FluKB: A Knowledge-Based System for Influenza Vaccine Target Discovery and Analysis of the Immunological Properties of Influenza Viruses

Citation

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Accessibility
FluKB is a knowledge-based system focusing on data and analytical tools for influenza vaccine discovery. The main goal of FluKB is to provide access to curated influenza sequence and epitope data and enhance the analysis of influenza sequence diversity and the analysis of targets of immune responses. FluKB consists of more than 400,000 influenza protein sequences, known epitope data (357 verified T-cell epitopes, 685 HLA binders, and 16 naturally processed MHC ligands), and a collection of 28 influenza antibodies and their structurally defined B-cell epitopes. FluKB was built using a modular framework allowing the implementation of analytical workflows and includes standard search tools, such as keyword search and sequence similarity queries, as well as advanced tools for the analysis of sequence variability. The advanced analytical tools for vaccine discovery include visual mapping of T- and B-cell vaccine targets and assessment of neutralizing antibody coverage. FluKB supports the discovery of vaccine targets and the analysis of viral diversity and its implications for vaccine discovery as well as potential T-cell breadth and antibody cross neutralization involving multiple strains. FluKB is representation of a new generation of databases that integrates data, analytical tools, and analytical workflows that enable comprehensive analysis and automatic generation of analysis reports.

1. Introduction

An estimated 250,000–500,000 people die from seasonal influenza infection each year. The economic impact of influenza is immense due to the large number of lost working hours, hospitalizations, further medical complications, and treatment costs. Although vaccines against influenza exist, the rapid mutation of influenza virus calls for constant surveillance and annual vaccine reformulation [1]. A huge body of sequence data, annotations, and knowledge is available in the literature, online resources, and biological databases such as GenBank [2], UniProt [3], Protein Data Bank [4], EpiFlu Database [5], OpenFlu Database [6], Influenza Research Database (IRD) [7], and the Immune Epitope Database (IEDB) [8]. However, the underlying mechanisms of host/pathogen interaction are still not completely understood. The lack of a “universal” or broadly neutralizing influenza vaccine can be attributed to, among other factors, combinatorial complexity of the host immune system and the highly variable nature of viral antigens leading to
immune escape of the emerging influenza variants [9, 10]. One approach, in an attempt to overcome challenges of immune escape, is to raise a T-cell response against class I or class II epitopes conserved among viral strains [11, 12]. Public databases represent valuable resource for the study and development of broadly protective T-cell vaccines, but our ability to analyze these data falls behind the pace of data accumulation.

Numerous computational analysis tools that are useful for vaccine target discovery are available. They include keyword and text search tools, sequence comparison tools such as the BLAST algorithm [13] or multiple sequence alignment tools such as MAFFT [14], MUSCLE [15], and the Clustal [16], 3D structure visualization tools [17, 18], HLA binding prediction algorithms [19–21], and conservation analysis tools [22, 23], among others. The application of these tools in discrete steps can yield valuable information; however the extraction of higher-level knowledge requires integrating data from multiple databases and employing various analytical tools to answer specific questions. For example, when a new infectious influenza strain emerges (such as H9N7 avian flu [24] or a new seasonal flu) it is desirable to rapidly investigate its similarities and dissimilarities with known sequences, its epidemic or pandemic potential in humans, how different it is from the past vaccine strains, and its T- and B-cell epitopes from previously circulating strains and estimate its immune escape potential. Additionally, for new pandemic strains (such as 2009 swine flu [25]) it is desirable to establish origin and identify strains that are useful vaccine candidates. Well-defined workflows enable rapid extraction of such knowledge and automated generation of reports that contain such information, for which knowledge-based systems have previously been utilized [26, 27]. The need for integration and advanced analysis of available data is rapidly increasing. The integration of multistep analysis of multidimensional data for vaccine analysis and discovery requires the automation of analytical workflows [28].

FluKB is a knowledge-based system that integrates multiple types of influenza data and analytical tools into such workflows to support vaccine target discovery. The datasets in FluKB consist of curated, enriched, and standardized protein sequence data, immunological data from multiple data sources, and a set of modular analysis tools. The analysis tools infrastructure comprises a library of individual tools along with standard (applicable to multiple pathogens) and specific influenza vaccine target discovery workflows. Furthermore, we developed a standardized nomenclature to enable and speed up data mining using automated workflows. FluKB has a user-friendly web-based interface to access the data, tools, predefined workflows, and workflow reports. The overall architecture of FluKB is shown in Figure 1.

2. Materials and Methods

2.1. KB-Builder. FluKB was implemented using the KB-builder framework [29]. Briefly, KB-builder consists of seven major functional modules that enable automated data extraction from multiple sources, data cleaning, import to a central repository, integration of basic analysis tools, integration of advanced analysis tools, workflow definition, and update and maintenance. The KB-builder framework enabled setting up a web-accessible knowledge base and the analysis workflows. A workflow takes the user request, performs complex analyses which combines specific data and analytical tools, and feeds the results into subsequent analyses to produce a comprehensive report. The web-accessible interface uses a set of graphical user interface forms. These interfaces access search routines, analytical tools, and workflows that use a combination of PHP, Perl, Common Gateway Interface (CGI), and C background software. Development of KB-build and FluKB was carried out in CentOS 5.11 Linux environment. The web server used is Apache HTTP server 2.2.3 so the access to the web server is per default parallelized for each user. The Linux server is a 16-core server with 32 GB ram and should be able to handle the traffic to the website.

2.2. Data Sources for FluKB Data Repository. The data repository within FluKB contains four types of data: protein sequence data, HLA-related data (T-cell epitopes and HLA ligands), 3D crystal structures, and neutralizing antibody-related data. Protein sequences available from Influenza A, B, and, C viruses were collected from IRD [7] and GenBank [2], while HLA-related data were collected from IEDB [8]. The complex structures of neutralizing antibodies against influenza virus hemagglutinin (HA) were collected from the Protein Databank (PDB) [4]. The binding and neutralization assays of each neutralizing antibody were collected from primary literature, as described in [30].

2.3. Data Collection, Cleaning, and Enrichment

2.3.1. Protein Data Entries and Their Updates. The following sequence data and annotations were extracted from the IRD: protein sequence, GenBank identifiers (GI and Accession Numbers), UniProt ID, and nomenclature (type, host, location, ID, year, and subtype) when available. The metadata included a vocabulary that comprises the instances of the following terms: protein names, host names, geographic locations, years, and subtypes. The vocabulary was generated from the collected entries and enriched using primary literature. The vocabulary comprised correct terms as well as variants of the terms, and erroneous terms. During updates, each new entry is checked against the list of correct and erroneous terms. If all terms within the entry are matched to the correct terms, the entry is automatically annotated and converted into the FluKB format. If an erroneous term is identified, the curator is alerted and the correct term is proposed for the entry. The curator then approves the change or manually corrects the entry and updates the vocabulary, when needed. Each new term identified in the update dataset is inspected by the curator and added to the vocabulary. The vocabulary iteratively increases in size with each update and less than 20 new terms are usually identified for each update. This approach enables data curation that is of extremely high quality and can be performed very fast (Figure 2 upper path). We converted existing nomenclatures, whenever possible, into standardized formats. To verify the protein annotations of newly added entries, protein
assignment of the strain proteomes was done by aligning the entry proteome sequence to a representative influenza reference sequences using the BLAST algorithm. We selected 10 proteins from UniProt that have detailed annotations as reference sequences for each protein of the two influenza types (A and B). The reference sequences are shown in Supplemental Table S1 (see Supplementary Material available online at http://dx.doi.org/10.1155/2015/380975).

2.3.2. Immunological Data. The immunological data extracted from IEDB include T-cell epitope sequence, epitope type (HLA binding, naturally processed, or T-cell epitope), PubMed references, experimental methods and results, and HLA-allele restriction (Figure 2 lower path). FluKB includes entries from IEDB that bind HLA class I or II allele with at least one positive result, or those reported as T-cell epitopes.

2.3.3. Neutralizing Antibody Data. For each neutralizing antibody, the following descriptive information is provided in the FluKB: isolation information (from human samples or from antibody phage-display libraries) from the primary
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Raw influenza data entries

Initial public data source

Temporary repository

List of metadata terms

Redundancy reduction and data enrichment

Dictionary of standardized items

Vocabulary consolidation for standardized nomenclature

Automated collection of data and annotations

Updates from public sources

Find all IEDB influenza epitopes

Extract epitope files

Select data subsets

Filter data subsets

Populate epitope repository

FluKB

Figure 2: Semiautomated generation and updating of protein sequence repository of FluKB (upper path) and data extraction and repository creation of the influenza specific epitopes from IEDB (lower path).

2.4. Influenza Nomenclature Standardization and Definition of Data Mining Keys. Each influenza strain sample is annotated using the nomenclature originally proposed by the WHO [32] providing a shorthand description of influenza virus strains. However, the lack of a standardized vocabulary has made these nomenclatures error prone. For example, the nomenclature of an influenza isolate of type A, subtype H2N3, isolated from a duck in Heinersdorf in 1986, is written, using the original nomenclature as

A/Peking duck/Heinersdorf/648-4/1986(H2N3)

The lack of standardization of nomenclature has led to inconsistent nomenclature and incomplete metadata, thus increasing the difficulty in extraction of specific data subsets for analysis. The nomenclature in the Heinersdorf 1986 strain 648-4 has two issues. First the organism name term is erroneous (“Peking duck” is a traditional Chinese dish, while “Pekin duck” is the organism). Second, it is unclear where Heinersdorf is located. To ensure a complete access control over the sequence data within FluKB each entry was given a standardized data mining key. This key converts the nonstandardized nomenclature fields into a fully standardized format. The key is represented in FluKB as a standardized FASTA header that provides a condensed and detailed summary of the sample’s information. The FluKB data mining key for the Heinersdorf sample HA protein is


This search key compresses detailed sequence annotations into the FASTA format enabling easy combination of the results of sequence comparison with the analysis of annotations. Standardized formats for host and geographic location enable proper grouping and mapping of results. For host species, the NCBI-Taxonomy IDs [33] and Bird Life International taxonomy (http://www.birdlife.org/datazone/info/taxonomy) names were used as standard terms, including the NCBI taxonomy number. For the geographical locations, two-letter ISO codes for the countries and provinces were used (ISO-3166, 2012) [33]. This allows for each of the host species and each geographic term to be described in nonambiguous terms. Examples of corrected ambiguities are shown in Table 1. The FASTA format is easily understandable because of the descriptive nature of the fields in the FASTA header. Finally we use the term key:yes for all entries that could be fully annotated allowing them to be utilized within the analysis framework of FluKB, while those that could not be fully annotated are assigned with key:no and can only be found by a search.

2.5. Implementation of Analytical Tools. A set of analytical and visualization tools have been integrated within the FluKB. These tools include a selection of keyword searching tools: MAFFT [14] for multiple sequence alignment (MSA) and BLAST [13] for sequence similarity search. Specialized
tools for the analysis of variability include sequence conservation metrics and their visualization using block entropy analysis [34]. The T-cell epitope prediction tools for HLA Class I and Class II have been integrated within FluKB for vaccine-related analyses. WebLogo [22] and BlockLogo [23] tools are used for visualization of results.

2.5.1. Sequence Conservation Metrics. FluKB enables conservation analysis of single positions within protein sequences, of linear blocks of amino acids extracted from multiple sequence alignments (MSA) of proteins using block entropy [34]. In addition, virtual peptides can be constructed from discontinuous epitopes within MSA and can be analyzed using block entropy, enabling the variability analysis of B-cell epitopes [30]. All these calculations are based on the calculation of Shannon entropy [35].

2.5.2. Visualization of Sequence Variability. WebLogo enables fast and easy interpretation of the position specific variability in an alignment. It displays amino acids by their corresponding one-letter code on a graph where all the amino acids present in one position are stacked on top of each other, and the frequency in the position is then based on their individual height in the graph. BlockLogo enables variability analysis of peptides (either linear peptides or discontinuous strings of amino acids), rather than single residues. The combinatorial number of possible blocks that can be created from a WebLogo can quickly become very large because of variation in individual positions. BlockLogo only shows the exact peptides that are most frequent for the investigated positions.

Table 1: All the alternative instances of the host Mallard in all of the entries in the knowledge base of FluKB. The standardized name for the search key is Mallard; Anas platyrhynchos; 8839.

<table>
<thead>
<tr>
<th>Alternative names for Mallard</th>
<th>Number of times present in dataset</th>
<th>Status</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anas domesticus</td>
<td>1</td>
<td>Ambiguous</td>
</tr>
<tr>
<td>Anas platyrhynchos</td>
<td>9</td>
<td>Error</td>
</tr>
<tr>
<td>Anas platyrhynchos</td>
<td>39</td>
<td>Correct</td>
</tr>
<tr>
<td>Domestic duck</td>
<td>58</td>
<td>Variant</td>
</tr>
<tr>
<td>Domestic Mallard</td>
<td>11</td>
<td>Variant</td>
</tr>
<tr>
<td>Feral duck</td>
<td>1</td>
<td>Ambiguous</td>
</tr>
<tr>
<td>Khaki campbell duck</td>
<td>12</td>
<td>Variant</td>
</tr>
<tr>
<td>Mallard duck</td>
<td>25,844</td>
<td>Correct</td>
</tr>
<tr>
<td>Mallard duck</td>
<td>1,172</td>
<td>Redundant</td>
</tr>
<tr>
<td>Pekin duck</td>
<td>137</td>
<td>Variant</td>
</tr>
<tr>
<td>Peking duck</td>
<td>92</td>
<td>Error</td>
</tr>
<tr>
<td>Sentinel duck</td>
<td>13</td>
<td>Error</td>
</tr>
<tr>
<td>Wdk</td>
<td>11</td>
<td>Ambiguous (abbreviation)</td>
</tr>
<tr>
<td>Wild duck</td>
<td>952</td>
<td>Ambiguous</td>
</tr>
<tr>
<td>Total</td>
<td>28,352</td>
<td></td>
</tr>
</tbody>
</table>

2.5.3. Prediction of MHC Class I and II Binders. For class I HLA binding prediction, the NetMHCpan 2.8a algorithm [19] was implemented. The alleles available for predictions are HLA-A*02:01, -A*03:01, -A*11:01, -A*2402, -B*07:02, -B*08:01, and -B*15:01, since prediction accuracies for these alleles are relatively high [36]. Additionally, we predict binders to HLA-A*01:01 since this allele is of very high frequency in the human population, although these predictions are of slightly lower accuracy than the seven benchmarked alleles. Collectively these alleles cover approximately 82% of human population [37]. For class II MHC predictions we used NetMHCIIpan 3.0 [21], with which users can predict binders to HLA-DRB1*01:01, *03:01, *04:01, *07:01, *11:01, *13:01, and *15:01, as these have been benchmarked and validated for high accuracy [21]. These alleles cover approximately 40% of the human population [37].

2.5.4. Analysis of Sequence Similarity and Geographical Mapping (Strain Mapper). The standardized data mining key in FluKB enables sequence similarity searching and the display of a sequence's origin on the world map. For this purpose, we developed the specialized tool, Strain Mapper. The query sequence is entered in the search window and optionally the maximal number of amino acid mismatches can be selected. We used Google Earth software (http://www.google.com/earth/) for geographic map display. It is combined with sequence similarity analysis to allow user to select a query sequence and find its closest matches in FluKB and visualize them on the world map. This feature provides visualization of epidemiological information useful for evaluating the spread of a given virus and closely related variants.

3. FluKB Database

3.1. Data Repository. The FluKB sequence repository as of December 2014 contains 402,306 sequence entries from 75,426 unique strains of influenza. There are 67,907 type A, 70,28 type B, 194 type C, and 297 unknown type sequence entries. Out of 330,435 full-length sequences and 71,501 fragments, 370 protein sequences failed to align well to any of the template alignments during the annotation process. Each sequence entry contains information about location, host, and time of isolation, as well as a standardized nomenclature for identification of strains. Each entry contains a protein sequence with standardized, curated, and enriched annotations (Table 2).

The epitope repository contains a total of 357 unique T-cell epitopes (194 class I and 163 class II) and 685 unique HLA binders (572 class I and 113 class II). Each record describes the type of epitope (T-cell epitope, naturally processed, or HLA binder), epitope sequence type (only exact epitopes are included in the repository), experimental method used for validation, HLA restriction, and literature references.

Twenty-eight neutralizing antibodies against influenza virus have crystal structures of HA/antibody complexes available in PDB. The functional data and neutralizing specificity of these antibodies were collected from published articles. Twenty of these antibodies target the globular head of the HA
3.2. Data Cleaning, Quality Control, and Enrichment. The sequence data collected from IRD were subject to extensive cleaning, quality control (QC), and enrichment of annotations. We found that 142,232 (38.25%) of the 402,306 entries contained at least one type of error, ambiguity, or missing data. Most errors were in the geographic location fields where 72,340 (17.9%) had an error and 6,821 (12.1%) had missing information in the entry (see Table 3). In the initial screen of the data we found 2,977 entries that did not conform to nomenclature standard, including 305 entries that lacked information about host species, 867 entries that lacked separation fields within the nomenclature, and 1,805 other deficiencies. These entries were manually corrected and their nomenclatures were updated.

Furthermore, abbreviations, alternative, and misspelled names constituted the largest proportion of errors and were present in more than 10,000 entries. All name-related errors were corrected by the dictionary consolidation using the dictionary of standardized metadata terms. An example of the redundancy is shown in Table 1 where the host “Mallard” is found in 16,457 of the FluKB entries described by 14 different terms. In total, 96.41% of errors of various types described above were corrected and 469 standardized forms of missing data (such as location and host species) were added by manually searching the original literature.

We plan to have yearly updates of FluKB moving forward as new data and tools become available.

### Table 2: The data fields in each protein entry.

<table>
<thead>
<tr>
<th>Field title</th>
<th>Field content</th>
</tr>
</thead>
<tbody>
<tr>
<td>GenBank GI</td>
<td>Accession number unique to GenBank</td>
</tr>
<tr>
<td>NCBI accession</td>
<td>Accession number unique to NCBI</td>
</tr>
<tr>
<td>CVC accession</td>
<td>Accession number unique to FluKB</td>
</tr>
<tr>
<td>Protein name</td>
<td>The protein name, based on template BLAST</td>
</tr>
<tr>
<td>Sequence type</td>
<td>Full or fragment of the protein sequence</td>
</tr>
<tr>
<td>Predict HLA binders</td>
<td>Predict HLA binders to the sequence</td>
</tr>
<tr>
<td>Blast sequence</td>
<td>Blast the sequence for similar sequences of FluKB</td>
</tr>
<tr>
<td>Sequence</td>
<td>The amino acid sequence of the entry</td>
</tr>
<tr>
<td>Epitopes in sequence</td>
<td>IEDB epitopes found in the protein sequence</td>
</tr>
</tbody>
</table>

### Table 3: The types of errors found for the geographical location in the metadata of the entry.

<table>
<thead>
<tr>
<th>Type of error</th>
<th>Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Case errors</td>
<td>26,335</td>
</tr>
<tr>
<td>Redundant information</td>
<td>14,165</td>
</tr>
<tr>
<td>Alternative name and abbreviations</td>
<td>11,208</td>
</tr>
<tr>
<td>Misspellings</td>
<td>2027</td>
</tr>
<tr>
<td>Alternative spellings</td>
<td>9,112</td>
</tr>
<tr>
<td>True location could not be determined</td>
<td>9,493</td>
</tr>
<tr>
<td>Total</td>
<td>72,340</td>
</tr>
</tbody>
</table>

3.3. Standardized Nomenclature. To enable automated data mining and workflows, we created data mining keys from the original nomenclature of influenza viruses with standardized terms. The data mining keys utilized NCBI’s taxID database for host species [32] and the ISO codes for geographical location (ISO-3166, 2012) [33]. A total of 398,078 sequences (98.95%) were assigned the new developed nomenclature, while 4,228 (1.05%) could not be assigned. The original standard nomenclature is included in the data mining key as a reference for additional literature searches and text mining of article databases [37]. Data mining for vaccine targets often requires the analysis of subsets of data, for example, patient data such as specific HLA profile, age group, phenotypes, or other factors. Similarly, epidemiological modelling may need analysis of sequences from certain hosts, for example, specific migrating birds, or limited to geographical locations. The host, time, and location of collection are key information that help determine the spread of specific influenza strains and are central for better understanding of influenza outbreaks [38]. The data mining keys enable such analyses by having a standardized nomenclature, which pattern recognition algorithms can utilize as labels. Entries without the data mining key are made unavailable to the analysis on FluKB as inclusion of entries that lack data could affect the reliability and the outcome of the results. The data mining keys are furthermore nomenclature crucial for the automation of computational analyses; standardization of nomenclature fields allows the computer to interpret the data automatically, which previously was limited. For instance, the taxonomy ID of hosts enables host specific
analyses that can potentially reveal features important for interspecies transmission of influenza. Proper organization of data allows for grouping of data by ancestral species and the variability can be followed over time. Furthermore the ISO codes for the geographical location by country and provinces enhance analyses in, for instance, epidemiological studies where an increased resolution in terms of actual spread can be analysed. This information can be used for the analysis of changes in T-cell and B-cell epitopes.

4. FluKB Tools

4.1. Database Searching and Querying. In FluKB, two search strategies can be deployed for sequence search: annotation-based or epitope-based. The first is a keyword search that enables the user to extract the data of FluKB into specific subsets and the second is a sequence similarity search by BLAST [13]. These search types are vital for the following data analysis as they enable the user to select the needed datasets based on specific scientific questions.

4.1.1. Keyword Search. The user can query the sequence entries for information such as the type, protein, subtype, year range, country, province, host, original nomenclature, and sequence type (fragment or full protein) by keyword search. The sequence entry database is indexed in order to decrease the retrieval time. An example of entry page retrieved by ID FLU0306481 or Strain Name A/Guangdong/1/2013, protein HA, is shown in Figure 3.

4.1.2. Sequence Similarity Search. FluKB has an indexed database generated from sequence entries that can be searched for sequence similarity using BLAST algorithm. The standard parameters are used: E value (10), word size (≥2), substitution matrix (BLOSUM62), gap cost (11) and extension (1), size of the result list (500), and pairwise list (250). Besides sequence search, the FluKB entries can be searched for T-cell epitopes and B-cell epitopes.

4.1.3. Stand-Alone Tools. FluKB can be queried using a selection of analytical tools under the tab “Tools.” Sequence alignment by MSA can be performed under the “Sequence alignment” tab by entering either a list of sequence IDs in the query window or a selection of subsets of proteins by name, influenza type, subtype, range of years of identification, country, province, host, or complete sequences/fragments. The protein subsets selection window (Supplemental Figure S1) can be used for sequence variability analysis and block entropy calculation under appropriate tabs. Epitope block entropy (T-cell epitopes) uses protein subsets selection window that enables the input of epitope sequence. The strain mapping tool is described in Section 2.5.4.

4.2. Variability Analysis. The analysis of variability of viral sequences is important for understanding the emergence of new strains, immune escape, changes in pathogenicity, the extent of spread of viral strains, and vaccine design. The variability analysis can be performed interactively, but the variability analysis tools are also integrated in the T-cell and B-cell mapping tools and relevant workflows. The interactive variability analysis can be performed using individual sequences or sets of sequences. The tools used in variability analysis are shown in Table 4. The main tools for the analysis of variability are BLAST search that can be accessed from individual entries (Figure 3, “find similar sequences”) or from the “Sequence alignment” under the “Tools” tab. The MSA can be performed from the results of BLAST search by selection of sequences and clicking the “Align them...” key. The positions of variability within the MSA results are color-coded for better visual inspection and each sequence is hyperlinked to its record (Supplemental

<table>
<thead>
<tr>
<th>Tool</th>
<th>URL</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>MAFFT MSA</td>
<td><a href="http://mafft.cbrc.jp/alignment/software/">http://mafft.cbrc.jp/alignment/software/</a></td>
<td>[12]</td>
</tr>
<tr>
<td>NetMHCpan 2.8a</td>
<td><a href="http://www.cbs.dtu.dk/services/NetMHCpan/">http://www.cbs.dtu.dk/services/NetMHCpan/</a></td>
<td>[15]</td>
</tr>
<tr>
<td>NetMHCIIpan 3.0</td>
<td><a href="http://www.cbs.dtu.dk/services/NetMHCIIpan/">http://www.cbs.dtu.dk/services/NetMHCIIpan/</a></td>
<td>[19]</td>
</tr>
<tr>
<td>Block entropy</td>
<td><a href="http://research4.dfci.harvard.edu/cvc/flukb/HTML/blockentropy.php">http://research4.dfci.harvard.edu/cvc/flukb/HTML/blockentropy.php</a></td>
<td>ISO-3166</td>
</tr>
<tr>
<td>BlockLogo</td>
<td><a href="http://mafft.cbrc.jp/alignment/software/">http://mafft.cbrc.jp/alignment/software/</a></td>
<td>[21]</td>
</tr>
</tbody>
</table>

Table 4: The sources of the integrated tools in FluKB and URL for their stand-alone versions web services.

FIGURE 3: Record FLU0306481, Protein HA representing the 2013 H7N9 outbreak of bird flu in China.
Figure 4: T-cell epitope EPI102 and the T-cell epitope analysis from the record FLU0021633. (a) Entry EPI150 with the graphical display of present T-cell epitope variants. (b) Graphical display of predicted T-cell epitopes (HLA-B*07:02, length 10, IC_{50} < 1000 nM).

The “Sequence variability analysis” tool plots entropy (red curve) and the percentage of sequences (blue curve) containing the consensus amino acid at all positions along with the consensus sequence (Supplemental Figure S3). Further visualization can be achieved by “Block entropy” calculation, which visualizes the conservation of peptides of lengths appropriate for immune recognition, rather than individual residues [34]. The “Epitope block entropy” calculation displays variability of a specific epitope across the selected subset of sequences.

4.3. T-Cell Epitope Search. T-cell epitope analysis can be performed directly from the protein entry. For example, three T-cell epitope entries are displayed on the record entry page (Figure 3). The epitope entry page for EPI150 is shown in Figure 4(a). In addition, the prediction of T-cell epitopes can be performed by selecting allele and peptide length in the “Predicted HLA Binders” field followed by the “Submit” action. The visual display of experimentally verified T-cell epitopes is shown in Figure 4(b).

T-cell epitope search can also be initiated from the “Search” tab, where epitope can be searched by the sequence. The results will appear as a list of epitopes along with their binding or T-cell restriction specificities. The list is hyperlinked to the epitope record, an example of which is shown in Figure 4(a). Finally, T-cell epitope search can be performed using the workflow titled “Vaccine targets” under the Workflows tab. After selection of input parameters, for example, Allele “HLA-A*0201,” Protein “HA,” Influenza type “A,” subtype “H7 N9,” year(s) “2013 2014,” Affinity threshold “500 nM,” and Conservation Threshold “95%” (and remaining values “default”), 98 sequences will be selected and the report will be generated.

4.4. B-Cell Epitope Search. B-cell epitope analysis can be performed from the Search tab by selecting “Antibodies list.” By the end of May, 2014, 28 antibodies and their detailed neutralizing and structural information have been deposited in FluKB. All of these antibodies are neutralizing antibodies against hemagglutinin protein on influenza virus. These antibodies are listed on the webpage, while their respective B-cell epitopes can be displayed on three interactive structures: X-31 strain-specific antibodies on HA structure from 1KEN, broadly neutralizing antibodies on HA structure from 1EO8, and influenza B virus antibodies on HA structure from 4FQK. This feature enables visual comparison of antibody-specific B-cell epitopes.

For each neutralizing antibody, the isolation information, structure information, and computational identified B-cell epitope information can be accessed. Also, the neutralized motifs and escape motifs extracted from experimentally validated strains from the primary literature are presented as well [30]. In addition, two workflows have been implemented for further analysis: the neutralization coverage estimation and B-cell epitope mapping (Supplemental Figure S4). The neutralized/escape coverage by a specific existing neutralizing antibody is calculated for the complete population of influenza strains. The strain population coverage by a neutralizing antibody can be assessed within any selected subset of influenza strains, such as year range, specific subtype, and geographic coverage. The B-cell epitope mapping is performed by submitting a query hemagglutinin sequence. Cross neutralization coverage of a known neutralizing antibody can be estimated based on sequence comparison to the known neutralizing epitopes. A discontinuous peptide is extracted based on epitope positions determined from crystal structures. This tentative discontinuous peptide is then compared
Figure 5: The results of B-cell epitope analysis II. Analysis of entry FLU0099373 for broadly neutralizing antibody F10 interaction. (a) The BLAST result of the query sequence to the sequence with highest identity in FLUKB, (b) the discontinuous peptide extracted from the query sequence and respective residue positions highlighted in full sequence, and (c) the summary information of neutralizing antibody. The discontinuous peptide from the query sequence is compared to all discontinuous peptides generated from FLUKB (with their neutralizing status listed). The identical sequence is highlighted in yellow with estimated status of neutralization given.

to the B-cell epitopes of experimentally validated strains. An example of B-cell epitope analysis is shown in Figure 5.

FluKB offers the capability to address complex questions relating to sequence variability on very specific subsets, identification of potential T-cell epitopes, and selection and combination of these epitopes into polyvalent vaccine constructs. The modular structure of the workflow renders FluKB highly flexible. The tools and data can be reorganized and more tools can be created to answer additional questions, for example, relating to epidemiological modeling and analysis of cross protective potential of neutralizing antibodies. The overall architecture can be viewed in Supplemental Figure S5.

5. Conclusion

Publically available influenza data are a valuable resource for computational analyses with applications in vaccine design. Similarly, existing bioinformatics tools provide the means for extraction of information and new knowledge. However, to utilize the full potential of these resources, data preprocessing must be performed and analytical tools must be carefully combined into well-defined workflows. These workflows allow users to ask specific questions (scientific, technical, and clinical) and provide means for systematic data analysis.

These workflows can automatically generate comprehensive analysis reports. The infrastructure of data and tools is the backbone of FluKB and similar knowledge-based systems [26, 27].

Despite many years of research and available vaccines, influenza remains a major public health burden and a threat of a major new pandemic. Multiple data sources provide information on protein and nucleotide sequences and immune epitopes in influenza [2, 5–8, 39, 40]. They represent well-maintained catalogues of influenza sequences and annotations, along with a selection of basic search tools. They focus mainly on providing access to data, extraction, and simple analyses. The FluKB was developed focusing on a different purpose, the facilitation of data mining for influenza vaccinology and immunology of influenza infection. The FluKB has very clean and standardized data, integrating information on antigen sequences, and immunological epitopes. The set of integrated analysis tools and workflows are designed to aid rational vaccine design. This includes the discovery of vaccine targets, assessment of variability, and in-depth analysis of immune epitope. FluKB is a unique data mining system for largely automated knowledge discovery from the ever-increasing body of influenza data with applications in both T-cell and B-cell immunology and vaccinology.
Systematic discovery of influenza vaccine targets requires highly accurate, up-to-date, and standardized data of influenza antigens and immune epitopes. The sequence and epitope data available through publications, various reports, and databases vary in quality, granularity, and data formats. The extraction of knowledge and discovery of vaccine targets from diverse and scattered data sources are a challenging and time-consuming task. FluKB integrates the content and the analytical tools in a unified system that enables the automation of complex queries and discovery. FluKB is a contribution to the long-standing quest for universal the automation of complex queries and discovery. FLUKB and the analytical tools in a unified system that enables from diverse and scattered data sources are a challenging and time-consuming task. FluKB integrates the content and the analytical tools in a unified system that enables the automation of complex queries and discovery. FluKB is a contribution to the long-standing quest for universal the automation of complex queries and discovery.

Conflict of Interests
The authors declare that there is no conflict of interests regarding the publication of this paper.

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