Community-driven roadmap for integrated disease maps

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Abstract

The Disease Maps Project builds on a network of scientific and clinical groups that exchange best practices, share information and develop systems biomedicine tools. The project aims for an integrated, highly curated and user-friendly platform for disease-related knowledge. The primary focus of disease maps is on interconnected signaling, metabolic and gene regulatory network pathways represented in standard formats. The involvement of domain experts ensures that the key disease hallmarks are covered and relevant, up-to-date knowledge is adequately represented. Expert-curated and computer readable, disease maps may serve as a compendium of knowledge, allow for data-supported hypothesis generation or serve as a scaffold for the generation of predictive mathematical models. This article summarizes the 2nd Disease Maps Community meeting, highlighting its important topics and outcomes. We outline milestones on the roadmap for the future development of disease maps, including creating and maintaining standardized disease maps; sharing parts of maps that encode common human disease mechanisms; providing technical solutions for complexity management of maps; and Web tools for in-depth exploration of such maps. A dedicated discussion was focused on mathematical modeling approaches, as one of the main goals of disease map development is the generation of mathematically interpretable representations to predict disease comorbidity or drug response and to suggest drug repositioning, altogether supporting clinical decisions.

Key words: disease maps; molecular biology; mathematical modeling; knowledge repository; biocuration; translational medicine; pathway representation

Introduction

The concept of disease maps emerged to bridge the domains of biological and computational research on various human disorders. In essence, these maps are representations of disease mechanisms that are both human and machine-readable [1–4]. Visual representation allows clinical and life sciences researchers to explore charted disease mechanisms, which are often complex and interconnected. Computer-tractable, standardized representation of the underlying information creates an interface to a broad range of bioinformatic workflows. As such, disease maps are an important platform with the potential to link the domains of biomedical knowledge and data, providing an intermediate step between a conceptual and an executable model.

In the recent years, the members of the Disease Maps Community (DMC) developed various disease maps resources, hand in hand with other groups around the globe. The community held its initial meeting in February 2017, hosted by the European Institute for Systems Biology and Medicine in Lyon, France. There we recognized a great potential in such type of exchange, especially because, despite different disease contexts, we face similar challenges, ranging from establishing proper tools and standards for knowledge encoding, through visualization of multidimensional data sets, to handling large and complex maps. We decided to meet regularly to help shape the direction where the project is heading. In October 2017, we held the 2nd DMC meeting, hosted by the Luxembourg Centre for Systems Biomedicine in Belval, Luxembourg. Here, we summarize this meeting, highlight important topics and outcomes of our discussions and propose a roadmap for the development of disease maps.

In this article, we first introduce the DMC and describe its rationale, mode of operation and spectrum of expertise. Next, we overview the 2nd DMC meeting, highlighting important topics and discussions of special focus. Then, we describe the milestones on the ‘Disease Maps Roadmap’, identified during a dedicated, extended discussion session during the meeting. In the last chapter of the article, we briefly summarize the outcomes and discuss further steps, including necessary standards and tools.

The Disease Maps Community

The DMC (http://disease-maps.org/) is a group of developers and users of disease maps of various human disorders, including cancer, neurodegenerative and immune diseases. The community formed to exchange experiences and to establish best practices for creation, maintenance and application of disease maps. The group is composed of biomedical and clinical researchers with expertise on particular diseases [2, 3, 5], but also of bioinformaticians, computer scientists and mathematicians working on technologies supporting curation and exploration of the maps [6–8]. Because the community involves projects at different stages of development, upcoming disease maps can benefit from the experience of developers at the advanced stage. At the same time, new disease maps bring their own unique use cases providing new perspective for the adoption of curation standards and required technology developments. At the time of writing, researchers from France, Germany, Luxembourg, UK, Portugal, Spain and Turkey take part in the DMC. The participation in the community is voluntary.

Regular meetings help to catalyze the exchange between the community members. The 1st DMC meeting allowed us to identify challenges shared across different disease maps’ projects and recognize the value of exchanging best practices. Moreover, it was apparent that we need to keep track of our efforts to best align them. Therefore, the main objective of the 2nd DMC meeting (http://disease-maps.org/events) was to bring the community up to speed about the ongoing activities, introduce new members with their projects and engage into deep discussion on challenges, potential solutions and the next steps to take. This discussion was at the heart of our meeting, and is described in detail in the following section. Participants engaged in extensive discussions on critical topics for tools, applications, curation standards and complexity management. Moreover, an entire session was dedicated to the topic of
mathematical modeling. Based on the outcome of our discussions, we outline the roadmap for disease maps development (Figure 1).

**Milestones on the ‘Disease Maps Roadmap’**

The community discussed five aspects of the disease maps, namely: (i) tools supporting the development and use of the maps, (ii) standards needed for biocuration of the content, (iii) management of complex content, (iv) application of the maps in the biomedical domain and (v) the predictive modeling of disease mechanisms. We defined a number of milestones, summarized in Figure 1. Some of them span multiple aspects of disease maps. For instance, ‘encoding and use of models’ need to be solved at the levels of tools, biocuration and modeling methodology. Complexity management and tools share milestones for ‘dynamic network layout’, while biocuration and applications both define ‘quality indicators of encoded knowledge’ as a milestone.

**Tools for map creation, visualization and exploration**

Disease maps are an emerging concept, bridging bioinformatics, molecular biology and clinical research. Appropriate tools are needed to support creation and use of the maps, including handling relevant standards for knowledge encoding, annotation and exploration. It is crucial to align new developments in this area with concrete use cases. In fact, the development of many available tools was initiated to directly address the needs of the DMC, and their further development takes into account the emerging challenges. Table 1 summarizes the tools discussed in the following text, both those already used for disease maps development and analysis, and those that offer new important functionalities.

**Constructing maps**

A key challenge in the field is the lack of tools tailored exactly to develop content for a disease map. Visual pathway editors [13, 17] that offer significant level of compatibility with Systems Biology Graphical Notation (SBGN) [11] are often used for this purpose, contributing to content reuse. Other solutions like Cytoscape-based Biological Network Manager (BiNoM) [9] or PathVisio [15] allow for importing, manipulating and exporting SBGN or CellDesigner formats. An interesting case is a graph editor yED [18] that introduced an SBGN palette, allowing drawing of graphs that look like SBGN diagrams.

Still, disease maps are frequently updated and extensively annotated knowledge repositories, and the mentioned editors have limited capabilities to support for such resources. Harmonization of curation standards (see section ‘Biocuration and knowledge representation standards’) is also difficult, as each of the mentioned tools uses its own encoding of the content, risking an inexact translation when transferring information between sources. An important development addressing this problem is the Web-based editor of diagrams encoded in SBGN: Newt [8]. The creators of Newt actively participate in the DMC, helping to shape and benefiting from the discussed roadmap. A milestone on the road toward mechanistic, modeling-oriented curation will be enabling support for the Systems Biology Markup Language (SBML) [12] (see section ‘Use of maps for mathematical modeling’) during the curation of disease maps.

**Maps exploration via Web platforms**

We also discussed how to explore and analyze the content of the disease maps. In this area, one of the first platforms for sharing disease maps as CellDesigner diagrams was Payao [16], followed by iPthways+ [4]. Their functionality was extended by tools like Molecular Interaction NETworkks VisuAlization (MINERVA) platform [6] and NaviCell [14], developed by the DMC members. They allow for visualization of large CellDesigner and SBGN diagrams using the Google Maps Application Programming Interface (API) to provide interactive annotation to maps’ elements and enable overlay of experimental data on top of these maps. Another solution for browsing large maps are various complexity management techniques such as expand–collapse and hide–show featured by the Newt pathway editor [8]. However, an open issue is the exploration and integration of simulation results from the associated models. A rough shortcut is currently available via visualization:

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**Figure 1.** The milestones of the DMC roadmap. Five groups of topics are highlighted. Tools: Software and methods supporting the development and maintenance of the maps; Biocuration standards: standards for knowledge gathering and encoding in the maps; Complexity management: methods that handle inherent complexity and facilitate visual exploration of the contents of the maps; Applications: workflows where maps can be applied to support knowledge exploration, generation of new hypotheses or support clinical decisions; and Modeling: standards and tools allowing to refine the maps into executable mathematical models.
Table 1. Summary of tools for creation and exploration of disease maps

<table>
<thead>
<tr>
<th>Tool description</th>
<th>Role</th>
<th>Web-oriented</th>
<th>Scale of maps</th>
<th>Data overlay</th>
<th>Supported standards</th>
<th>Active</th>
<th>Kinetics support</th>
<th>Used for disease maps</th>
</tr>
</thead>
<tbody>
<tr>
<td>BiNoM⁶: Manipulating disease map diagrams, Cytoscape plugin</td>
<td>Explore</td>
<td>No</td>
<td>Large</td>
<td>Yes</td>
<td>BioPAX⁹, CellDesigner, SBGN, SBMLⁱ¹</td>
<td>No</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>CellDesigner⁸⁰: Construction of process diagrams and simulations for molecular biology</td>
<td>Construct</td>
<td>No</td>
<td>Large</td>
<td>No</td>
<td>CellDesigner, CellDesigner, CellDesigner</td>
<td>No</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>iPthways⁶⁶: Visualization of pathways and process diagrams</td>
<td>Explore</td>
<td>Yes</td>
<td>Large</td>
<td>No</td>
<td>CellDesigner</td>
<td>Yes</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>MINERVA⁵⁶: Visualization and exploration of disease map diagrams</td>
<td>Explore</td>
<td>Yes</td>
<td>Large</td>
<td>Yes</td>
<td>CellDesigner, CellDesigner, CellDesigner, SBGN</td>
<td>Yes</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>NaviCell⁵⁴: Visualization and exploration of disease map diagrams</td>
<td>Explore</td>
<td>Yes</td>
<td>Large</td>
<td>Yes</td>
<td>CellDesigner</td>
<td>Yes</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>Newt⁷⁰: Construction of pathways and process diagrams</td>
<td>Construct</td>
<td>No</td>
<td>Medium</td>
<td>No</td>
<td>SBGN</td>
<td>Yes</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>PathVisio⁵⁵: Construction of pathways and process diagrams</td>
<td>Construct</td>
<td>No</td>
<td>Small</td>
<td>Yes</td>
<td>SBGN</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Payao⁴⁰: Visualization of pathways and process diagrams</td>
<td>Explore</td>
<td>Yes</td>
<td>Large</td>
<td>No</td>
<td>CellDesigner</td>
<td>No</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>yED⁵⁸: Construction of pathways and process diagrams</td>
<td>Construct</td>
<td>No</td>
<td>Medium</td>
<td>No</td>
<td>SBGN</td>
<td>Yes</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>BioPAXViz⁶⁸: Visualization of metabolic pathways</td>
<td>Explore</td>
<td>No</td>
<td>N/A</td>
<td>Yes</td>
<td>BioPAX</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>COBRA Toolbox⁶⁰: Simulation and visualization of pathways</td>
<td>Explore</td>
<td>No</td>
<td>Medium</td>
<td>No</td>
<td>CellDesigner, CellDesigner, SBML</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Escher⁵⁹: Construction and simulation of metabolic pathways</td>
<td>Construct</td>
<td>Yes</td>
<td>Medium</td>
<td>Yes</td>
<td>SBML, SBGN</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>iVUN⁷²: Visual analysis and simulation of kinetics in pathways</td>
<td>Explore</td>
<td>No</td>
<td>Small</td>
<td>Yes</td>
<td>SBML</td>
<td>No</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>NDex⁷¹: Sharing of network data for computational biology</td>
<td>Explore</td>
<td>Yes</td>
<td>N/A</td>
<td>No</td>
<td>Cytoscape⁷⁴</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Physiome Model Repository⁷³: Sharing of cellular models</td>
<td>Explore</td>
<td>Yes</td>
<td>N/A</td>
<td>No</td>
<td>CellML⁷⁵</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>SEEK⁷⁷ (FAIRDOMHub): Sharing of SBML models and datasets</td>
<td>Explore</td>
<td>Yes</td>
<td>N/A</td>
<td>No</td>
<td>SBML</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
</tr>
</tbody>
</table>

Notes: The table lists the tools that support construction and exploration of the disease maps' content, highlighting their role in the process. We indicate their capability to work over the Web (‘web-oriented’ column) and the size of the maps that they can handle (‘scale of maps’ column): large: over a thousand of elements, medium: hundreds of elements, small: under a hundred of elements. ‘Data overlay’ column indicates which tools can overlay external data set on their content. ‘Supported standards’ column lists which standard data formats are supported by a given tool. Even though ‘CellDesigner’ format is only a de facto standard, based on early versions of SBGN and SBML formats, we list it because of the popularity of the tool. Finally, we indicate which of the tools are actively developed, support reaction kinetics and are currently used for disease maps’ creation and exploration. SEEK and NDex platforms provide an automated layout of uploaded models, while BioPAXViz and Physiome model repository use layoutless formats (BioPAX and CellML), making the assessment of the scale imprecise.
e.g. the outcomes of flux balance analysis can be shown by different thickness and color of corresponding reactions on the map, as in Escher [21]. Another example, the iVUN system (interactive Visualization of Uncertain biochemical reaction Networks) [22], uses the kinetic parameters encoded in the map directly via the visualization interface to run simulations. Finally, the recently upgraded Constraint-Based Reconstruction and Analysis (COBRA) Toolbox [20] introduces a built-in visualization functionality for constraint-based modeling results and enables visualization of modeling results via the MINERVA platform. Overall, current platforms for analysis and visualization are Web-based, and with the increasing size of disease maps, it is important to ensure scalability of expensive operations such as layout and simulation. The increase of client-side computing power allows to use local resources for some work and use the Web server for heavy computations like graph layout. A milestone in the direction of in-depth map exploration will be Web-based visualization of simulation results together with the contents of a disease map, or its parts, used for the simulation.

**Integrating maps in a shared repository**

Another challenge that requires proper tools is the integration of maps into a repository. As disease maps projects mature, it is natural to break up large complex maps into smaller modules, which can be used independently or composed into the full map. This asks for a platform to manage multiple maps simultaneously, and cross-link their content. Currently, MINERVA and NaviCell offer support in creating a single hierarchical multi-modal disease map. A challenge that remains to be addressed is a repository spanning multiple disease maps, allowing us to query resources of various disease domains, either by keyword or by network neighborhood. For this to happen, we need to propose solutions for versioning and comparing different maps, also taking into account different annotations and context of particular projects with the aim to converge into the common standard of disease maps annotations and representation. Often, the lossless conversion between formats like SBML, SBGN or Biological Pathway Exchange (BioPAX) [10–12] is not possible. Therefore, it is crucial to develop a framework for a unifying notation for encoding the disease mechanisms and annotating them (discussed in the section ‘Biocuration and knowledge representation standards’), supported by converters minimizing the information loss on translation. A good step in this direction may be a repository of uniform, reusable modules and models of pathways that are common for multiple disorders, and can be used across many projects (discussed in the section ‘Map complexity management’). Efforts like FAIRDOMhub, the NDex platform and the Physiome Model Repository go in a similar direction [23, 25, 27]. The effective use of a shared repository is only possible with a powerful set of queries including graph-based ones such as shortest paths between a specified set of molecules and common target of a gene set [28]. Here, a milestone will be a translation of one or more common modules between different disease maps. Another important goal to be reached is enabling communication between different disease maps, allowing to query their resources.

**Biocuration and knowledge representation standards**

Biocuration of a disease map is a difficult task that heavily depends on the expertise of the curator. A clearly defined set of best practices can facilitate this process, similarly to protocols for construction of biomodels [29]. External resources like Gene2Disease or MalaCards, and tools like Integrated Network and Dynamical Reasoning Assembler (INDRA) [30–32] can help in organizing and referencing the disease-related knowledge integrated into a map.

**Curation standards**

A number of curation standards can help with harmonizing the content in various disease maps. Graphical notation and modeling languages like SBGN, SBML or CellML [11, 12, 26] offer good guidance in encoding molecular networks, while annotation of biological entities according to the Minimal Information Requested In the Annotation of biochemical Models (MIRIAM) guidelines is supported by the Identifiers.org infrastructure [33]. Whenever modeling-level description is available, the model structure can be automatically checked for consistency, e.g. to detect divergent reactions, or negative concentrations of molecules [34]. Continuous checks for correctness against these standards and resources are a key activity for developing useful disease maps. However, the specificity of certain disease mechanisms is often difficult to describe in a standardized manner. Encoding and annotating complex structures or specific post-translational modifications in a diagram may be challenging for the curator, when the proper balance between clarity and precision is not obvious. Thus, it is important to establish a set of quality indicators for the curated mechanisms indicating their usefulness and the precision of the underlying information.

**Map updates**

The standards mentioned above describe the format of the content. Another important aspect that requires attention is the relevance to the disease area—keeping the content up-to-date and relevant for current and upcoming analytical challenges. This requires dedicated curation effort, but also a community of users in the field who evaluate the content and assess its relevance for the disease of interest. Thus, supporting a given disease map by accompanying social networking tools, like discussion forums, may help catalyze the communication. From the computational point of view, text mining solutions may be used to identify potentially relevant mechanisms to include or review. These suggestions can be in turn discussed openly with the community, encouraging discussion and engagement. Testing such a text mining-based update system and comparing it across different disease maps may provide new ideas how to accelerate the time-consuming curation process. Additionally, this may lead to improvement of the algorithms of text mining supporting the curation, as they are tested against manually curated information.

**Knowledge representation consistency**

The DMC projects cover various pathologies and are at different stages of development. This diversity results in varying depth of curation for particular diseases and their mechanisms. For instance, knowledge about specific mutations and their mechanisms is important for the cancer field, while chronic disorders may put less emphasis on it. For this reason, the content of different disease maps should be reused with care. Molecular pathways implemented in a map for neurodegenerative diseases may be relevant in inflammatory disorders, but they might have to be modified or extended. Therefore, consistent and precise annotation is necessary for both appropriate use and successful reuse of curated content. Although platforms like MINERVA offer an annotation consistency check, the verification takes place after the curated content is uploaded to the platform. A curation tool checking for annotation consistency on-the-fly would help to avoid errors and omissions, improving
the quality of generated content and reducing the curator’s burden.

**Connecting maps to disease hallmarks**

Another challenge curators face is to design the map in such a way that end users can recognize the mechanisms of the disease and tell them apart from the normal, physiological function of a given pathway. Also, users often interpret the map based on their individual data sets, for instance for subgroups of patients, or specific cell lines.

While curating the map’s content, it is important to evaluate it methodically for the relevance to each disease area. Replication of hallmark findings in a given domain is often tangible, as many appropriate data sets are now publicly available, either via general repositories of molecular phenotypes, such as Gene Expression Omnibus [35] and the Expression Atlas [36] or disease-specific resources such as the Genomic Data Commons [37] and the Human Protein Atlas [38] in the case of cancer. Identification of differentially expressed molecules and their visualization on the map will help to refine the map’s content, but also will be a demonstration of its utility. A series of such analyses may help to calculate significance and vulnerability scores, describing how strongly a given mechanism is implicated in the disease, and how often it is perturbed. Benchmarking scenarios, describing these in silico validation experiments, are a necessary component of disease map development. Such scenarios and benchmark data sets will have to take into account the disease heterogeneity and differences in statistical approaches used for data preparation across studies.

**Map complexity management**

Disease maps aim to describe disease mechanisms, which often span across multiple scales of human physiology and involve numerous cross-talking pathways. This comes with the challenge of meaningful organization of such complex knowledge. Thus, complexity management in our case aims to resolve the perception difficulty of different scales and mechanisms without losing the understanding of the disease as a whole.

Complexity management foundations for disease maps are distilling the relevant content to the disease context, highlighting the mechanisms critical for the pathology, categorizing the mechanisms based on their general biological relevance and creating high-level, abstract views of relationships between key concepts. These approaches are used already at the stage of curating the maps’ content.

**Network complexity**

Densely connected biological networks are impossible to draw without edge crossing (nonplanar graphs). A currently applied approach is to create multiple instances of (to clone) a molecule in various contexts (different compartments, pathways or modifications), which reduces visual clutter. This task can be automated by an algorithm suggesting when to clone a certain molecule to improve overall graph perception [39]. Similarly, clearly separable modules of a disease map can be transformed into submaps, linked hierarchically to the overview map. At the same time, visualization and management of such distributed content become more difficult, as different instances of the same molecule, or separate submaps, have to be meaningfully searched and explored. Development of tools for exploration of hierarchically abstracted and modularized networks is an important milestone on the road toward managing network complexity. Testing the existing functionality of Newt for collapsing subnetworks, especially for large-scale disease maps, will help to better specify challenges in front of such tools.

Finally, we noticed that in the field of electrical engineering, which was a source of inspiration for developing standards for graphical network representations, established conventions exist for representing crossing wires on the electrical diagrams. As creating network diagrams completely free of edge crossing does not seem to be possible or useful, developing standards on resolving possible misinterpretations would be a useful step in managing complexity of large disease maps.

**Scale complexity**

Another group of complexity management techniques concerns map visualization. These include semantic zooming into diagrams [6, 7], collapsing and expanding subnetworks in a diagram [8] or bundling edges to discover structure of dense networks [40]. One important type of semantic zooming subdivides different content types among multiple layers, where the zoom level defines the level of complexity seen by the user. For instance, the highest zoom level could show the most generic physiological view, e.g. the tissue or organ affected by the disease, the zoom layer below would show cell type relationships in the tissue, while subsequent zooms would show different levels of complexity of underlying cellular and molecular networks. Visualization of these complex networks at low granularity can be facilitated by representing network motifs (commonly encountered graph structures, like phosphorylation or complex formation) as recognizable symbols, or highlighting the most relevant molecules for the disease. This hierarchical way of layered display can be complemented by ‘vertical’ layers, showing separately different classes of molecular processes, e.g. transcription, signaling or metabolism.

**Layout complexity**

Hierarchical layers allow complexity management at the overview level for easier navigation to a particular area of the map. However, when examining details of molecular processes, users need tools to disentangle dense bundles of interactions and relate the content in front of their eyes to the rest of the disease map. Display of such local views can be implemented with the help of dynamic layouts, where the wiring of the diagram is temporarily changed in the area examined by the user to better reflect current context. Interactively changing the layout on-the-fly can be foreseen for the local views because of their small size. For instance, the technique of hyperbolic trees may allow us to remove local edge crossings in an area of the map, which would be infeasible for the entire map [41]. The local topology of the network can also be adapted to minimize the curvature of locally viewed edges [42], or it can be modified to reflect the uploaded data sets. In these data-driven layouts differentially regulated molecules can become larger and more central, while flux balance analysis results may change the length of the edges to reflect the reaction rate. There are alternative methods for creating data-driven layouts of biological networks, based on nonlinear dimension reduction constrained by the network structure [43]. These and other complex graph visualization methods such as hierarchical bundling of smoothed edges [44] can greatly facilitate understanding the complex structure of connections between the objects on the map and its relation to the studied data sets.

**Managing technical complexity**

A less conceptual but not less important aspect of managing complexity of disease maps is related to technical problems, i.e.
it concerns a set of questions related to performance and interoperability.

Despite the development of a new generation of network editors, efficient manipulations needed for constructing and maintaining disease maps with thousands of nodes remain challenging. Here, one could explore the possibilities of existing approaches for complex and multiscale visualizations used in other domains such as the Web Graphics Library (WebGL). For instance, while dealing with large and complex networks, one can reuse existing methods of advanced memory caching that avoid keeping the whole complex network in memory, like it is done in Google Maps for smooth browsing of huge raster geographical images.

The interoperability between existing standards approved by the community, such as SBGN Markup Language (SBGN-ML), SBML 3.0 with Layout and Render extension and de facto standards used to construct most of disease maps, like the CellDesigner proprietary SBML extension, remains a challenge. However, this aspect happened to be relatively inexpensive to improve. For instance, at the time of writing, a new fully functional bidirectional converter from CellDesigner to SBGN-ML has been developed as a collaborative effort between DMC members (https://github.com/royludo/cd2sbgnml). Such tools will allow the use of the rich computational systems biology toolkit to analyze the existing collections of disease maps.

Applications of disease maps
The way disease maps are used drives the curation of the content and indicates directions for technology development [45]. Disease maps are created for various purposes, for instance as a didactic resource, a knowledge repository, a platform to visualize data or a collection of predictive molecular signatures. These use cases reflect different stages of development of a disease map, when its contents are continuously refined from a collection of most known mechanisms of a given disease (‘hallmarks’) through verification against established expertise and available experimental data.

Access to bioinformatic databases
Disease maps applications that focus on knowledge exploration require easy and direct access to various data resources. MINERVA and NaviCell platforms provide such access to a number of annotation sources, like HUGO Gene Nomenclature Committee (HGNC), UniProt, Chemical Entities of Biological Interest (ChEBI), PubChem or Gene Ontology [46–49]. From our experience, users can better understand representations of particular disease mechanisms if they can cross-check descriptions of the included molecules. However, advanced data interfaces are needed, such as querying pathway databases for entire sequences of reactions from Reactome or WikiPathways [50, 51]. Netw implement such functionality for drawing interactions. A corresponding feature for visual exploration of disease maps remains to be implemented.

Tissue and disease specificity
Visual navigation through complex content will be greatly facilitated by introducing visual tags for cell or tissue types on the maps. Highlighting elements or interactions unique for certain physiological environments is needed for users to disentangle complex bundles of reactions, and to understand them. Semantic zoom functionalities, already implemented to a certain degree in disease maps platforms (discussed in the section ‘Map complexity management’), need to be extended. When zooming into complex networks, the content should be presented with gradually increasing number of details, based on the complexity of underlying physiology and on the density of explored molecular networks.

Individual disease maps represent contextualized pictures of various pathologies. Comparing disease maps’ contents will help to identify deregulation of mechanisms specific to a given disorder, as well as pathways implicated in a number of pathologies. Such comparisons become tangible thanks to pipelines for data cross-linking and visualization of complex networks. Combined with patient-specific data, such exploratory analysis in maps of overlapping pathologies, like cancer subtypes, may support personalized medicine by facilitating interpretation of patient-specific drug resistance.

Health and disease data interpretation pipelines
Clinical applications of disease maps [45, 52] are close to the role of a Clinical Decision Support System, with an emphasis on exploration and interpretation of medically relevant data. Big health data, collected in great amounts by health-care providers and pharmaceutical companies, need to be structured and interpreted through visualization. This is a scenario where disease maps may provide a valuable context to large data sets, allowing meaningful filtering and summary of otherwise indigestible numbers. Initial steps in creating big health data pipelines to disease maps have been taken [45], where a disease map is used to visualize gene expression based on patients’ demographic data.

In the end, disease maps may be a great support to knowledge-based drug discovery using patients’ data, but only after drug databases can be linked with the maps’ content and supported by dedicated analytical pipelines. For instance, disease maps may become a platform for network data-driven drug response prediction. This will require identification and assessment of disease-rewired pathways, network analysis to identify a desired intervention set (target interactions or elements in the network) and mapping this intervention set back to drug databases, looking for secondary use of existing medications (drug repositioning).

The final goal of a disease map development is to become mathematically interpretable and to support clinical decisions in a given domain. Importantly, the process of refining and exploring a disease map itself provides knowledge building, even without an immediate clinical application. Although the map is created to be quantified and analyzed with data to predict a clinically relevant outcome, its qualitative interpretation can have a great value in hypothesis generation and for guiding experimental design. This is an important note to take into account when managing expectations about applications of a disease map.

Use of maps for mathematical modeling
Disease maps are currently used to organize knowledge and to visualize data. The ultimate goals are however the generation of testable hypotheses, the identification of actionable targets and the support of clinical decision making. To achieve this, executable mathematical models are required. Depending of the required level of resolution, qualitative models (e.g. logical or Boolean models), or quantitative models (e.g. ordinary differential equations, stochastic differential equations or Markov jump processes) can be used. Yet, the formulation of mathematical models requires more information than the use of maps for
visualization, and this generates additional challenges to address.

**Construction of executable mathematical models from disease maps**

The formulation of executable mathematical models requires information on molecular species and their interactions. For the formulation of qualitative models, information about the mode of interaction between molecular species is required (e.g. activating or inhibiting). This information can be extracted from SBGN Activity Flow maps [53, 54]. However, most of the available disease maps use SBGN Process Descriptions or a combination of SBGN Process Descriptions and SBGN Activity Flow diagrams. This complicates an automatic construction of a logical model substantially. For the formulation of quantitative models, information about the properties of reactions is necessary, including stoichiometry and reaction kinetics [55]. While stoichiometry should be encoded in SBGN Process Descriptions, the kinetic rate laws are usually missing. The definition of rate laws requires additional information or assumptions, e.g., that a reaction follows the law of mass action kinetics. Some efforts have been launched to generate logical and numerical models from pathway maps [56]. For instance, the ongoing work on automated translation of SBGN and CellDesigner formats into logical models may help to bridge the quantitative and qualitative applications of disease maps. However, this remains a challenging task, providing results of mixed quality. To support the construction of executable mathematical models from disease maps, the first milestone would be the definition of a standard operating procedure (SOP), which informs biocurators about the minimal information, which has to be implemented in the disease maps. In this context, the use of SBML for the model formation and automatic checking of model consistency might be more appropriate. An important issue is therefore to ensure a proper link between molecular processes and the phenotype of interest.

**Parameterization or executable mathematical models**

Quantitative mathematical models usually possess unknown parameters, e.g. binding affinities and degradation rates. To ensure that the models are predictive, these parameters have to be estimated from experimental data. This requires comprehensive data sets as well as computational methods for statistical inference.

Data sets are available in the literature and in established databases, such as BRENDAL [57] and SABIO-RK [58]. However, most literature-based data sets are unstructured and difficult to assess. Furthermore, the quality of experimental data varies heavily. A milestone for any disease map project aiming at quantitative models therefore is the establishment of a database of general and disease-specific data. The databases could be created together with the disease maps, and encode essential qualitative properties as well as quantitative data. The databases established for different projects should ideally follow common standards.

To estimate the unknown parameters from the available data, an efficient computational pipeline is required. As disease maps usually possess hundreds or even thousands of state variables and parameters, the resulting computational complexity might be challenging for established toolboxes such as COmplex P Athway Simulator (COPASI) [59], Data2Dynamics [60], Parameter EStimation Toolbox (PESTO) [61] or PottersWheel [62]. Moreover, such a large number of variables will require an automated procedure to check parameter identifiability. A milestone is the establishment of a scalable computational pipeline, which is applicable to the standardized models and databases established in the disease map projects. Such a pipeline could combine efficient objective function and gradient evaluation methods [63] with advanced parallel optimization schemes [64].

**Personalization of models using data**

A parameterized quantitative model can in principle be used for decision support in the clinic. To provide patient-specific predictions, the model needs to be personalized with patient-specific information. While this is a procedure fairly easy to do with small models, such as the ones used in pharmacokinetic/pharmacodynamic modeling, it is much less so in the case of large maps with a great number of molecular partners. In recent studies, exome and transcriptome sequencing data of cancer cell lines have been used to set cell line-specific translation rates [65, 66]. In a similar study, the mRNA expression was used to predict the survival of individual neuroblastoma patients [67]. While both approaches were successful in the respective applications, transcription rates and mRNA levels can change in response to treatment. For an analysis of the long-term response of patients, alternative strategies may be necessary. A milestone in this respect will be to develop different individualization approaches and then assess them in a range of applications. In addition, disease-related functional variants need to be implemented to benefit from comprehensive sequencing and genome-wide association studies (GWAS).

**Summary**

A ‘disease map lifecycle’, as shown in Figure 2, starts with curation and integration of knowledge about disease mechanisms. This collected knowledge, combined with experimental data and annotations from bioinformatics databases, supports better understanding of the disease and formulation of systems-level, data-driven hypotheses. The ‘disease map lifecycle’ is a dynamic process, as feedback from the interpretation of such contextually driven knowledge leads to the design of further, tailored data interfaces, permits better consolidation of knowledge within the repository and may, if validated experimentally, introduce new knowledge about disease mechanisms for further curation and incorporation into the map. The milestones of the community-driven roadmap (Figure 1) are indicated in Figure 2.

**Application example: drug repositioning**

Signaling pathways implicated in human diseases create a complex network with redundant pathways. This complexity explains frequent failure of one-drug-one-target paradigm of treatment, resulting in drug resistance in patients. To overcome the robustness of the cellular signaling network, the treatment should be extended to a combination therapy scheme [68].

Disease maps allow integrating patient high-throughput data together with the information about biological metabolic and signaling machinery specific to a given disease. This in turn may help deciphering molecular patterns specific to each patient and finding the best combinations of candidates for therapeutic targeting. A simple drug repositioning scenario may involve creating data overlays for tissue-specific gene and protein expression and their visual analysis for spatial and temporal patterns in signaling cascades encoded in a given map. As disease maps platforms [6] provide a direct interface to DrugBank [69] and ChEMBL [70], the user can browse for drugs targeting the most interesting elements of the network directly.
via the visual interface. With a number of other such resources available, like STITCH [71], KEGG Drug [72], Cancer Therapeutics Response Portal [73], Kinome NetworkX [74] or NCGC pharmaceutical collection [75], this data interface can be extended to provide more extensive drug target search results.

Moreover, the digital and standardized form of disease maps enables their network structure to be easily extracted for high-throughput computational analysis, following the workflow established by the steps of visual exploration and analysis. The members of DMC performed such analyses to find synthetically interacting genes [76], predict drug synergy [77] or suggest complex intervention sets that open a possibility of drug repositioning [52, 78].

Thematic highlight: mathematical modeling in human diseases research

The thematic highlight of the 2nd DMC meeting was mathematical modeling and disease maps. Building a computational model from a disease map is a process of transformation of a static literature-based representation into a dynamic executable format. This is important for a better understanding of how a disease progresses over time. It is also an environment where hypotheses and assumptions can be added and tested. Here, the prior knowledge (literature curation) can be integrated with newly generated data including omics data. Different types of computational models can be developed on the basis of the same pathway-based disease map. During the community meeting, we started reviewing and discussing possible approaches.

N. L. N. focused his presentation on the representation and modeling of allosteric proteins sensing calcium signals. Proteins with multiple binding sites, multiple independent features (such as binding partners, domains, conformations) and multi-subunit complexes are difficult to represent, let alone model. Trying to enumerate all molecular states leads to a combinatorial explosion of entities to model, and an even greater explosion of reactions to include. Some avenues allow to circumvent the problem, from rule-based modeling to abstract proteins representing probabilistic populations, or even implicit representations, e.g. Hill functions. Some of these approaches were illustrated by modeling Calmodulin, Calcineurin and CaMKII responses during synaptic plasticity.

J. H. presented parameter estimation methods based on adjoint sensitivities. These methods possess much better scalability properties than state-of-the-art approaches and facilitate the parameterization of large-scale models, potentially also executable models derived from disease maps. An application to a large-scale model of cancer signaling—essentially a disease map—was presented with more than a thousand chemical species and several thousands of unknown parameters [65]. J. H. demonstrated that the mechanistic model provides more accurate prediction for cell proliferation than statistical approaches.

R. M. T. F. discussed important differences between the notions of a reconstruction, a model and a map of molecular
mechanisms in human physiology. He presented the Recon re-
source [79], the most complete reconstruction of human metab-
olism to date, and how in combination with constraint-based
modeling it is used in systems-level biomedical research. The
latest version of the reconstruction, called Recon3D [80], intro-
duces structures of proteins and metabolites to the encoded re-
actions, and can be an important support to the canonical
metabolic pathways in various disease maps. As an example, he
discussed a map of mitochondrial metabolism, developed on the
basis of Recon3D, that can support Parkinson’s disease research.
A. Z. challenged the possibility of immediate use of disease
maps in mathematical modeling, suggesting that they are cur-
rently playing a role of interactive encyclopedias rather than
blueprints for chemical kinetics-based modeling of large reac-
tion networks (structural network models). He argued that the
disease maps rather reflect our knowledge in the corresponding
domains together with its incompleteness and controversy.
Thus, A. Z. coined a notion of executable encyclopedia as oppos-
ite to structural model, as a hypothetical approach based on
pragmatic middle-out mathematical modeling as opposite to
the pure bottom-up approach.

Outcomes and outlook

The 2nd DMC meeting brought together curators of disease
maps, developers of methodologies and tools and users. This
allowed us to clarify objectives and use cases, and aligned them
into a multi-lane roadmap for disease maps. The DMC will pro-
gress in parallel on several different lanes: tools, applications,
curation standards, complexity management and mathematical
modeling, at different paces, but in the same direction and with
the same goal. Importantly, there are stages of the roadmap
where the milestones align across the lanes. These will be
treated with priority by the community.

Our discussions brought up a number of resources that we,
disease maps curators and users, can benefit from. Participation
of leaders of the Physiome and Recon projects [79, 81] led to
ideas on how to capitalize on existing and well-structured
knowledge and methods they developed. We reviewed current
and upcoming interfaces to pathway databases and data ana-
lysis pipelines that will help us to curate and interpret the
maps’ content.

This productive series of meetings will continue. The 3rd
DMC meeting is scheduled for June 2018 in Paris, hosted by
Institut Curie (http://disease-maps.org/events). We aim to re-
view and update the roadmap and enlarge the community.

Most importantly, we would like to maintain the atmosphere of
collaboration and open exchange within the community, which
is the key to improvement and further development of the
Disease Maps Project. There are several tools, approaches and
platforms developed by DMC members. Exposure of the partici-
pants to these resources will allow active exchange of know-
how, and parallel hands-on tutorials will be provided.

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