PDBe-KB: a community-driven resource for structural and functional annotations

PDBe-KB consortium*†

Received August 14, 2019; Revised September 11, 2019; Editorial Decision September 20, 2019; Accepted October 01, 2019

ABSTRACT
The Protein Data Bank in Europe-Knowledge Base (PDBe-KB, https://pdbe-kb.org) is a community-driven, collaborative resource for literature-derived, manually curated and computationally predicted structural and functional annotations of macromolecular structure data, contained in the Protein Data Bank (PDB). The goal of PDBe-KB is two-fold: (i) to increase the visibility and reduce the fragmentation of annotations contributed by specialist data resources, and to make these data more findable, accessible, interoperable and reusable (FAIR) and (ii) to place macromolecular structure data in their biological context, thus facilitating their use by the broader scientific community in fundamental and applied research. Here, we describe the guidelines of this collaborative effort, the current status of contributed data, and the PDBe-KB infrastructure, which includes the data exchange format, the deposition system for added value annotations, the distributable database containing the assembled data, and programmatic access endpoints. We also describe a series of novel web-pages—the PDBe-KB aggregated views of structure data—which combine information on macromolecular structures from many PDB entries. We have recently released the first set of pages in this series, which provide an overview of available structural and functional information for a protein of interest, referenced by a UniProtKB accession.

INTRODUCTION
Since 1971, experimentally determined 3D structures have been deposited to the Protein Data Bank (PDB)—the single global archive for macromolecular structures (1). As of August 2019, the PDB contains more than 150,000 entries, referencing over 47 500 unique protein sequences in the Universal Protein Resource (UniProt) (2), with ~12 000 new PDB structures added each year. The continuous improvement of experimental methods drives the expansion of the protein structural space covered by known structures. Ultimately, the goal of structure determination is to gain insights into the function of macromolecules (3), and to advance this goal, it is essential to place structural data in a biological context (4).

The wealth of structural data from the PDB is utilised by hundreds of data resources and scientific software. Many of these resources add valuable annotations and thereby, enhance the biological context of macromolecular structures. Such annotations include catalytic sites (5), ligand binding sites (6–8), molecular channels (9), post-translational modification sites (10,11) and other functional sites (12–14), context-dependent roles of small molecules (15,16), effects of genetic variability or mutations (17,18), dynamical properties and flexibility of biopolymer chains (19) and other biophysical parameters (20,21). Currently, the impact of these valuable annotations is limited by the following three factors: (i) the data is fragmented over a large number of distinct resources, each with its own data structure, formats and access mechanisms, making it difficult to compare or aggregate even similar types of annotations; (ii) many of these specialist resources typically reach only a comparatively small section of the scientific community; (iii) even expert users may not be aware of the full extent of the expanding ecosystem of these resources (22).

In an effort to align the management of these valuable data with the FAIR principles of Findability, Accessibility, Interoperability and Reusability (23), we have launched in 2018 the Protein Data Bank in Europe-Knowledge Base (PDBe-KB, pdbe-kb.org), a community-driven, collaborative resource, whose aim is to place macromolecular structures in their biological context by bringing together the various resources providing pieces of this context (Figure 1).

To facilitate this data integration, PDBe-KB partners have defined a common data exchange schema and format (available at https://gitlab.ebi.ac.uk/pdbe-kb/funpdb/funpdb-schemas) for functional annotations of PDB data. The schema focuses on the commonalities of the annotations, capturing the minimal required information that can describe them, and provides links to more comprehensive views of the data hosted by the contributing partners, allowing users to explore the complete data available at the specialist data resources. This arrangement ensures that PDBe-KB remains scalable and maintainable while also increasing
Traditionally, a PDB entry represents structures based on a single set of experiments, where each structure may represent only a segment of the full-length protein. However, PDB entries that describe the structure of the same protein are not interconnected. Furthermore, there is a rich ecosystem of resources and scientific software providing added value annotations based on the structures archived in the PDB, and when combined, these annotations provide evidence for the biological context of the protein. Therefore, the aim of PDBe-KB is to integrate these annotations and interconnect the various PDB entries in order to provide comprehensive, aggregated views of biologically meaningful entities, such as full-length proteins.

In 2018, PDBe-KB launched a deposition system accepting annotations from partners, and an infrastructure to store and expose these data via a distributable database and an Application Programming Interface (API). Furthermore, in March 2019, PDBe-KB introduced a website for accessing and displaying the contributed annotations and PDB data. To date, 18 partner resources from 8 countries collectively contributed over 520 million manually curated or predicted residue-level annotations for PDB structures (Table 1).

PDBe-KB is managed by the Protein Data Bank in Europe team (PDBe; pdbe.org) (25) at the European Bioinformatics Institute. PDBe-KB is also one of the activities of the ELIXIR 3DBioInfo community (26), which aims to further integrate annotations and bioinformatics data related to macromolecular structure and to act as a forum for structural bioinformatics resources and research teams. PDBe-KB partners have agreed resource governance guidelines (http://pdbe-kb.org/guidelines), whose key points are summarised below: (i) the data contributed to PDBe-KB is free from any restrictions on distribution and re-use; (ii) the partners are responsible for the quality of the data they contribute; (iii) protocols for data generation must be described in peer-reviewed publications; and (iv) in case of predicted/calculated annotations, the contributing partner commits to submit data updates at least once a year, i.e. to provide annotations for newer PDB entries and/or to update the existing annotations when the underlying algorithms change. Literature-based manually curated resources may contribute data updates less frequently. PDBe-KB convenes annual general meetings where these guidelines are reviewed, new/prospective members present their work, and the overall progress and plans are discussed. New areas of future research are also identified with regards to gaps in annotations, improved data visualization and benchmarking of computational methods that provide annotations.

**IMPLEMENTATION**

The PDBe-KB infrastructure and workflow has two main parts (Deposition and Access) and can be divided into five components, described below and schematically depicted in Figure 2: (i) the data exchange JSON schema; (ii) the data deposition and validation system; (iii) the graph database that stores the aggregated data; (iv) the API enabling internal and external data access and (v) the reusable web components for data visualization.

**Data deposition**

To facilitate data transfer and enhance interoperability, the consortium members have collaboratively designed a lightweight data exchange schema, available at https://gitlab.ebi.ac.uk/pdbe-kb/funpdbe/funpdbe-schema. The schema is a JSON specification and it defines the mandatory and optional fields required for deposition. The schema represents PDB entries hierarchically, with entry information at the root, and chains, and residues on subsequent levels. Residues may have multiple associated annotations, grouped together in ‘sites’ or ‘residue groups’ which provide labels for residues that belong together. For example, certain residues may together compose a druggable pocket (7), where each residue would have its distinct associated probability score, yet would be referenced to the same site, labelled ‘pocket 1’.

It is mandatory to provide residue-level annotations using the PDB author residue identifiers (i.e. the residue numbering provided by the original authors of the PDB entry).
Table 1. Partner resources contributing annotations to PDBe-KB

<table>
<thead>
<tr>
<th>Partner resource (Reference)</th>
<th>Resource leader</th>
<th>Type of annotations</th>
<th>Number of PDB entries</th>
</tr>
</thead>
<tbody>
<tr>
<td>COSPI-Depth (21)</td>
<td>M. S. Madhusudhan</td>
<td>Residue depth</td>
<td>141 097</td>
</tr>
<tr>
<td>P2rank (6)</td>
<td>D. Hoksza</td>
<td>Binding site predictions</td>
<td>138 892</td>
</tr>
<tr>
<td>Arpeggio (15)</td>
<td>T. Blundell</td>
<td>Ligand interactions</td>
<td>117 023</td>
</tr>
<tr>
<td>3DComplex (14)</td>
<td>E. D. Levy</td>
<td>Interaction interfaces</td>
<td>111 355</td>
</tr>
<tr>
<td>DynaMine (19)</td>
<td>W. Vranken</td>
<td>Backbone flexibility predictions</td>
<td>98 548</td>
</tr>
<tr>
<td>POPSCOMP (20)</td>
<td>F. Fraternali</td>
<td>Solvent accessibility</td>
<td>77 578</td>
</tr>
<tr>
<td>AKID (11)</td>
<td>M. Helmer-Citterich</td>
<td>Kinase-target predictor</td>
<td>41 492</td>
</tr>
<tr>
<td>ChannelsDB (9)</td>
<td>R. Svobodova</td>
<td>Molecular channels</td>
<td>25 351</td>
</tr>
<tr>
<td>CATH-FunSites (13)</td>
<td>C. Orengo</td>
<td>Functional site predictions</td>
<td>23 975</td>
</tr>
<tr>
<td>canSAR (7)</td>
<td>B. al-Lazikani</td>
<td>Druggable pocket predictions</td>
<td>17 804</td>
</tr>
<tr>
<td>FoldX (17)</td>
<td>L. Serrano</td>
<td>Energetic consequences of mutations</td>
<td>3778</td>
</tr>
<tr>
<td>ProKinO (10)</td>
<td>N. Kannan</td>
<td>Curated regulatory sites</td>
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</tr>
<tr>
<td>14–3-3-Pred (12)</td>
<td>G. Barton</td>
<td>Binding site predictions</td>
<td>1941</td>
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<tr>
<td>CaMKinet (in preparation)</td>
<td>M. Kumar</td>
<td>Curated PTM sites</td>
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<tr>
<td>M-CSA (5)</td>
<td>J. Thornton</td>
<td>Curated catalytic sites</td>
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</tr>
<tr>
<td>3DLigandSite (8)</td>
<td>M. Wass</td>
<td>Binding site predictions</td>
<td>910</td>
</tr>
<tr>
<td>Missense3D (18)</td>
<td>M. Sternberg</td>
<td>Mutations in Human Proteome</td>
<td>0*</td>
</tr>
<tr>
<td>MetalPDB (16)</td>
<td>A. Rosato</td>
<td>Curated metal binding sites</td>
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<tr>
<td>ELM (24)</td>
<td>T. Gibson</td>
<td>Short linear motifs</td>
<td>0*</td>
</tr>
</tbody>
</table>

PDBe-KB integrates data contributed by partner resources who provide a wide array of functional and biophysical annotations for PDB structures. Transfer of annotations is in progress.

to map annotations to residues. In case of predicted annotations, the raw scores and confidence scores are also required. The depositors are encouraged to provide URLs linking to the original, complete datasets of which the deposited annotations are often a subset, although in some cases a generic URL to the resource or software can be provided.

The schema is maintained by the PDBe team and is updated according to the requirements of the consortium, tracking the changes in a changelog file. The repository of the schema also contains example JSON files that can serve as templates for prospective collaborators.

Deposition of annotations to PDBe-KB starts with discussions between new partners and PDBe, and after agreeing to the PDBe-KB guidelines, the prospective contributors convert their data to the PDBe-KB data exchange format described above. To facilitate the data conversion, a standalone Python package is provided to check if these files comply with the schema specification, as well as to perform all the consistency checks that the server-side deposition pipeline would perform. The package is available at https://gitlab.ebi.ac.uk/pdbe-kb/funpdbe/funpdbe-validator.

After the annotations are converted to compliant JSON files, they are transferred to private FTP areas provided
by EMBL-EBI, and that serve as the authoritative sources for the annotations. The PDBe team runs an internal high-throughput validation and data processing pipeline weekly to perform all the checks and post-processing necessary for integrating the submitted annotations into the PDBe-KB graph database, described in the next section.

Data access

One of the major advantages of building a knowledge base such as PDBe-KB is the co-location and interconnectedness of diverse types of information in a single environment. To capitalise on this advantage, all the annotations provided by PDBe-KB partners are integrated with the core PDBe data in a graph database powered by Neo4j (https://neo4j.com). Each PDB structure is represented as a tree in this graph, with the root being the PDB entry, connected to entities (unique molecules) and chains (instances of these molecules). Entities and chains are connected to residues. Where the mapping is available through the SIFTS data (27), PDB residues are connected to the corresponding residues in UniProtKB sequences. PDB residues (∼150 million) are also linked with all the available PDBe-KB annotations, e.g. whether the given residue is part of a catalytic site or belongs to a macromolecular interaction interface.

Integrating PDB and PDBe-KB data as a graph allows the straightforward transfer of annotations from PDB entries to both directly mapped and to highly similar (90% identity) UniProtKB sequences and by extension to other PDB entries connected to those UniProt accessions. Weekly snapshots of the graph database index are made available on the PDBe FTP area at ftp://ftp.ebi.ac.uk/pub/databases/msd/graphdb/. The compressed snapshot required 102 Gb of storage (∼500 Gb uncompressed) as of August 2019.

To ensure consistent and robust access to all the PDBe-KB data stored in the graph database, the PDBe team has developed a REST API (implemented in the Flask framework for Python - https://palletsprojects.com/p/flask/), which contains 50 public endpoints. Detailed documentation and examples for each endpoint are available at https://pdbc-ke.org/api. This API powers the new PDBe-KB web components and pages and is also used by PDBe query pages, PDBe query system, and has already been integrated into some of the other EBI resources.

Web components

Clear and intuitive visualisation of structural data is crucial for conveying scientific information to the diverse user community. PDBe-KB presents the core PDB data and the biological context annotations on novel aggregated views for structural data. The first set of such views focus on full-length proteins (UniProtKB accessions). These views are assembled from modular, reusable and customisable web components which retrieve the data via a REST API (see above in the ‘Data access’ section). These components are implemented in the Angular 6+ framework (https://angular.io) and include textual and pictorial data representations and interactive visualizations. Specifically, to display PDBe-KB annotations we use the new version of the ProtVista sequence feature viewer (28) (Figure 3A), which is a visualization tool co-developed by UniProt (29), InterPro (30) and PDBe (25) (available at https://github.com/ebi-webcomponents/nightingale) and, in order to display annotations in 3D, we use the macromolecular viewer LiteMol (31) (Figure 3B, available at https://www.ebi.ac.uk/pdbe/pdb-component-library).

AGGREGATED VIEWS OF PROTEINS

To display the functional annotations and structural data in a useful manner, we have developed a set of novel web pages (aggregated views) that provide an overview of all the structure data related to a full-length protein (UniProtKB accession), as opposed to the conventional pages that focus on single PDB entries. These novel views were designed following extensive gathering and analysis of user requirements and were launched in March 2019.

The aggregated views for proteins contain seven sections, which aim to answer specific scientific questions: (i) summary: how many PDB entries map to the UniProtKB accession? How many ligands and macromolecular partners are observed in the PDB interacting with the protein in question? How many types of annotations and how many publications are available? Which PDB entries contain representative structures of the protein, and what do these representative structures look like?; (ii) structural coverage and domains: what PDB entries are available for a given protein and their mapping onto to the UniProt sequence, and what domains are present in these entries?; (iii) small molecules: which small molecules are observed to either interact directly with the protein in question or are observed in PDB entries containing it?; (iv) macromolecular interactions: which macromolecules does the protein interact with according to the available data in PDB entries?; (v) functional annotations: what are the annotations deposited by PDBe-KB partner resources for the protein?; (vi) similar proteins: For proteins with structures available in the PDB, a list of similar proteins, with high sequence identity, from UniProtKB without any structure data and if there are no data in PDB for the protein of interest, are there proteins with high sequence identity that have structural data available in the PDB? and (vii) publications: which are the primary publications citing the PDB entries related to the protein? What reviews are available either based on PDB or UniProtKB accessions for this protein?

Summary and structural overview

The summary section provides an overview of all the available data using icons that also serve to navigate between the various sections (Figure 4A). Basic information such as the gene name, species and biological function of the protein is provided by the Proteins API (32). Known enzymes and proteins implicated in diseases are labelled accordingly. The section also provides a gallery of representative structures available for the various segments of the full-length protein. These representative PDB structures are selected based on sequence coverage, resolution and data quality (33). Structural studies of large proteins tend to yield multiple structures of shorter (often non-overlapping) segments of the primary sequence that are described in different PDB entries.
Figure 3. The aggregated protein views are built using several web components, the main components being ProtVista and LiteMol. ProtVista (A) is a sequence feature viewer co-developed by UniProt, PDBe and InterPro. It can be used to display residue-level information mapped to sequences. Our implementation of LiteMol (B) is a lightweight molecular viewer wrapped into a reusable web component that allows the visualisation of biological assemblies, complexes and ligand binding sites.

This section aims to simplify the navigation between the various entries. Clicking on the images of any of the representative PDB structures launches an instance of the interactive 3D visualization tool LiteMol (31).

The Structures and Domains section provides an overview of all (i.e. not just the representative) PDB entries available for the protein and displays which region of the protein these PDB entries cover (Figure 4B). It also indicates which residues are modeled with atomic coordinates (i.e. observed), and which ones are not (i.e. unobserved), and allows identification of PDB entries containing particular domains from Pfam (34), CATH (35) and SCOP (36). Additionally, it displays the secondary structure content of the representative PDB chains and predictions of intrinsic disorder provided by MobiDB (37). All this information is displayed using an implementation of the ProtVista sequence feature viewer (28).

Galleries and annotations

The ligands and environments section displays a gallery of all the unique small molecules that are observed to be interacting with the protein of interest in any of the PDB entries (Figure 5A), as well as any small molecules observed in the same PDB entry as the protein of interest (i.e. not necessarily interacting directly). This gallery can be filtered using molecule names, three-letter molecule (chemical compound) identifiers and PDB identifiers, and clicking on each
Figure 4. The first two sections of the aggregated views of proteins provide an overview of all the data available in PDB for a protein of interest. The view includes the number of interacting small-molecules, macromolecular interaction partners and functional annotations, as well as the number of publications related to the PDBs, mapped to the protein (A). It also offers visual help for identifying all the PDB entries that cover various segments of the protein, as well as showing representative non-overlapping structures both as static images and interactive 3D viewer (B).

image displays the ligand and its environment in 3D using LiteMol (31). If the molecule acts as a cofactor (38), this is also indicated in this section.

Below the gallery, ProtVista is used to highlight all the residues of the protein that directly interact with each of the small molecules, as determined by the Arpeggio tool (15). Each row shows the residues interacting with a particular small molecule, while collapsing the category can provide an overview of all the binding sites, which can be used to find protein sequence patterns in protein–ligand interactions. This overview can thus help in differentiating between sporadic interaction sites of solvent molecules and functionally relevant binding pockets.

The macromolecular interactions section has a similar layout to the ligands and environments section and uses the same web components to display different data (Figure 5B). The gallery in this section shows all the macromolecular partners which are directly interacting with the protein of interest, in at least one of the PDB entries and highlights how many PDB entries this interaction is observed. It also provides links to the aggregated views for these partner proteins. The images can be clicked to display the complexes in
Figure 5. The aggregated protein views provide information on all the observed interactions between the protein of interest and ligands or macromolecular interaction partners. The gallery of ligands (A) and macromolecules (B) can be used to display their interactions in an interactive LiteMol instance, and to navigate to the corresponding PDBs and PDB-KB entry pages to get more information about the partner molecules. Additionally, all the available functional annotations and biophysical parameters provided by PDB-KE partner resources are being displayed using ProtVista (C).

3D using LiteMol (31). Again, ProtVista is used to highlight all the residues that are directly interacting with the macromolecular partners, i.e. all the interaction interfaces are identified and listed.

The subset of the annotations provided by PDB-KE partners are displayed in the additional annotations section of the first version of the aggregated views for proteins (Figure 5C). This section shows all the predicted and experimentally determined functional sites and biophysical parameters contributed by the partner resources listed in Table 1. Curated annotations are colored blue, while prediction-based annotations are colored grey, to allow clear visual distinction between the two categories. Additionally, all the available variation data from UniProt (29) are displayed. All these annotations (as well as any data displayed with ProtVista from any of the sections) can be downloaded in CSV and JSON formats.

Similar proteins

The similar protein section shows all the UniProtKB accessions that are in the same UniRef90 cluster as the protein of interest, i.e. those proteins that share >90% sequence identity, further restricted to a minimum of 70% coverage by a single PDB structure (27). Further developing this section can help to transfer annotations and structural information from approximately 47 500 structurally characterised proteins (i.e. UniProtKB accessions represented in the PDB) to more than 2.2 million UniProtKB accessions reached via the UniRef90 clusters.
CONCLUSIONS AND OUTLOOK

PDBe-KB is an open-access, collaborative and integrated resource managed by the PDBe team at EMBL-EBI. The goal of PDBe-KB is to place macromolecular structure data in their biological context, and therefore to help the broader scientific community to utilize these data in both fundamental and applied research and education, as well as to reduce the fragmentation of the valuable annotation data derived by many structural bioinformatics resources. PDBe-KB was established in 2018, and since then it has gathered contributions from 18 resources from 8 countries, providing over 520 million residue-level annotations. PDBe-KB recently became an activity of the ELIXIR 3D BioInfo community, which aims to bring together structural bioinformatics developers and scientists to work towards FAIRness of structural annotations data (23).

PDBe-KB aims to provide novel aggregated views of PDB data and their annotations, with the first set of such aggregated views, launched in March 2019, focusing on full-length proteins (i.e. UniProtKB accessions). We continue to work on displaying additional functional and structural annotations contributed by the PDBe-KB partner resources and by transferring annotations between proteins which are in the same UniRef90 cluster.

Recently, we have started designing the next set of aggregated views that will focus on small molecules in the PDB and will integrate the structural data from the PDB with the biological context annotations pertaining to small molecules. These small-molecule-centric views are expected to be released in the first half of 2020. Concurrently, in collaboration with the Complex Portal team (39) at EMBL-EBI we have started investigating the data and visualization requirements for creating aggregated views aimed at macromolecular complexes.

To further improve data accessibility, we packaged the PDBe-KB graph database and made it distributable so that the complete database can be installed and queried locally by the scientific community and industry partners alike.

We extend an invitation to explore the possibility of joining the PDBe-KB consortium to new partners who can fill the gaps in current PDBe-KB biological context annotations, in particular pertaining to nucleic acids, membrane proteins, peptides, antibodies and biophysical parameters derived from experimental characterization of small molecule binding sites and protein complexes.

In conclusion, PDBe-KB brings structural data and annotations closer to the broader scientific community by allowing convenient access to data that would have been otherwise available in fragments from many specialist data resources. PDBe-KB also serves as a platform that helps specialist data resources to increase their visibility and helps them to increase the FAIRness of their data.

ACKNOWLEDGEMENTS

We would like to thank all the members of PDBe team for their continued support of the design and development of PDBe-KB. We would like to thank the UniProt and InterPro teams for their contributions on the ProtVista sequence feature viewer as well as with certain annotations.

FUNDING

BBSRC via the FunPDBe [BB/P024351/1]; India Partnering Award [BB/P025846/1]; SIFTS [BB/M011674/1]; ELIXIR supporting with multiple implementation studies that have contributed annotations; EMBL-EBI: BBSRC grant [BB/N019172/1 to T.P.L.]; NIH [1U01CA239106-01 to N.K.]; NIH [R01GM1149]; AIRC [IG 16895 to M.H.C.]; Research Foundation Flanders (FWO) [G.0328.16N to W.V.]; Hermesfonds for ELIXIR Belgium [AH.2017.051, IO 17001360]; Czech Science Foundation [17-21122S to K.B.]; ELIXIR CZ Research Infrastructure Project (MEYS) [LM2015047 to K.B.]; R.S. was also funded by European Regional Development Fund – projects ELIXIR-CZ [CZ.02.1.01/0.0/0.0/16_013/000177]; Wellcome Trust Doctoral Training [100150/Z/12/Z to F.M., G.J.B.]; Wellcome Trust Strategic Awards [WT09230, WT083481, WT09745, BBSRC BB/R014752/1, BB/J019364/1]; Wellcome Trust for support through an Investigator Award 2016–2021 [200814/Z/16/Z to T.L.B.]; E.D.L. is incumbent of the Recanati Career Development Chair of Cancer Research. S.D. acknowledges support from the Koshland Foundation. Funding for open access charge: BBSRC grant “FunPDBe” [BB/P024351/1].

Conflict of interest statement. None declared.

This paper is linked to: https://doi.org/10.1093/nar/gkz990.

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