RCSB Protein Data bank: Tools for visualizing and understanding biological macromolecules in 3D

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Abstract
Now in its 52nd year of continuous operations, the Protein Data Bank (PDB) is the premiere open-access global archive housing three-dimensional (3D) biomolecular structure data. It is jointly managed by the Worldwide Protein Data Bank (wwPDB) partnership. The Research Collaboratory for Structural Bioinformatics Protein Data Bank (RCSB PDB) is funded by the National Science Foundation, National Institutes of Health, and US Department of...
1 | INTRODUCTION

The Protein Data Bank (PDB) is in its 52nd year of continuous operation. Established in 1971 as the first open-access digital data resource in biology, it currently houses ~200,000 experimentally determined 3D structures of proteins and nucleic acids (DNA and RNA) and their complexes with one another and with small-molecule ligands (e.g., enzyme cofactors, inhibitors, drugs). Since 2003, the PDB archive has been jointly managed by the Worldwide Protein Data Bank (wwPDB, wwpdb.org) partnership. Full members of the wwPDB include three founding members—US-funded RCSB Protein Data Bank (RCSB PDB), PDB in Europe (PDBe), and Protein Data Bank Japan (PDBj)—plus the Electron Microscopy Data Bank (EMDB) and the Biological Magnetic Resonance Bank (BMRB). Protein Data Bank China (PDBc) was recently admitted to the wwPDB as an Associate Member. The wwPDB is committed to managing PDB data for users around the world at no charge for data deposition or egress, with no limitations on data usage. All PDB data are made available by wwPDB partners under the most permissive Creative Commons CC0 license (creativecommons.org/publicdomain/zero/1.0/). It is no exaggeration to say that the PDB was “walking the walk” decades before scholars began “talking the talk” regarding the principles emblematic of responsible data stewardship in the modern era: FAIR (Findability, Accessibility, Interoperability, and Reusability) and FACT (Fairness, Accuracy, Confidentiality, and Transparency).

Figure 1a documents the relentless growth of the PDB archive from its modest beginning with just seven X-ray crystal structures of proteins purified from natural sources. Technical advances in preparative biochemistry, exogenous protein production using bacterial and eukaryotic expression hosts, sample preparation and cryogenic preservation, structural biology instrumentation and facilities, methods of structure determination, and computational power have transformed structural biology (reviewed in Reference 12). Since publication of the first X-ray crystal structure of a protein (sperm whale myoglobin), the discipline has transformed itself from a small number of dedicated teams laboring for years, sometimes decades, to determine a single structure to a core element of the biological sciences. At the time of writing, PDB data depositors numbered more than 58,000 working on all permanently inhabited continents. Figure 1b enumerates contributions from macromolecular crystallography, nuclear magnetic resonance (NMR) spectroscopy, 3D electron microscopy (3DEM), and multiple methods to the growth of the PDB archive. For
macromolecular crystallography and 3D electron microscopy structures (now ~93% of the PDB archive), resolution distributions are illustrated in Figure 1c, d. Although a majority of PDB structures determined at better than 2.5 Å resolution come from macromolecular crystallography (~68% of the archive), 3D electron microscopy is now capable of delivering structures in the most favorable cases at nearly 1 Å resolution (e.g., 1.15 Å resolution structure of apoferritin, PDB ID 7a6a). The impact of PDB data spans fundamental biology, biomedicine, bioenergy, and bioengineering and biotechnology. The PDB has even found its way into the field of economics. Literature citations running from Agriculture to Zoology attest to the breadth of impact,
traversing all of the natural, mathematical, and engineering sciences. It is no exaggeration to state that recent advances in de novo protein structure prediction by AlphaFold2 and RoseTTAFold were only possible because of existence of the PDB as an enormous public archive of rigorously validated and expertly biocurated 3D biostructures. These “gold-standard” data served as a reliable source of training and testing datasets for development of artificial intelligence/machine learning-based tools. Within biomedicine, PDB data have transformed our understanding of human health and disease and continue to provide valuable starting points for structure-guided discovery of small-molecule drugs and design of new biologics agents.

United States federal funders have supported the PDB since its inception. As the US wwPDB data center, RCSB PDB is jointly funded by the National Science Foundation, the National Institutes of Health, and the US Department of Energy. Safeguarding and nurturing the PDB archive and providing open access to PDB data is the work of four coordinated RCSB PDB services, encompassing data deposition; archive management and access; data exploration; and training, outreach, and education.

This invited Tools Issue contribution describes how RCSB PDB (i) is organized to deliver services efficiently to many millions of users worldwide; (ii) works with wwPDB partners to process, validate, and biocurate ~16,000 new depositions annually; (iii) acts as the wwPDB-designated Archive Keeper; (iv) enables exploration and 3D visualization of PDB data integrated with more than 50 external data resources through its research-focused web portal RCSB.org; and (v) supports training, outreach, and education through its introductory PDB101.RCSB.org web portal.

2 | RESULTS

2.1 | RCSB PDB organizational structure

Four RCSB PDB services support transformation of global data into global knowledge (Figure 2).

2.1.1 | Service 1: data deposition

The wwPDB OneDep software system is used for global management of deposition, validation, expert biocuration, and remediation of macromolecular crystallography, 3D electron microscopy, nuclear magnetic resonance spectroscopy, and micro-electron diffraction structures, experimental data, and related metadata. RCSB PDB does not charge for data deposition.

2.1.2 | Service 2: archive management and access

As the wwPDB-designated Archive Keeper, RCSB PDB safeguards the PDB archive and maintains the PDBx/
mmCIF data dictionary (see below) that enables organization and searching of archived data. Programmatic access to PDB data is made available through multiple application programming interfaces (APIs).\textsuperscript{28} Strict adherence to the PDBx/mmCIF data standard enables facile integration of 3D structure information with ~50 trusted external data resources (Table 1). RCSB PDB does not charge fees for data egress and there are no limitations on data usage.

2.1.3 | Service 3: data exploration and delivery

The RCSB PDB research-focused RCSB.org web portal supports PDB data searching and download, browsing, 3D visualization, custom report generation, and analyses, again at no charge to users with no limitations on data usage.\textsuperscript{5,29}

2.1.4 | Service 4: outreach and education

PDB-101 delivers training, outreach, and education resources at no charge via its introductory PDB101.RCSB.org web portal focused on structural biology and its impact across the sciences.\textsuperscript{30}

Additional elements of RCSB PDB operations include a Customer Service Help Desk, supporting 3D structure depositors and PDB data consumers around the world, and an Infrastructure Team, working to ensure >99% 24 × 7 × 365 service availability uptime. Status of RCSB PDB servers, microservices, and APIs is monitored by NS1 (ns1.com) and made publicly available on a real time basis at status.rcsb.org.

RCSB PDB operations currently spans three performance sites, including its headquarters at Rutgers, The State University of New Jersey, plus smaller teams housed at the University of California San Diego-San Diego Supercomputer Center (UCSD-SDSC) and the University of California San Francisco (UCSF). RCSB PDB sub-teams have complementary roles within the organization. All Service 1 team members are based at Rutgers. A bi-coastal team of data managers and software developers supports both Services 2 and 3. Service 4 training, outreach, and education team members are based at Rutgers. The Customer Service Help Desk is coordinated from Rutgers. Infrastructure Team members are divided between Rutgers and UCSD-SDSC. The RCSB PDB team contains specialists in method-specific biocuration, software development, IT infrastructure, training, outreach, and education who work together to ensure that the RCSB PDB delivers high quality data and services.

Projects developed across all services receive guidance from the RCSB PDB Advisory Committee (recognized experts in fields, including but not limited to, structural biology, cell and molecular biology, computational biology, information technology, and education). The wwPDB Advisory Committee, an international team of experts in macromolecular crystallography, 3D electron microscopy, nuclear magnetic resonance spectroscopy, bioinformatics, and data science additionally provides guidance on matters surrounding the PDB archive.

All four RCSB PDB Services can be accessed from the RCSB.org home page depicted in Figure 3. Navigation menus arrayed across the top of the page provide access to drop-down menus, with a selection echoed in the tabs in the center panel. Deposit provides access to all Service 1 features. Services 2 and 3 can be accessed from: Search, Visualize, Analyze, and Download. Learn provides access to all Service 4 resources. Miscellaneous information regarding RCSB PDB is available from the More navigation menu. Documentation provides access to General Help and documentation for the entire web portal. Documentation and news updates are fully text searchable using the pull-down option from the search bar. Careers advertises current job openings across the organization.

2.2 | RCSB PDB Service 1: Data deposition

All PDB structures are deposited through the wwPDB OneDep software system (deposit.wwpdb.org), which supports structure deposition,\textsuperscript{26} rigorous validation,\textsuperscript{31–33} expert biocuration,\textsuperscript{34} and occasional remediation.\textsuperscript{35–37} Experimental methods supported by OneDep include macromolecular crystallography, 3D electron microscopy, nuclear magnetic resonance spectroscopy, and micro-electron diffraction. In addition to OneDep, RCSB PDB offers GroupDep\textsuperscript{38} for deposition of large numbers (>10) of closely related macromolecular crystallography structures (e.g., results from crystallographic fragment screening\textsuperscript{39}) via requests sent to deposit@deposit.rcsb.org.

Development of OneDep enhanced the rate at which the international team of wwPDB biocurators can process incoming PDB data, permitting scaling of Service 1 operations without the need of additional personnel.

During 2021, wwPDB partners processed 14,573 experimental structure depositions, a number slightly lower than the record of 15,436 set in 2020 during the COVID-19 lockdown but higher than in 2019 (13,377). During this same period, RCSB PDB processed ~39% of the global depositions (from the Americas and Oceania, and GroupDep\textsuperscript{38} users globally), PDBj
In response to community input, RCSB PDB continues to integrate new trusted external data resources as updated at [rcsb.org/docs/general-help/data-from-external-resources-integrated-into-rcsb-pdb](http://rcsb.org/docs/general-help/data-from-external-resources-integrated-into-rcsb-pdb).

| Resource | Description |
|----------|-------------|
| AlphaFold DB<sup>19,62</sup> | Computed structure models by AlphaFold2 |
| ATC | Anatomical therapeutic chemical (ATC) classification System from World Health Organization |
| Binding MOAD<sup>63</sup> | Binding affinities |
| BindingDB<sup>64</sup> | Binding affinities |
| BMRR<sup>65</sup> | BMRR-to-PDB mappings |
| Catalytic site Atlas<sup>66</sup> | Active sites and catalytic residues in enzymes |
| CATH<sup>67</sup> | Protein structure classification |
| Cambridge structural Database<sup>68</sup> | Crystallographic small molecule data from the Cambridge crystallographic data Centre |
| ChEBI<sup>69</sup> | Chemical entities of biological interest |
| ChEMBL<sup>70</sup> | Manually curated database of bioactive molecules with drug-like properties |
| DrugBank<sup>71</sup> | Drug and drug target data |
| ECOD<sup>72</sup> | Evolutionary classification of protein domains |
| EMDB<sup>8</sup> | 3D electron microscopy density maps and associated metadata |
| ExplorEnz<sup>73</sup> | IUBMB enzyme nomenclature and classification |
| Gencode<sup>74</sup> | Gene structure data |
| Gene Ontology<sup>75</sup> | Gene structure data |
| Genotype-tissue expression (GTEx)<sup>76</sup> | Tissue-specific gene expression data |
| GlyCosmos<sup>77</sup> | Web portal integrating the glycosciences with the life sciences |
| GlyGen<sup>78</sup> | Data integration and dissemination resource for carbohydrates and glycoconjugates |
| GlyTouCan<sup>79</sup> | Glycan structure repository |
| Human gene nomenclature committee ([genenames.org](http://genenames.org)) | Human gene name nomenclature and genomic information |
| IMGT<sup>80</sup> | International ImMunoGeneTics information system |
| Immune epitope Database<sup>81</sup> | Antibody and T cell epitopes |
| International mouse phenotyping consortium (IMPC, [mousephenotype.org](http://mousephenotype.org)) | Mouse gene phenotype data |
| MemProtMD<sup>82</sup> | Database of membrane proteins embedded in lipid bilayers |
| ModelArchive ([modelarchive.org](http://modelarchive.org)) | Computed structure models (e.g., by RoseTTAFold) |
| Mpstruc ([https://blanco.biomol.uci.edu/mpstruc/](http://blanco.biomol.uci.edu/mpstruc/)) | Classification of transmembrane protein structures |
| NCBI Gene<sup>47</sup> | Gene info, reference sequences, et al. |
| NCBI Taxonomy<sup>47</sup> | Organism classification |
| NDB<sup>83</sup> | Experimentally determined nucleic acids and complex assemblies |
| OPM<sup>84</sup> | Orientations of proteins in membranes database; classification of transmembrane protein structures and membrane segments |
| PDBbind-CN<sup>85</sup> | Binding affinities |
| PDBFlex<sup>86</sup> | Protein structure flexibility |
| PDBTM<sup>87</sup> | Protein Data Bank of Transmembrane Proteins |
| Pfam<sup>88</sup> | Protein families |
| Pharos<sup>89</sup> | Drug targets and diseases |
| ProteinDiffraction.org ([proteindiffraction.org](http://proteindiffraction.org)) | Diffraction images |
| PubChem<sup>90</sup> | Chemical information |
processed ~32% of global depositions (from Asia and the Middle East; reflecting significant year-on-year growth in the number of depositions coming from the People’s Republic of China). A total of 12,593 new structures were publicly released into the PDB during 2021, which is lower than the record of 14,009 set in 2020 but again higher than in 2019 (11,497). Year-to-date deposition trends are on track (as of early October) for the number of new depositions in 2022 to exceed 16,000.

Service 1 support for global PDB data deposition embodies four elements, including Prepare Data, Validate Data, Deposit Data, and related Help & Resources, all accessible from the Deposit menu and home page panel at RCSB.org.

i. Prepare Data provides access to information about the PDBx/mmCIF data dictionary underpinning the entire PDB archive, plus software tools developed by RCSB PDB (e.g., pdb_extract, SF-Tool, Ligand Expo, Maxit)\(^{30-43}\) that are used frequently by structural biologists for organizing information prior to data deposition.

ii. Validate Data provides access to the stand-alone, anonymous wwPDB Validation Server and APIs, which should be used to identify possible issues with experimental data and/or atomic coordinates during structure determination. Depositors are strongly encouraged to resolve any issues identified by the Validation Server before proceeding to structure submission. Additional resources available under Validate Data include Information for Journals, pertaining to use of wwPDB Validation Reports during manuscript review, and Validation Task Forces, describing related community engagement. Multiple descriptions of structure validation have been published in peer-reviewed journals.\(^{31-33,35,44,45}\)

iii. Deposit Data provides access to the wwPDB OneDep\(^{26}\) global system for structure deposition, validation,\(^{31-33}\) biocuration,\(^{44}\) and remediation.\(^{35-37}\) This software tool was developed jointly by wwPDB partners. It is used by all wwPDB data centers worldwide to ensure data uniformity, completeness, and consistency across the PDB archive. Data depositors now benefit from a streamlined deposition process, reducing the time required for data entry while ensuring complete data capture. Every few years (or when new features are added to validation software), wwPDB Validation Reports are recomputed for all structures in the archive to stay abreast of improvements in average structure quality and support statistics-based detection of outliers. wwPDB biocuration processes scrutinize biopolymers, small molecule ligands, and carbohydrates through different OneDep Modules. The Sequence Module cross-checks biopolymer sequences against UniProt\(^{46}\) and NCBI\(^{47}\) databases for taxonomy and sequence reference annotation. Small molecule ligands are extracted from the coordinates, searched for matched ligands in the wwPDB Chemical Component Dictionary (Chemical Component Dictionary),\(^{48}\) and standardized with nomenclature in the Ligand Module. This module also allows biocurators to create Chemical Component Dictionary definitions for new ligands. Saccharide units covalently bound to a biopolymer are represented as branched carbohydrates based on connectivity (e.g., N-glycosylation).

Depositors are encouraged to submit macromolecular crystallography data in bulk using RCSB PDB Group Deposition capabilities,\(^{38}\) where appropriate (e.g., crystallographic fragment screening). The efficiency of GroupDep, as it is informally known, encourages capture of valuable data that might
otherwise remain locked up within individual laboratories. Integrative or hybrid methods structures determined with more than one experimental technique (e.g., 3D electron microscopy combined with chemical cross linking) should be deposited to the wwPDB PDB-Dev prototype repository (pdb-dev.wwpdb.org).49–54

iv. Deposition Help & Resources provides access to Deposit FAQs, Tutorials, Deposition Policies, Annotation Procedures, the PDBx/mmCIF Dictionary (see below), Chemical Component Dictionary, PDB File Format Guides, BioSync Beamlines and Facilities information,55 and Related Tools.

2.3 RCSB PDB Service 2: Archive management and access

PDBx/mmCIF Data Standard: PDB data architecture is defined by the PDBx/mmCIF dictionary.56–58 It is the macromolecular extension of an earlier community data standard, the Crystallization Information Framework.59
(cif.iucr.org), developed by the International Union of Crystallography for small molecule X-ray diffraction. The macromolecular data standard is maintained by the wwPDB partners in collaboration with wwPDB PDBx/mmCIF Working Group domain experts (wwpdb.org/task/mmci).58 wwPDB partners and the Working Group collaborate on developing terminologies for emerging and rapidly evolving methodologies (e.g., X-ray Free Electron Laser Serial Crystallography and 3D electron microscopy), and remediating (or enhancing) representations for existing data content (e.g., carbohydrates).

Within the PDBx/mmCIF dictionary, PDB data follow a strict hierarchy. An individual PDB structure is referred to as an Entry, identified with a unique PDB ID (currently four alphanumeric characters, for example, 1q2w). Within each PDB Entry, every chemically unique molecule is defined as an Entity (including Polymer, Branched, or Non-polymer), denoted by a numeric Entity ID. Polymer Entities (including proteins or short polypeptide chains, DNA, and RNA) are made up of covalently linked chemical building blocks as described in the Chemical Component Dictionary and individually numbered according to position within the Polymer sequence. Branched Entities are either linear or branched carbohydrates, composed of saccharide units covalently linked via one or more glycosidic bonds. Non-polymer Entities are small molecules (enzyme cofactors, ligands, water molecules, etc.) and ions (e.g., Na\(^{+}\), Cl\(^{-}\), Zn\(^{2+}\)). Every Non-polymer Entity has a unique Chemical Component Dictionary ID (one to three or more character alphanumeric code). The Chemical Component Dictionary provides nomenclature standards and chemical descriptions for all Non-polymer Entities and every Polymer Entity component occurring within the PDB archive. Within a PDB Entry, there can be multiple Instances (or copies) of any particular Entity, each labeled with a unique Chain ID (one or more alphanumeric characters, for example, A, AA, ...). In nature, multiple proteins and/or nucleic acid chains frequently occur as supramacromolecular complexes. Within the PDBx/mmCIF hierarchy, they are referred to as Assemblies, each with a unique numeric Assembly ID. The standardized nomenclature of the PDBx/mmCIF dictionary also provides the backbone of the Advanced Search interface, which is explained in greater detail below and in RCSB.org Advanced Search documentation.

Archive Keeping: As wwPDB-designated PDB Archive Keeper, RCSB PDB is responsible for safeguarding and managing the PDB, and making its contents freely available. At the time of writing, the size of the archive was ~1 TB (excluding 3D electron microscopy experimental data stored in EMDB, which currently totals ~6 TB). The entire corpus of PDB data is safeguarded in multiple copies in secure institutional computer machine rooms located on both US coasts, and backed up twice weekly (i.e., before and after each weekly update).

In its role as wwPDB-designated PDB Archive Keeper, RCSB PDB releases an updated version of the archive every week using a two-stage process. For each new PDB Entry slated for the upcoming weekly release, Stage One includes sequence(s) (amino acid or nucleotide) for each distinct Polymer Entity; InChI string(s) for each distinct ligand; and crystallization pH value(s), where appropriate. They are released at the wwPDB web portal (see www wwpdb.org ftp pdb ftp-sites) every Saturday by 03:00 Universal Time Coordinated (UTC). Stage One supports weekly blind challenges for in silico prediction of protein structure (Continuous Automated Modeling Evaluation or CAMEO, cameo3d.org) and small-molecule docking (Continuous Evaluation of Ligand Protein Prediction or CELPP, drugdesigndata.org/about/CELPP). Stage Two completes the process every Wednesday at 00:00 UTC by releasing the updated PDB archive in full (currently adding ~300 new structures/week, updating previously released entries, and occasionally removing obsoleted structures).

PDB data are freely distributed via File Transfer Protocol (FTP) over Hypertext Transfer Protocol (HTTP) and remote sync (rsync), providing universal open access to the archival information in two forms (latest archive, ftp.wwpdb.org/pub/pdb/data; and latest and prior versions archive, ftp-versioned.wwpdb.org).

PDB data are also made available without fees or egress charges by Amazon Web Services (AWS) through its Open Data Sponsorship Program registry.opendata.aws/pdb-3d-structural-biology-data/. Download tests conducted from Asia, the United Kingdom, and the United States documented significantly faster download speeds for larger PDB structure data files hosted by AWS versus delivery of the same information for the RCSB PDB web portal (and those maintained by both PDBe and PDBj). PDB data consumers wishing to carry out large-scale data downloads are strongly encouraged to do so using AWS (s3.rcsb.org).

During 2021, a total of ~2.36 billion PDB data files were downloaded from the archive, which is nearly double the previous record of 1.32 billion set in 2020. Historically, requests for data downloads have come from every sovereign country recognized by the United Nations. Approximately two-thirds of data downloads come from Asia, the United Kingdom, and the United States documented significantly faster download speeds for larger PDB structure data files hosted by AWS versus delivery of the same information for the RCSB PDB web portal (and those maintained by both PDBe and PDBj). PDB data consumers wishing to carry out large-scale data downloads are strongly encouraged to do so using AWS (s3.rcsb.org).

External Data Integration: Service 2 also integrates PDB data with information from ~50 trusted external resources (Table 1). Integration of information from
external resources is a critical element of an exhaustive weekly process that loads the updated PDB archive into the RCSB PDB data warehouse. Thereafter, it is made available for our public data access APIs. At the time of writing, the entire corpus of data managed by RCSB PDB occupied >100 TB of digital storage.

To provide users with the most current information about a structure entry, information from trusted external resources is integrated on a weekly basis. RCSB.org is a “living data resource,” while scholarly publications describing PDB Entries are static documents that reflect what was known about the biomolecule(s) at the time of study. Thereafter, it is not uncommon for new biological or biochemical functions of a macromolecule to come to light, or new disease-causing mutations to be identified. Such new findings are integrated with PDB data every week, thereby ensuring that RCSB.org users have access to the most current information pertaining to every 3D biostructure in the public domain.

2.4  |  RCSB PDB Service 3: Data exploration and delivery

Most RCSB PDB users access the archive through our RCSB.org research-focused web portal. In 2021, more than 6.8 million unique internet protocol (IP) addresses from around the world were used to access RCSB.org, which exceeded our 2020 record of nearly 6.7 million (during the pandemic lockdown). We estimate that ~99% of these PDB data consumers are not experts in structural biology. Their research interests are very broad, encompassing fundamental biology, biomedicine, energy sciences, and bioengineering and biotechnology. Service 3 is tasked with creation and maintenance of a public-facing web portal to serve the diverse needs of these multiple communities of users.

As described earlier, the RCSB.org home page (Figure 3) displays navigation menus that provide access to Deposit, Search, Visualize, Analyze, Download, Learn, More, Documentation, and Careers resources. Immediately below these menus, users will find a top bar Search box that supports text searching (based on ElasticSearch, elastic.co) of the entire PDB archive or RCSB PDB Documentation, depending on the option selected. Immediately below the Search box, single click access is provided to Advanced Search and Browse Annotations features. The remainder of the home page is devoted to a description of the PDB archive and RCSB PDB, COVID-19 Coronavirus Resources, Careers, Molecule of the Month, Latest Entries, Features & Highlights, and News.

The flexibility, generality, and utility of RCSB PDB RCSB.org web portal tools are exemplified below with a case study focused on human programmed cell death protein 1 (PD-1), cancer immunotherapy, tumor neoantigens, monoclonal antibodies, and small-molecule drugs.

Therapeutic anti-PD-1 antibodies are used clinically to reduce the likelihood of “tumor sparing” by T-cells when they encounter malignant cells expressing program death-ligand proteins PD-L1 or PD-L2. Anti-cancer immune surveillance mechanisms, relying on detection of tumor neoepitopes (i.e., oligopeptide segments of tumor neoantigens) presented to the immune system by major histocompatibility complex (or MHC) classes 1 and 2, usually detect and kill nascent tumors before they begin to proliferate and cause disease. However, T-cell responses to tumor neoepitopes can be down regulated by a so-called “immune checkpoint” when PD-1 (normally displayed on T-cell surfaces) binds to PD-L1 (or PD-L2) present serendipitously on the surface cancer cells. Targeting this immune checkpoint with monoclonal antibodies is revolutionizing treatment of solid tumor cancers.

President Jimmy Carter, for example, benefitted from intravenous infusions of pembrolizumab (Keytruda.com, trade name Keytruda), which put his late-stage melanoma into long-term remission and may well have cured him of his widely metastatic cancer. Pembrolizumab and other immune checkpoint therapies are thought to be most effective against high mutation burden cancers. Both late-stage melanoma and tumors with defects in DNA mismatch repair (e.g., a subset of rectal cancers) carrying large numbers of tumor neoantigens can be treated successfully using antibodies that block interactions between PD-1 and PD-L1 (or PD-1 and PD-L2). Targeting PD-L1 with monoclonal antibodies has also proven effective in some individuals for some cancers (e.g., durvalumab, Imfinzi.com, trade name Infizi). James Allison (MD Anderson Cancer Center) and Tasuku Honjo (Kyoto University) shared the 2018 Nobel Prize in Physiology or Medicine for their “discovery of cancer therapy by inhibition of negative immune regulation.” Honjo is credited with the discovery of PD-1, the target of pembrolizumab and nivolumab (Opdivo.com, trade name Opdivo).

Structure Summary Page: For a topic like this, users often begin by viewing individual PDB entries, typically described in peer-reviewed publications that enumerate PDB structure IDs explicitly. When a single PDB ID (e.g., PDB structure 5jze101) is entered into the RCSB.org top bar Search box and users press the Return key or click the Magnifying Glass icon, they are taken directly to the relevant Structure Summary Page. Each Structure Summary Page provides summary information for that entry, with “tabs” linking to web portal pages displaying more detailed information specific to the entry. Figure 4a
FIGURE 4  Legend on next page.
shows the upper portion of the PDB ID 5jxe Structure Summary Page, which provides structure images of pembrolizumab anti-PD1 monoclonal antibody Fab recognizing the ectodomain of PD-1, a summary of structure contents, wwPDB Validation quality assessment, and the primary literature citation. The Macromolecules section (Figure 4b) contains three subsections: the first providing a detailed summary for the two copies of human PD-1 present in the PDB Entry, plus two subsections each for two copies of the light and heavy chain of the Fab. The bottom portion of the Structure Summary Page (Figure 4c) includes sections for Experimental Data & Validation (providing a summary of the X-ray crystallographic experiment plus a button to View more in-depth experimental information) and an Entry History section with deposition and revision information.

1D-3D Data Visualization: PDB structure 5jxe reveals at the atomic level how pembrolizumab recognizes the ectodomain of PD-1 on the surface of human T-cells. More details regarding the structure can be explored by clicking the 1D-3D View button in the upper section of the Structure Summary Page (Figure 4a, red ellipse), which takes the user to a simultaneous graphical display of the one-dimensional (1D) amino acid sequences (Figure 5, left) and the 3D structure (Figure 5, right) with various annotations incorporated into the 1D view, including identification of PD-1 residues involved in “Binding Chain F” (heavy chain of the pembrolizumab monoclonal antibody). The right-hand 3D view panel uses the Mol* molecular graphics open-source software system, first deployed on the RCSB.org web portal in 2020. Mol* operates entirely within a web browser without the need to download additional software. It also inter-operators with the PDBx/mmCIF data standard, allowing 3D graphical delivery of value-added annotations at the level of individual amino acid residues.

**Figure 4** Structure Summary Page for PDB ID 5jxe. (a) Upper section hosts structure images, summary of structure contents, wwPDB Validation quality assessment, and primary literature citation. The link to the 1D-3D viewer is highlighted with a red ellipse. (b) Macromolecules Section: First of three subsections providing detailed summary for the two copies of human PD-1 present in the structure. (N.B.: Additional subsections for two copies each of the heavy and light chains of the pembrolizumab monoclonal antibody Fab are not shown.) “Find similar proteins by:” can be invoked for Sequence similarity searching (magenta ellipse) from a pull-down menu to select desired percentage of sequence similarity. “Find similar proteins by:” can be invoked for 3D Structure similarity searching (red ellipse) with a single mouse click. The Entity Group Sequence Clusters 30% Identity search button is denoted with a magenta rectangle. (c) Experimental Data & Validation and Entry History Sections

**Figure 5** 1D-3D View for PDB ID 5jxe. In the left panel, clicking on Ser55 (highlighted with a red arrow) on the track CHAIN A (at the top of the page) invokes 3D graphical display of the view on the right, showing a hydrogen bond between the sidechains of Ser55 of human PD-1 (labeled Ser55) and Arg99 of the pembrolizumab Heavy Chain (not labeled). Atom color coding: O-red; N-blue, C-green (for PD-1), C-yellow (for pembrolizumab Heavy Chain), and C-pink (for pembrolizumab Light Chain)
residues. Extensive introductions to Mol* can be found in a 2021 Nucleic Acids Research Web Server issue article,\textsuperscript{104} the 2022 Protein Science Protein Tools special issue,\textsuperscript{29} and RCSB.org documentation.

Mousing over the interacting PD-1 residues in the 1D view highlights locations of these stabilizing interactions in the 3D view. Clicking on any one of the interacting PD-1 residues (e.g., Ser55, Figure 5, red arrow) within the 1D view causes the 3D view to center on the residue and display all non-H atoms within 5 Å, showing hydrogen bonds and other interactions as dashed lines (e.g., H-bond between PD-1 Ser55 and pembrolizumab heavy chain Arg99, Figure 5, right).

Structure-guided Data Exploration: The PDB archive often includes multiple entries related to a particular topic, which may be found using the “Find similar proteins by: Sequence” button on the Structure Summary Page. This feature is available for every Polymer Entity in the Entry (Figure 4b: magenta ellipse, with pull-down menu for 100% identity, 95%, ...). Starting from the Structure Summary Page for PDB structure 5jxe, invoking “Find similar proteins by: Sequence (100% identity)” returns a results page listing 10 Polymer Entities that are exact matches in amino acid sequence to human PD-1. Scrolling part way down this list reveals PDB structure 4zqk,\textsuperscript{105} entitled “Structure of the complex of human programmed death-1 (PD-1) and its ligand PD-L1.” Clicking on 4zqk takes the user to the Structure Summary Page for this related structure. Selecting the 1D-3D View button then allows the user to identify the residues in PD-1 involved in molecular recognition and explore interactions between PD-1 and PD-L1 at the atomic level, as described above.

Alternatively, clicking on the “Find similar proteins by: 3D Structure” button on the structure summary page of PDB ID 5jxe (Figure 4b, red ellipse) returns a results page listing >100 polymer entities with similar 3D structure (root-mean-square-deviation or RMSD values ranging from 0.0 Å for the exact match to ~2.1 Å). The large number of structurally similar proteins represented in the PDB archive reflects the ubiquity of the immunoglobulin-like beta sandwich fold, as classified by SCOP\textsuperscript{95}/SCOPe,\textsuperscript{96} SCOP2,\textsuperscript{106} ECOD,\textsuperscript{72} CATH,\textsuperscript{67} IMGT antibody Annotation,\textsuperscript{80} and SAbDab antibody Annotation,\textsuperscript{93} all of which are available from the annotations tab on the structure summary page.

Another option for exploring related proteins is supported by the newly introduced RCSB PDB Group feature, located immediately below the description of each protein Polymer Entity occurring in a given PDB structure (Figure 4b). For the human PD-1 Polymer Entity in PDB structure 5jxe, clicking on the 30% identity Sequence Cluster button (Figure 4b, magenta rectangle) delivers the user to the Group Summary page depicted in Figure 6a, which displays information for a cluster of 12 related proteins (encompassing all occurrences of human and murine PD-1 represented in the PDB archive). Clicking on the Group Sequence option reveals a multiple sequence alignment of all 12 related proteins (Figure 6b). Additional viewing options on the Group Sequence page include Structural Features (including a consensus sequence, secondary structural elements, and CATH and SCOP annotations) and Binding Sites (identifying amino acid residues in contact with various monoclonal antibody Fabs, PD-L1, and PD-L2).

Additional Functionality: Additional Structure Summary Page functionality can be accessed using the other top tabs shown in Figure 4a, which include 3D View, Experiment, Sequence, Genome, and Versions. Clicking on the 3D View tab takes the user to a stand-alone Mol* 3D graphics window, equivalent to the right-hand portion of the 1D-3D View. The Experiment tab (typically used only by structural biologists) displays a detailed summary of the X-ray crystallography study yielding the structure. The Sequence tab takes the user to the equivalent of the left-hand portion of the 1D-3D View. The Genome tab presents an alignment of the human PD-1 polypeptide chain with human chromosome 2. (N.B.: At the time of writing, this functionality was only supported for 1,614 organisms.) Clicking on the Versions tab displays information regarding any revisions to the PDB Entry following its initial release. Going beyond necessary minor revisions of a clerical nature made to individual PDB entries by wwPDB biocuration team members from time to time, depositors-of-record (i.e., principal investigators, research team leaders) are encouraged to “correct” their PDB entries (wherever appropriate) by providing updated atomic coordinate files. Since 2018, atomic coordinate versioning without triggering issuance of a new PDB ID has been supported by the wwPDB in response to depositor feedback.

2.5 | Searching the PDB archive

In cases where users do not know the PDB ID of a relevant Entry, multiple options for searching the archive are available on the RCSB.org web portal. The top bar Search box located in the upper section of the homepage is the workhorse of RCSB PDB search capabilities. When non-PDB ID text is entered into the Search box, a drop-down list appears with possible search refinements identifying the input text as being found within Additional Structure Keywords, Structure Title, Structure Author, Polymer Entity Description, Gene Name Source Organism, Taxonomy Name, etc. Clicking on one of the options in the
drop-down list takes the user to a combined results page. In most cases, scrolling down the results page will allow identification of relevant PDB structures for exploration via their Structure Summary Pages by clicking on a PDB ID. Another option supports free text searches, which are carried out most expeditiously when the phrase of interest is enclosed within double quotes. Otherwise, structures containing any of the text words in the query will be returned and may include false positives.

Directed searches using full Boolean logic are supported by the Advanced Search page, which can be reached from the top of any RCSB.org web portal page by

**FIGURE 6**  Entity Group Sequence Clusters 30% Identity for human PD-1. (a) Group Summary page, consisting of three components: ribbon representation drawings for individual Group Members with buttons provide access to the Structure Summary Pages (Summary) and a Mol* display (Structure); a graphical table showing Group Members; and bar-graph summarizing public release dates for individual Group Members. (b) Group Sequence page, providing a multiple sequence alignment of Group Members. Additional tabs on the Group Sequence page provide access to information about Structural Features and Binding Sites.
clicking on the Advanced Search button. The example depicted in Figure 7 was used to identify PDB structures of human PD-L1 (UniProt Accession Code Q9NZQ7) that also contain small-molecule ligands with molecular weight $\geq 300$ and $\leq 1000$ Da. At the time of writing, invoking this Advanced Search yielded 13 PDB structures containing human PD-L1 bound to a distinct small-molecule ligand. Toggling between Return Structures and Return Molecular Definitions (highlighted in figure) changes the output from individual PDB structures to individual small-molecule ligands.

These 13 PDB structures were described in six peer-reviewed journal publications by a number of research groups focused on discovering and developing orally bioavailable, small-molecule drugs as alternatives to monoclonal antibody immune checkpoint anti-cancer therapies.\textsuperscript{107–113} Such ligands are thought to reduce tumor immune evasion via small molecule-induced dimerization PD-L1 and subsequent internalization. This effect on the immune checkpoint is mechanistically distinct from that of anti-PD-1 and anti-PD-L1 monoclonal antibodies, which interdict PD-1/PD-L1 mediated interactions between T-cells and tumor cells. Disappearance of PD-L1 from the surface of the tumor cell prevents them from engaging with T-cells in a manner that could down regulate the immune response to the tumor. A comparable Advanced Search intended to identify 3D structures of human PD-1 (UniProt Accession Code Q15116) bound to small-molecule ligands in PDB detected zero matches at the time of writing. Effective small-molecule drugs targeting immune checkpoints would provide access to cancer immunotherapies for millions of patients around the world unable to afford monoclonal antibody treatments (for reference, the current list price for a full 2-year course of pembrolizumab is ~US $360,000, plus ancillary charges for intravenous infusions occurring every 3 or 6 weeks).

2.6 | Pairwise structure comparison

The Pairwise Structure Comparison tool is one of the most popular RCSB.org web portal features and was described in detail in the 2022 Protein Science
Available from the Analyze navigation menu, Pairwise Structure Comparison supports superposition and quantitative comparison of 3D structures using Java implementations of well-established tools (e.g., Combinatorial Extension or CE,\textsuperscript{114} CE with Circular Permutations,\textsuperscript{115} FATCAT,\textsuperscript{116} TM-align,\textsuperscript{117} and superposition guided by Smith-Waterman local sequence alignment\textsuperscript{118}) provided by the BioJava project.\textsuperscript{119} This feature has also been implemented within the stand-alone version of the Mol* 3D Viewer, which can be accessed from the Visualize navigation menu. Users can open atomic coordinate files and import them into the Mol* session in either legacy PDB or PDBx/mmCIF formats, or simply download them directly from the PDB archive. Once the two PDB structures are visible on the Mol* 3D-Canvas, clicking on the Superposition Panel allows the user to select displayed objects (i.e., by chains or by atoms) and superpose them.

Figure 8, downloaded by activating the Screenshot feature (camera iris icon) available on the Mol* 3D-Canvas and clicking the Download button, illustrates the result of superposing Chain A of PDB ID 5jxe (human PD-1 bound to pembrolizumab) with Chain B of PDB ID 4zqk (human PD-1 bound to human PD-L1), which gave RMSD ~1.0 Å for common Cα atom pairs. The image of the superposed structures reveals the mechanism of action of pembrolizumab at the atomic level. Binding of the Fab would sterically interfere with PD-L1 binding to PD-1, thereby preventing down regulation of the T-cell when the T-cell receptor detects a neoepitope displayed on the surface of the malignant cell by major histocompatibility complex classes I and II.

A detailed guide to the superposition process can be accessed by using the path documentation navigation menu \(\rightarrow\) Mol* \(\rightarrow\) FAQs/scenarios leading to the FAQ/scenario, entitled “How do I compare/superpose multiple structures?” an accompanying FAQ/scenario, entitled “How can I save a Mol* session or state and return to it at a later time?” explains how to generate and download files for saving a Mol* session or Mol* state for later use.

**2.7 RCSB PDB Service 4: Training, outreach, and education**

RCSB PDB has a long-standing commitment to training and community outreach and education. The dedicated web portal PDB-101 (denoting an introductory course; PDB101.RCSB.org) was launched in 2011 to support PDB archive exploration by teachers, students, and the general public. A detailed description of PDB-101 activities was provided most recently in the 2022 Protein Science Tools Issue.\textsuperscript{30} Service 4 resources help train the next generation of PDB users and promote the importance of structural biology and protein science to nonexperts. Regularly published features include the highly popular Molecule of the Month (MotM) series,\textsuperscript{120} 3D biostructure-related activities, molecular animations and videos, and educational curricula. Materials are organized into various categories (Health and Disease, Molecules of Life, Biotech and Nanotech, and Structures and Structure Determination) and searchable by keyword (e.g., cancer, checkpoint therapy, and antibody).

“Cancer Biology and Therapeutics” is the biennial health focus of PDB-101 for 2022 and 2023. Recently completed and ongoing activities related to cancer are as follows:

1. Publication of six new MotM articles that were coauthored by under-represented minority undergraduate students participating in the Rutgers University Institute for Quantitative Biomedicine 2022 Winter Boot Camp (iqb.rutgers.edu), entitled “Science Communication in Biology and Medicine.” Article topics include Vascular Endothelial Growth Factor (VegF)
and Angiogenesis (Figure 9; March 2022); Nicotine, Cancer, and Addiction (May 2022); Pyruvate Kinase (June 2022); HER2/neu and Trastuzumab (April 2022); Non-Homologous End Joining Supercomplexes (July 2022); and Secretory Antibodies (August 2022).

2. RCSB PDB hosted the 2022 High School Video Challenge (pdb101.rcsb.org/events/video-challenge/the-challenge). Participating students were asked to explore two cancer-related molecular signaling systems: the p53/p21 pathway and the Epidermal Growth Factor Receptor (EGFR)/Ras pathway. In all, 38 teams from around the US created short videos, each telling a coherent story explaining underlying scientific concepts and touching on public health aspects of cancer, such as screening, prevention, and awareness. External expert reviewers selected the winning videos based on scientific content and communication effectiveness, while the public was invited to vote on a “viewer’s choice” video.

Previously published PDB-101 activities related to cancer included MotM articles on p53 Tumor Suppressor (July 2002); Estrogen Receptor (September 2003); Epidermal Growth Factor (June 2010); RAS Protein (April 2012); RAF Protein Kinases (March 2016); Programmed Cell Death Protein 1 (December 2016); Chimeric Antigen Receptors (October 2017); MDM2 (June 2019); Cyclin and Cyclin Dependent Kinase (August 2019); and Cisplatin and DNA (March 2021). Given the public health importance of human papilloma viruses (HPV, a cause of genital warts and cervical, genital, anal, and oropharyngeal cancers), PDB-101 developed related content in the form of (i) a molecular origami paper-folding model of the reconstituted HPV capsid that has been used successfully for vaccine design, (ii) a Human Papillomavirus and Vaccines MotM article (May 2018); and (iii) a Nature Oncogene review article focused on the biology of HPV viewed through the lens of the PDB.

Going beyond our biennial theme of Cancer Biology and Therapeutics, we use our PDB101.RCSB.org web portal and Service 4 social media channels (Twitter: @build-models; Facebook: RCSBPDB) to empower educators, students, and the general public to explore and understand how 3D biostructures are being used in research to address global health challenges of immediate concern. Compelling SARS-CoV-2 images created by Service 4 team member Professor David S. Goodsell (pdb101.rcsb.org/sci-art/goodsell-gallery) and introductory materials related to coronaviruses were provided in the early days of the COVID-19 pandemic. Up-to-date information on PDB structures and resources related to COVID-19 is available at RCSB.org/covid19, and new MotM articles, illustrations, animations, and flyers are created and presented on the PDB101.RCSB.org web portal as important new findings emerge from COVID-19 researchers. A recently published RCSB PDB article described PDB-related resources related to structure-facilitated messenger RNA vaccine design. Similarly, new PDB-101 resources are being used to explain what we know about the structural biology and vaccinology of poxviruses in response to concerning reports of outbreaks of monkeypox in non-endemic areas (accessible by searching PDB101.RCSB.org with the keyword “poxviruses”).

![Artistic conception of VegF signaling from the Molecule of the Month article Vascular Endothelial Growth Factor (VegF) and Angiogenesis.](doi: 10.2210/rcsb_pdb/goodsell-gallery-041)

**FIGURE 9** Artistic conception of VegF signaling from the Molecule of the Month article Vascular Endothelial Growth Factor (VegF) and Angiogenesis. Two neighboring cells are shown with cell membrane in green and cytoplasm in blue, and blood plasma at the top left in tan. VegF (magenta) arrives through the blood to the potential site of a new blood vessel and brings together two copies of VegF receptor (top center, lavender with glycosylation in yellow) to form an active dimer. Activated VegF receptor then initiates a signal cascade of kinases (yellow molecules attached to the membrane and pink molecules in the cytoplasm), leading to phosphorylation of many proteins, including cadherin (green). The phosphorylated cadherins separate making room for new blood vessels. Cytoplasmic proteins (blue) in the two cells include long actin filaments, L-shaped tRNA, a small ribosomal subunit, and many metabolic enzymes. The full image is available at PDB-101 (doi: 10.2210/rcsb_pdb/goodsell-gallery-041)
101 features on Ebola Virus published in 2014, including a MotM article, an award-winning Ebola Virus Illustration, and an Ebola Virus Protein Video, are likely to attract renewed interest in the wake of recent outbreaks in Uganda.

3 | DISCUSSION

RCSB PDB is currently in its 24th year of continuous operations, making it nearly half as mature (as opposed to old) as the PDB itself. The 50th anniversary of the PDB was celebrated in 2021 with numerous peer-reviewed publications and journal special issues and virtual scientific meetings hosted by the wwPDB Foundation and wwPDB members. Throughout its golden anniversary celebrations, there was universal acclaim for the pioneering efforts of the PDB as a vanguard of the open-access data movement and global sharing of research findings.

Today, there is broad appreciation that advances in basic and applied research depend critically on unfettered access to the research findings of the broader scientific community. Given that the vast majority of 3D biostructures currently in the public domain were generated with governmental or private philanthropic support, it is only right that this information be made freely available with no limitations on usage. Promulgation of the FAIR and FACT principles by government funders in many nations and the efforts of non-governmental organizations such as the CoreTrustSeal and the Global Biodata Coalition are playing critical roles in raising awareness of the value of open data sharing. Equally important, going forward, will be development of sustainable funding mechanisms to support global core biodata resources such as the PDB.

Looking ahead, perspectives on the intertwined futures of structural biologists, the PDB archive, the RCSB PDB, and the Worldwide PDB partnership are informed by the COVID-19 experience. Less than 1 month after release of the viral genome sequence in early 2020, the first experimentally determined structure of a SARS-CoV-2 protein—that of the main protease—was deposited to the PDB and made publicly available. Thereafter, structural biologists and wwPDB data centers worked in concert to make 3D structures of viral proteins freely available to basic and applied researchers, vaccine designers, and drug hunters. At the time of writing, there were ~2700 structures of SARS-CoV-2 proteins and hundreds of SARS-CoV protein structures archived in the PDB. Together, they helped fuel an immense effort in basic and applied virology and immunology research; supported design of two widely available, life-saving messenger RNA vaccines; and facilitated structure-guided discovery and development of a highly effective main protease-targeting drug (i.e., Pfizer’s nirmatrelvir, co-administered with ritonavir as Paxlovid) and multiple monoclonal antibodies approved for passive immunization (e.g., Evusheld, a fixed-dosed combination of tixagevimab and cilgavimab).

The words “what’s past is prologue” spoken by Antonio in Act 2 Scene 1 of Shakespeare’s The Tempest ring as true today as they did when the play was first performed on November first 1611. Evolution of structural biology as a scientific discipline and growth PDB as the first open access digital data resource in biology served us remarkably well as preparation for the pandemic. With the advent of artificial intelligence/machine learning-based de novo protein structure prediction, there is every reason to believe that structural biologists will be even more prolific, particularly when it really matters while SARS-CoV-2 and its variants of concern continue to threaten global public health. Their generous contributions of new information will ensure that the PDB continues to grow in importance as the single global open access data resource for experimentally determined 3D biostructures. As the RCSB PDB enters its 25th year of continuous operations, it remains committed to meeting ongoing challenges and delivering the ever-increasing corpus of structure information in myriad ways that will meet the needs of its diverse community of users around the world.

AUTHOR CONTRIBUTIONS

Stephen K. Burley: Funding acquisition (lead); project administration (lead); supervision (lead); writing – original draft (lead). Charmi Bhikadiya: Software (supporting). Chunxiao Bi: Software (supporting). Sebastian Bittrich: Software (supporting). Henry Chao: Software (supporting). Li Chen: Software (supporting). Paul A. Craig: Writing – review and editing (supporting). Gregg V. Crichlow: Data curation (supporting); validation (supporting). Kenneth Dalenberg: Software (supporting). Jose M. Duarte: Project administration (supporting); software (lead); supervision (supporting). Shuchismita Dutta: Visualization (equal); writing – review and editing (supporting). Maryam Fayazi: Software (supporting). Zukang Feng: Project administration (lead); software (supporting); supervision (supporting); validation (supporting). Justin W. Flatt: Data curation (supporting); validation (supporting). Sai J. Ganesan: Validation (supporting). Sutapa Ghosh: Data curation (supporting); validation (supporting). David S. Goodsell: Visualization (equal); writing – review and editing (supporting). Rachel Kramer Green: Project administration (supporting). Vladimir Guranovic: Software
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CONFLICT OF INTEREST
The authors declare no conflict of interest with this publication. The content is solely the responsibility of the authors and does not necessarily represent the official views of the National Institutes of Health.

DATA AVAILABILITY STATEMENT
wwPDB policy states that data files contained in the PDB archive are available under the CC0 1.0 Universal (CC0 1.0) Public Domain Dedication.

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