous protein families was a special feature of a paper about how the first 43 newly detected PE1 proteins from Chromosome 17 were found as of 2018 in the neXt-MPSO Challenge.⁸⁵

Major efforts have been directed at membrane proteins.^{86,87} Hydrophobic proteins account for about 40% of missing proteins; Lys and Arg residues tend to be few and either too close together in hydrophilic loops or too far apart for tryptic digestion to be useful. Other proteases have similar problems, generally generating higher sequence coverage but few additional protein identifications. Large families or groups include GPCR, zinc finger, homeobox, keratin-associated, and coiled-coil domain proteins.^{88,89}

We expect all of these strategies for finding missing proteins, characterizing the functions of already-detected proteins, and utilizing proteogenomics in precision medicine to be fruitful in the coming years.

■ ASSOCIATED CONTENT

SI Supporting Information

The Supporting Information is available free of charge at <https://pubs.acs.org/doi/10.1021/acs.jproteome.0c00485>.

Supplementary Table S1: neXtProt Analysis of the 17 874 MS-based PE1 Proteins from PeptideAtlas and MassIVE (organized by PeptideAtlas only, MassIVE only, additional neXtProt entries, and All) (XLSX)

Supplementary Table S2: Evidence for the 950 non-MS-based PE1 neXtProt Proteins (ordered by neXtProt accession code, Column A, for 906 with retrievable data); Column headings nearly match columns A to O in Supplementary Table S3 (XLS)

Supplementary Table S3: Human Protein Atlas Immunohistochemistry and Transcript Expression Findings for 820 of the 950 non-MS-based PE1 neXtProt Proteins (ordered alphabetically by tissue with highest expression, Column S) (XLS)

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Notes

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