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The SysteMHC Atlas project

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ABSTRACT

Mass spectrometry (MS)-based immunopeptidomics investigates the repertoire of peptides presented at the cell surface by major histocompatibility complex (MHC) molecules. The broad clinical relevance of MHC-associated peptides, e.g. in precision medicine, provides a strong rationale for the large-scale generation of immunopeptidomic datasets and recent developments in MS-based peptide analysis technologies now support the generation of the required data. Importantly, the availability of diverse immunopeptidomic datasets has resulted in an increasing need to standardize, store and exchange this type of data to enable better collaborations among researchers, to advance the field more efficiently and to establish quality measures required for the meaningful comparison of datasets. Here we present the SysteMHC Atlas (https://systemhcatlas.org), a public database that aims at collecting, organizing, sharing, visualizing and exploring immunopeptidomic data generated by MS. The Atlas includes raw mass spectrometer output files collected from several laboratories around the globe, a catalog of context-specific datasets of MHC class I and II peptides, standardized MHC allele-specific peptide spectral libraries consisting of consensus spectra calculated from repeat measurements of the same peptide sequence, and links to other proteomics and immunology databases. The SysteMHC Atlas project was created and will be further expanded using a uniform and open computational pipeline that controls the quality of peptide identifications and peptide annotations. Thus, the SysteMHC Atlas disseminates quality controlled immunopeptidomic information to the public domain and serves as a community resource toward the generation of a high-quality comprehensive map of the human immunopeptidome and the support of consistent measurement of immunopeptidomic sample cohorts.

INTRODUCTION

T cells have the ability to eliminate abnormal cells through recognition of short peptides presented at the cell surface by major histocompatibility complex (MHC) molecules (human leukocyte antigen [HLA] molecules in human). In mammals, cells are decorated by thousands of such peptides, which are collectively referred to as the MHC class I and class II immunopeptidome (1–3). The MHC class I immunopeptidome is composed predominantly of peptides of 8–12 amino acids in length that are mainly found on the surface of virtually any cell- and tissue-type in the body. The MHC class II immunopeptidome is composed of peptides of 10–25 amino acids in length that are mainly found on a subset of professional antigen presenting cells, reviewed in (4,5). The amino acid sequence of these peptides is not random. In fact, individual peptides have the ability to bind MHC molecules via specific anchor residues that define a MHC binding motif (6). Such motifs are generally MHC allelic-specific, thereby limiting the pool of peptides that can be presented on the surface of a specific cell for scrutiny by T cells. In humans, this limitation is countered by the high diversity of HLA alleles. In fact, each individual can express up to six different HLA class I allotypes and typically eight different HLA class II allotypes, and more than 16 700 allelic forms are expressed at the human population level (http://www.ebi.ac.uk/ipd/imgt/hla/stats.html; May 2017). Thus, the composition of the human immunopeptidome is tremendously complex (7). Describing and understanding the complexity of the immunopeptidome and its functional implications is a central and fundamental challenge of immunology with important clinical implications in precision medicine (8).

Mass spectrometry (MS) is a powerful unbiased method to explore the composition of the immunopeptidome (9). Following pioneering work by Hans-Georg Rammensee (10) and Donald Hunt (11) in the early 90’s, the analytical performance of this technique has rapidly evolved and currently enables the identification of thousands of HLA-associated peptides from a single MS measurement (12–22). Notably, the use of MS techniques to conduct ‘immunopeptidomic’ studies has become increasingly popular over recent years, thanks to technical advances and breakthroughs in the field of immuno-oncology (23). As a consequence, huge amounts of immunopeptidomic data have been and continue to be generated at significant expense.

Immunopeptidomics is an expanding field driven by a rapidly growing community of researchers and deep technology platforms. In 2015, a Human Immuno-Peptidome Project (HIPP; https://www.hupo.org/Human-Immuno-Peptidome-Project) was created as a new initiative of the Biology/Disease-Human Proteome Project (B/D-HPP) — a program under the umbrella of the Human Proteome Organization (HUPO) (24). The long-term goal of this initiative is to make the robust analysis of immunopeptidomes accessible to any immunologist, clinical investigator and other researchers by the generation and dissemination of new methods/technologies and informational resources (25–27). Participants in this initiative recognized the need...
for an open immunopeptidomics repository in which output files of mass spectrometric measurements of immunopeptidome samples would be annotated, stored and shared without restriction. Here, we introduce the SysteMHC Atlas project, the first public repository devoted to MS-based immunopeptidomics. In brief, the SysteMHC Atlas uploads raw immunopeptidomics MS data originally deposited into the PRIDE database along with the metadata associated with the experiment (Figure 1) (28). Each project is labeled with the HIPP tag as a B/D-HIPP subproject. Raw MS data are then processed through a uniform computational pipeline for MHC peptide identification, annotation (29) and statistical validation (30,31) (Figure 1B). Lists of MHC peptide ligands as well as sample/context- and allele-specific peptide spectral libraries (32) are generated and presented in the Atlas in a way that they can be searched and browsed by researchers via a web interface. Allele-specific peptide spectral libraries can be further converted into formats that are compatible for uploads into the SWATHAtlas database in order to support immunopeptidomic analyses by SWATH-MS/DIA (Data-Independent Acquisition) methods. Importantly, the SysteMHC Atlas aims to be an open and active repository in which raw MS data can be periodically reprocessed with more advanced informatics tools for peptide identification, statistical validation, HLA peptide annotation and library generation, as these become available to the community—a procedure that has been successfully applied in the field of proteomics to ensure high-quality peptide identification with well-understood false discovery rates (FDR) and quality controls (33). The community is expected to benefit from the SysteMHC Atlas at various levels: (i) basic scientists and clinicians can navigate within a large catalog of high-quality context-specific HLA-associated peptides to gain new insights into the composition of the immunopeptidome, (ii) computational scientists find a rich source of data to develop or test new algorithms for immunopeptidomic analyses and (iii) access to HLA peptide assay spectral libraries facilitates next-generation MS analysis of immunopeptidomes (i.e. SWATH-MS/DIA) (34).

CONTENT AND FUNCTIONALITIES OF THE ATLAS

The first version of the SysteMHC Atlas (February 2017) contains raw and processed MS data derived from 16 published human immunopeptidomics projects/datasets (Figure 2). It also contains information from seven unpublished datasets that were released by the data producers. The number of MS output files per project ranges between 1 and 192 for a total of 1184 raw files. All datasets were generated in data-dependent acquisition (DDA) mode using different instruments and fragmentation methods, including collision-induced dissociation (CID), higher energy collisional dissociation (HCD), electron transfer dissociation (ETD) and electron transfer and higher energy collision dissociation (EThcD) (21). Several laboratories used the spiked-in landmark indexed Retention Time (iRT) peptides for retention time normalization (35,36). Each dataset is labeled with a unique and permanent SYSMHC number. Direct links to PubMed, PRIDE and Immune Epitope Database (IEDB) are also provided if applicable (Figure 2). We briefly describe below the content and functionalities of the SysteMHC Atlas.

A catalog of context- and allele-specific MHC class I and class II peptides

The SysteMHC Atlas contains mainly naturally presented human MHC class I and class II peptides. Natural MHC-associated peptides were extracted by immunofinity purification or mild acid elution from cell lines, primary cells and tissues—i.e. peripheral blood mononuclear cells (PBMCs), T cells, B cells, dendritic cells, macrophages, fibroblasts, colon carcinoma, breast cancer and glioblastoma. All biological sources were HLA typed and peptides from 67 HLA class I and 27 HLA class II alleles are represented in the current version of the database (February 2017). A full listing of all the samples and corresponding metadata (i.e. organism, tissue and cell type, culture conditions, disease state, HLA type, antibody used for immunofinity purification, LC-MS/MS parameters etc.) can be found next to the raw MS files at the project website.

In May 2017, ~29.5 million MS/MS spectra were searched using a uniform and well-tested computational pipeline and yielded 250,768 and 1458,698 distinct peptides with iProphet probability $P \geq 0.9$ and $P > 0.0$, respectively. After applying strict confidence filters for the identification of class I and class II peptides, 119 073 high-confidence HLA class I peptides (peptide FDR 1%, 8–12 amino acids) were identified and annotated to specific HLA-A, -B or -C alleles using an automated annotation strategy as described (34) (see Supplementary Figure S1 for statistics). For class II molecules, 73 465 high-confidence peptides were identified (peptide FDR 1%, 10–25 amino acids, belonging to groups of two or more peptides with an overlap of at least four amino acids). Of note, the assignment of peptides to specific HLA class II alleles will be considered in the future as soon as robust bioinformatics tools for class II peptide annotation become openly available (26). The high-confidence class I and class II peptides were mapped onto 13, 132 and 7704 of the human UniProtKB/Swiss-Prot proteins, respectively.

An important goal in MS-based immunopeptidomics is to assess the size of the human immunopeptidome at the population level. To answer this question, we plotted the cumulative number of distinct HLA class I peptides as a function of the addition of identified spectra at FDR 1% (Figure 3). Each data point on the curve represents an added experiment, and the experiments are presented in chronological order of data acquisition. When looking at the combined data from all HLA class I alleles in the Atlas, our analysis suggests that for the presently available technology the saturation level might already be reached (Figure 3A). However, this observation might be biased given the limited number of HLA alleles as well as the limited number of cell and tissue types that were sampled until now. In addition, when individual HLA class I alleles were considered, the number of distinct peptides continued to steeply increase for several HLA class I allele such as -A02, -C02 and C16 (Figure 3B) indicating that for these alleles, saturation had not yet been achieved as the curves are expected to reach saturation only.
Figure 1. Overview of the SysteMHC Atlas project. (A) The SysteMHC Atlas aims to be a long-term data-driven project that serves the community. It is linked to other repositories of proteomic data and consists of two main components: (i) a uniform computational pipeline for processing raw MS files and (ii) a web interface with storage, searching and browsing capabilities. First, shotgun/DDA-MS experimental data generated for specific projects are submitted by the data producers to PRIDE. Raw MS data are then uploaded into the SysteMHC Atlas and processed through a consistent and open computational pipeline (B) that controls the quality of peptide identification and peptide annotation to specific HLA alleles. Spectral libraries are generated and can be converted into high-quality HLA allele-specific peptide assay libraries, also available at SWATHAtlas. All the results generated by the computational pipeline are made available to the public domain via the SysteMHC Atlas web-based interface, which provides links to the Immune Epitope Database (IEDB) for accessing lists of peptides originally identified and published by the data producers. (B) Current computational pipeline used for generating the immunopeptidome- and spectral database for different HLA allotypes. MS output files generated from several types of instruments are first converted into mzXML file format and then searched using several open-source database search engines. The resulting peptide identifications are combined and statistically scored using PeptideProphet and iProphet within the Trans-Proteomic Pipeline (TPP) (30,31). The identified peptides are next annotated to their respective HLA allele in a fully automated fashion using the stand-alone software package of NetMHCcons 1.1 (29). Spectral libraries are generated using SpectraST (32). Allele-specific peptide spectral libraries are generated from multiple samples—an example for HLA-A03 is highlighted in red. Each HLA peptide is labeled with a unique and permanent library identifier (LibID). Details regarding the computational pipeline and how the data were processed are available at the SysteMHC Atlas website in the ‘ABOUT’ section.
Figure 2. Immunopeptidomics datasets used for building the first version of the SysteMHC Atlas. Data from 23 projects that collectively generated 1184 raw MS files constitute the initial contents of the SysteMHC Atlas. Each project is labeled with a unique SYSMHC identifier and linked to its corresponding PubMed, PRIDE and IEDB ID. For unpublished projects, IDs are not applicable (NA).

when most peptides observable with the applied technology will have been cataloged. Altogether, our current analysis suggests that the Atlas data are not yet comprehensive. In the future, collecting additional MS/MS data from new experiments—including from new HLA alleles, new cell origins, new experimental conditions, new protocols and new MS technologies—will be necessary to properly assess the size and complexity of the human immunopeptidome.

In addition to naturally processed ligands, the SysteMHC Atlas also contains data for synthetic peptides predicted to bind specific HLA alleles. Datasets generated from synthetic peptides might be particularly useful for benchmarking new software tools (37) and to extend the contents of libraries derived from native peptides for targeted analysis of immunopeptidomes (3,38,39). To date, SysteMHC Atlas contains four datasets composed of synthetic peptides: SYSMHC00001 contains data generated from a large collection of synthetic HLA class II peptides encoded by Mycobacterium tuberculosis (Mtb) (34,40); SYSMHC00020, SYSMHC00021 and SYSMHC00022 contain data obtained from synthetic HLA class I peptides encoded by Mtb (41,42), Epstein–Barr virus (EBV) and Homo sapiens, respectively.

The SysteMHC Atlas user interface

Researchers can browse, search and download information using query interfaces available at the website (https://systemhcatlas.org). In particular, the ‘EXPLORE’ link leads to a page where immunopeptidomic data are searchable on numerous levels, including peptide sequence, source protein, as well as HLA class and type. For instance, the user can query the data to specifically identify (i) all class I peptides derived from a specific source protein (e.g. BIRC6 in Figure 4), (ii) the repertoire of peptides presented by a specific HLA type and/or (iii) in which tissues or experimental conditions have specific peptides been observed etc. Thus, the SysteMHC user interface enables large immunopeptidomics datasets to be explored in a user-specifiable fashion.

An important function of the SysteMHC Atlas is to serve as a repository devoted to immunopeptidomics MS-related data at several levels of processing. Specifically, we provide raw and converted mzXML files, iProphet results and HLA peptide spectral libraries, all available for download at the website (Figure 5). In the current version of the Atlas, a total of 539 sample/context- and 37 HLA allele-specific peptide spectral libraries were made available and can be visualized using the respective links from the web interface. Three new allele-specific spectral libraries (i.e. HLA-B15, -C03 and -C07) were also converted into TraML files for SWATH-MS/DIA analysis of immunopeptidomes, as previously described (34,36). These standardized libraries contained the iRT peptides for retention time normalization and data analysis. TraML files are directly available for download at SWATHAtlas.
Figure 3. Cumulative number of MS/MS spectra versus cumulative number of distinct peptides for HLA class I alleles at FDR 1%. (A) All HLA class I peptides were combined. (B) HLA class I alleles that were frequently found in various datasets. Eventually, the curves are expected to reach saturation when most observable peptides will have been cataloged at 1% peptide FDR.
Figure 4. Explore page in the SysteMHC Atlas web-based interface. HLA allele-specific peptide spectral libraries can be downloaded here. The web interface can also be used to query the SysteMHC Atlas and find specific information. (A) As an example the source protein BIRC6 was searched and the Atlas returned back all HLA-associated peptides originating from this protein as well as the context (i.e. SysteMHC ID, Sample ID, iProphet score, HLA annotation score, spectral counts, assigned HLA type and class) in which this peptide was observed. Then, the user can click on a specific Sample ID hyperlink and be redirected to the corresponding raw MS files and metadata (e.g. tissue type, cell type, culture condition, purification method, antibody used, mass spectrometer used etc). (B) The peptide RLLDYVATV was searched and the Atlas returned back the datasets in which this peptide was observed. By clicking on the peptide sequence hyperlink, the user is redirected to a new page in which the LibID information is available for MS/MS spectra visualization. Information can be downloaded as .csv files for further analysis.
Figure 5. Data storage and visualization. To access information about specific datasets, the user selects a specific SYSMHC ID/Project name (e.g. SYSMHC00005) and clicks on ‘view dataset’ at the bottom left of the screen. The samples related to this project are then listed and linked to the number of replicates, organism, tissue, and cell type of origin as well as the HLA typing information (upper panel). The user can then click on a specific Sample ID to visualize the metadata and to download the raw or converted mzXML MS files (red squares). A list of sample-specific HLA-associated peptides can be visualized at 1% peptide-level FDR (green squares). Sample-specific spectral libraries, including consensus fragment ion spectra, can be visualized and downloaded (orange and blue squares). Heat maps (black squares) are used to visualize the annotation of individual peptides to their respective HLA allele (dark blue peptides are predicted to be strong HLA binders according to NetMHCcons).
FUTURE DIRECTIONS

Data sharing, public resources and large-scale/community projects are growing in popularity and necessity in life sciences (43–48), and specifically in proteomics (49,50) where public data sharing is growing exponentially in recent years. Along this line, the SysteMHC Atlas represents the first community-driven resource devoted to collect, store, organize and share large immunopeptidomics datasets generated by MS methods—an important contribution to the Human Immuno-Peptidome Project (25,27). The SysteMHC Atlas will be further developed and enhanced to enable public dissemination of uniform and high-quality immunopeptidome data generated by an open and ever-improving computational pipeline. To this end, raw MS data will be reprocessed periodically using novel high-performance software tools as they are made available to the community. Future software tools are expected to outperform current algorithms for (i) MHC peptide identification, (ii) MHC peptide FDR estimation in large immunopeptidomic datasets and (iii) class I and II peptide annotation to specific HLA alleles, as described (http://www.biorxiv.org/content/early/2017/05/13/098780) (51). In the near future, we aim at providing the necessary tools to retrieve information on post-translationally modified MHC-associated peptides: phosphopeptides, Arg-methylated peptides and proteasome-generated spliced peptides in particular, as those might be of particular relevance for the rational design of immunotherapeutic interventions (52–56). We also plan to identify the potential for large-scale integration and interoperability of all immunopeptidomic data with PRIDE (28), IEDB (57) and SWATHAtlas (34). Thus, we intend the SysteMHC Atlas to become a growing community-driven database and an interoperable, high-performance infrastructure for systematic analysis of terabytes of immunopeptidomic big data. If successful in longer term, we anticipate that the SysteMHC Atlas project will provide key insights to the immunology community and will foster the development of vaccines and immunotherapies against various immune-related diseases such as autoimmunity, allergies, infectious diseases and cancers.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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