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Introduction

In the 19th century, brilliant pioneers like Mendel, Darwin and Semmelweis required little more than meticulous observation of their surroundings for developing their theories on inheritance, evolution and infectiology that would become the foundations of modern biology and medicine. Since then, a range of sophisticated technologies have been developed that now allow us to observe biological systems at molecular resolution. However, our rapidly growing knowledge at the molecular level
also revealed the fundamental limitations of traditional reductionist approaches that aim to understand complex biological systems by dissecting their individual elements (Greene and Loscalzo, 2017; Stéphanou et al., 2018). It is becoming increasingly clear that many system-wide phenomena cannot be understood in this fashion, and that often the ‘whole is more than the sum of its parts.’ It is thus essential to systematically study not only the isolated elements of these systems, but also their interactions. These interactions are key to understand the emergence of novel properties and behaviors, in particular when moving across different scales, i.e., from molecules to cells, tissues, organs and organisms or entire populations (Fig. 1). Complex networks provide a natural framework for systematically investigating the various relationships between the constituents of biological systems within and between scales.

In the following we aim to provide an overview of how tools and concepts from network theory may help address important fundamental and practical challenges in biology and medicine. We start by reviewing key relationships between structural network properties and functional characteristics of the represented biological system. We then introduce frequently used networks, from the molecular level of protein-protein interactions within the cell, all the way to the level of transportation networks that span the globe. We conclude by highlighting a few future challenges in this highly active and dynamic research field.

**From Network Structure to Biological Function**

Network theory provides a versatile and general toolbox for investigating complex systems composed of interacting elements. In the most generic case, each element of the system is represented by a node and each interaction between a pair of nodes by a link (Fig. 2). In mathematical terms, the collection of all nodes and links is also called a ‘graph.’ This simple definition can be extended, for example by adding weighted or directed links, by including time-dependence or different layers of connectivity between the nodes, resulting in so-called weighted, directed, temporal or multi-layer networks, respectively.

Networks can be characterized at different levels, ranging from the level of individual nodes (e.g., their number of connections, or their centrality within the network) to the level of groups of nodes (e.g., their connection density), to the global level of the entire network (e.g., the distribution of the number of connections per node across all nodes in the network). The finding that these properties can be associated with important biological characteristics makes network theory a valuable tool in biology and medicine. For example, proteins that are located at a highly central position within molecular interaction networks have been shown to perform important roles in the cell, whereas more peripheral proteins are often less essential (Piñero et al., 2016; Costanzo et al., 2019). Densely interconnected groups of nodes correspond to functionally closely related groups of proteins (Barabási et al., 2011). Similarly, genes that are associated to the same disease tend to aggregate in specific disease modules within molecular networks (Menne et al., 2015; Chiassian et al., 2015).

**Random Networks as Reference Models**

In order to assess the magnitude and statistical significance of an observed network characteristic, suitable random controls are needed. Network theory provides a wide range of well-studied random graph models that can be used as reference (Albert and Barabási, 2002; Piñero et al., 2016). The most basic model is the classic random graph, in which a given number of links is distributed randomly among a given number of nodes (Erdős and Rényi, 1960). In contrast to many real world networks,
including most molecular networks, random graphs do not contain highly connected nodes, so-called hubs, underlying their importance as they could not have emerged by chance alone. More advanced reference models can be introduced by keeping additional properties of the original network constant during the randomization procedure. Rewiring algorithms, for example, keep the number of links per node \( f_i \) fixed and have been used to uncover basic design principles of gene regulatory networks, such as the tendency of highly connected hubs to avoid each other (Maslov and Sneppen, 2002) or the discovery of network motifs, i.e., recurrent connectivity patterns among small groups of nodes (Milo et al., 2002).

Also dynamic aspects of networks, such as their growth over time, can be assessed through random models. An important class of networks that are characterized by the presence of hubs are so-called scale-free networks (Albert and Barabási, 2002). These networks emerge by iteratively adding new nodes to the network, such that they have a tendency to form links with already highly connected nodes. The basic model can be extended by adding a latent fitness to each node (Bianconi and Barabási, 2001) to investigate the role of evolutionary processes and positive selection of genes with high fitness in shaping the structure of molecular networks.

### Molecular Interaction Networks

The first layer of information transfer and transformation from genotype to phenotype is mediated by molecular networks within the cell. In analogy to the genome representing the blueprint for all molecular components, the collection of all their interactions...
is referred to as the interactome. The interactome thus represents the blueprint for the collective functions that emerge from interactions between individual components. Most commonly, the term interactome is used specifically for protein-protein interaction (PPI) networks. Over the last two decades, genome-scale PPI networks have become available for a variety of species (Alanis-Lobato et al., 2017; Oughtred et al., 2019). PPIs can be mapped out systematically using yeast two-hybrid approaches (Rolland et al., 2014) or mass spectrometry based methods (Huttlin et al., 2017). The most comprehensive PPI networks also incorporate results compiled from numerous small-scale experiments from the literature (Oughtred et al., 2019) or computational predictions (Kovács et al., 2019).

PPIs are not the only molecular interactions within the cell that are biologically relevant and can be experimentally assessed. Other important molecular networks include metabolic networks and signaling cascades (Choudhary and Mann, 2010; Fabregat et al., 2016). Moreover, links may also represent indirect relations, for example in gene regulatory networks, where one gene can act on another via transcribed RNA or a translated protein, resulting from the binding of transcription factors and regulatory elements to the genetic material, which can be assessed experimentally through chromosome conformation capture techniques (Babaei et al., 2015).

Over the last two decades, numerous relationships have been uncovered between the structural characteristics of molecular networks and the function of the systems that they represent. In PPI networks, for example, connection patterns such as the number of interaction partners, network distance between proteins or densely interconnected network neighborhoods are directly related to biological functions in both healthy and disease states (Barabási et al., 2011; Caldera et al., 2017; Meyer et al., 2018). In a network context, disease states can often be identified with localized perturbations of the underlying molecular network. Such perturbations may be internal, for example genetic mutations associated with severe hereditary diseases (Köhler et al., 2019), or external, such as chemical or other environmental exposures (Kalia et al., 2019). Collectively, the set of all such external factors is called the exposome. Given the broad nature of this term, it is unclear whether a comprehensive mapping of the exposome and its impact on the interactome is at all achievable. First attempts in this direction focus on specific exposures, for example the impact of toxicants on metabolite networks (Kalia et al., 2019; Veneman et al., 2017). These approaches enable on the one hand the inference of which chemicals a system has encountered, and offer on the other hand an opportunity to elucidate the response mechanisms following a particular exposure.

**Beyond Physical Interactions**

The links in the molecular networks discussed above represent physical interactions that can be directly measured. In addition to these physical networks, we can also construct functional networks, where links represent more indirect relationships or similarities. The most commonly used functional networks are co-expression networks, where two genes are linked if their expression levels were found to be correlated across different experimental conditions (Saha et al., 2017). Other important examples are genetic interaction networks, where a link between two genes indicates that the phenotype of their combined knock-out deviates from the expectation based on the individual knockouts (Costanzo et al., 2016; Rauscher et al., 2018), drug-drug interaction networks, in which links connect non-additive drugs (Caldera et al., 2019) or chemical networks, in which compounds are linked based on structural similarity (Lo and Torres, 2016).

Functional networks may also contain various types of nodes, connecting for example genes and drugs: A genome-wide screen in Saccharomyces cerevisiae has recently been used to map out the interactions between 1377 chemical compounds and 177 genes (Piotrowski et al., 2017). Systematically exploring pairwise combinations of cellular perturbations has great potential for functionally annotating individual components, such as genes, drugs or environmental factors, as well as for identifying the involved molecular pathways and, more generally, for elucidating the fundamental rules that underlie the cellular response to combinations of perturbations.

Similarity networks were further used to study individual exposomes, by connecting co-occurring species or chemicals, revealing temporal and environmental patterns such as compounds that were released simultaneously during rainy days (Jiang et al., 2018). This exemplifies the potential to investigate different aspects of a biological concept (here the exposome) through complementary network approaches.

**From Molecules to Organisms**

While biological processes span a wide range from molecules to cells, tissues, organs and whole organisms, the networks at the molecular level are the most studied and best understood at this point. This reflects their importance as the primary interface between genotype and phenotype, but also the fact that they are more easily accessible experimentally compared to other relevant networks.

At the level of cellular organization, the neural networks that constitute the nervous system have probably received most attention (Bullmore and Sporns, 2009). Considerable efforts are made to systematically map out neural networks, in humans, as well as in model organisms. Similar to the different types of molecular networks introduced above, neural networks may also either represent direct cellular networks, where nerve cells are connected through synapses, or functional networks, in which regions of the brain are linked if they show correlated patterns of activity. The first complete direct neural network was resolved as
early as 1986 for the worm *Caenorhabditis elegans* (White et al., 1986). For higher organisms, only partial maps are available to date, for example in mice (Briggman et al., 2011; Bock et al., 2011), but also in human (Glasser et al., 2016), if only at a very coarse grained level. Mapping out the complete human ‘connectome’ of all our brain cells will likely remain out of reach for many years due to its staggering size (Sporns, 2013).

A similarly complex and important biological system is the immune system, whose primary objective is to maintain the normal function of an organism under constant threat by internal and external challenges, ranging from tumor cells to viral infections. Given the diversity of participating organs, cell types and molecules, it has been proposed to conceptualize the immune system as a multi-layered network (Bergthaler and Menche, 2017; Rieckmann et al., 2017; Kveler et al., 2018). The nodes in this network represent cells, links represent communication through signaling molecules, such as cell-surface receptors or secreted molecules. Different layers may represent different contexts, such as organs, tissues or activation status.

**Global Networks in Epidemiology**

An effective response to a viral or bacterial infection is not only critical for individual organism, but may also be seen in the much larger, potentially world-wide, context of epidemics. To accurately model the spread of a contagious disease, we must understand both the social networks of personal interactions, as well as the local and global transportation networks along which people travel (Pastor-Satorras et al., 2015). Interestingly, it was shown that not only infectious diseases are transmitted across networks of social interactions, but also other sociological and health-related conditions, such as smoking behavior (Christakis and Fowler, 2008), weight gain (Christakis and Fowler, 2007) or happiness (Fowler and Christakis, 2009).

Mathematical models have a long history in epidemiology and date back to the early 20th century (Kermack and McKendrick, 1927). Classical models divide a population into three compartments, in which people are either susceptible to an infection (S), currently infected (I), or recovered (i.e., immunized) or otherwise removed (R) from the susceptible pool. The temporal dynamics of these SIR models can be described by differential equations. Early models typically assumed ‘uniform mixing,’ i.e., an equal probability for any infected individual to contaminate any susceptible individual. More recently, these models have been significantly improved by considering the relevant social and transportation networks that underly the disease spreading process (Wang et al., 2017). For instance, the structure of the face-to-face contact network has a profound impact on how fast and how far a contagious disease may spread among a population (Pastor-Satorras et al., 2015). Likewise, the efficiency of different immunization strategies can only be fully understood when taking these networks into account.

Smallpox is currently the only human infectious disease that was successfully eradicated through immunization. There are two key aspects for the success of an immunization campaign: First, the effective access to immunization, which includes the availability of a vaccine, but also the individual willingness to get vaccinated. The latter may decline along with the disease prevalence, even in countries where immunization is compulsory. It has been shown that a better understanding of the herd effect improves the adherence to such programs (Brockmann, 2017; Betsch et al., 2017). Second, the impact that an immunization of a certain subpopulation has on the spreading of the disease, i.e., how it reduces the contagion rate in total, as well as within smaller subcommunities. This aspect can be studied from a network theory point of view. The structure of the social face-to-face contact network determines whether bottlenecks (such as high centrality nodes or links) could prevent a disease outbreak and whether some communities are more at risk than others. This enables the design of more efficient quarantine strategies with maximal impact on the social connectivity. In the past, epidemiological modeling was essential for example in handling the avian influenza outbreak in 2005 (Longini et al., 2005). More recently, great efforts have been made to profile and constrain the spread of the 2019-nCov virus, including charting the phylogeny of viral samples (Hadfield et al., 2018) and mapping the spreading risk based on global transportation networks, which allow for predicting disease spread much more accurately than maps based on geographic distance (Brockmann and Helbing, 2013).

**Summary and Outlook**

Networks provide a powerful framework for investigating biological systems ranging from the molecular to the global scale. A key factor for the success of network theory in biomedical applications is that many structural network characteristics can be related to functional properties of the respective biological system. In molecular networks, for example, densely connected node communities often correspond to proteins involved in a particular cellular process. Likewise, disease associated processes can be identified with specific connectivity patterns between groups of perturbed nodes.

An important open question in this context is how exactly different network perturbations influence each other. For example, it has been found that a network overlap between a drug-induced perturbation and a disease associated perturbation may either indicate an effective treatment of the respective disease, but also the opposite, namely that the disease may be a side effect of the treatment (Cheng et al., 2019; Guney et al., 2016). This highlights an important methodological and conceptual limitation of current network approaches. We are still lacking a systematic understanding of the combined effect of independent perturbations, in particular when considering complex phenotypes. To date, most large-scale experimental efforts for elucidating the combined effect of perturbations relied on relatively simple, one-dimensional readouts, such as
growth assays, for example in the characterization of genetic interactions (Kuzmin et al., 2018) and drug-gene interactions in yeast (Piotrowski et al., 2017). More recently, more informative readouts have been employed as well, such as high-content imaging or next-generation sequencing, which allow for a much more detailed assessment of the interactions between different perturbations. Using these high-dimensional readouts it is possible to identify different types of interactions between perturbations (positive, negative), as well as their direction. Furthermore, high-dimensional readouts allow for the identification of interactions that lead to the emergence of entirely new phenotypes. The first studies aiming to map out such high-resolution ‘perturbome’ networks were based on morphological changes induced by combinations of genetic perturbations in a model organism (Fischer et al., 2015) and combinations of drug perturbations in cell lines (Caldera et al., 2019), respectively.

Another major focus of recent network-based biomedical research is the integration of the diverse data describing different levels of biological organization. While combinations of different ‘omics’ data, e.g., genomics, transcriptomics, proteomics, metabolomics and microbiome data, are becoming more and more common in basic research, their translation into clinical applications is still scare (Karczewski and Snyder, 2018), despite their potential for applications in P4 medicine being widely recognized (Apweiler et al., 2018). Network approaches can offer valuable contributions to solving current technical and conceptual challenges in integrating multi-omics and multi-scale data (McGillivray et al., 2018). Indeed, concrete translational impact is the ultimate ambition of network biology and network medicine.

See also: Metabolic Systems. Structure and Function in Complex Biological Networks

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Useful resources to build or annotate biological networks

http://cbdm-01.zdv.uni-mainz.de/~mschaerfer/hippie/
https://thebiogrid.org/
http://www.disgenet.org/
https://gtexportal.org/
http://snap.stanford.edu/data/