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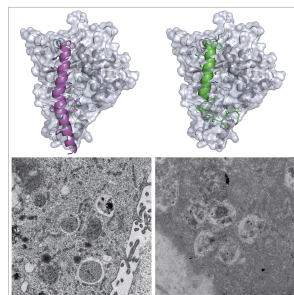
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Autophagy Regulatory Network — A systems-level bioinformatics resource for studying the mechanism and regulation of autophagy

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Abbreviations: miRNA, microRNA; NHR, nuclear hormone receptor; PPI, protein-protein interaction; TFs, transcription factors.

Autophagy is a complex cellular process having multiple roles, depending on tissue, physiological, or pathological conditions. Major post-translational regulators of autophagy are well known, however, they have not yet been collected comprehensively. The precise and context-dependent regulation of autophagy necessitates additional regulators, including transcriptional and post-transcriptional components that are listed in various datasets. Prompted by the lack of systems-level autophagy-related information, we manually collected the literature and integrated external resources to gain a high coverage autophagy database. We developed an online resource, Autophagy Regulatory Network (ARN; <http://autophagy-regulation.org>), to provide an integrated and systems-level database for autophagy research. ARN contains manually curated, imported, and predicted interactions of autophagy components (1,485 proteins with 4,013 interactions) in humans. We listed 413 transcription factors and 386 miRNAs that could regulate autophagy components or their protein regulators. We also connected the above-mentioned autophagy components and regulators with signaling pathways from the SignaLink 2 resource. The user-friendly website of ARN allows researchers without computational background to search, browse, and download the database. The database can be downloaded in SQL, CSV, BioPAX, SBML, PSI-MI, and in a Cytoscape CYS file formats. ARN has the potential to facilitate the experimental validation of novel autophagy components and regulators. In addition, ARN helps the investigation of transcription factors, miRNAs and signaling pathways implicated in the control of the autophagic pathway. The list of such known and predicted regulators could be important in pharmacological attempts against cancer and neurodegenerative diseases.

Introduction

Since the discovery of autophagy in the 1960s, and the discovery of autophagy-related genes in yeast in the 1990s, our knowledge of the regulation of autophagy expanded significantly. Major post-translational regulators of the autophagic machinery are well known, compared to the transcriptional and post-transcriptional regulators, where only limited information is available currently.^{1,2} Autophagy is essential in homeostasis

and stress-response as well as in macromolecular turnover and development.³ Both its insufficient and overdriven functions can hinder cell survival.⁴ Thus, the regulation of autophagy is critical, with high medical importance. The autophagic machinery, consisting of a complex interplay between more than 30 initiator and executor proteins, must be under constraints of precise, context-dependent and systems-level regulatory mechanisms at post-translational, transcriptional, and post-transcriptional levels.

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The proteins involved in the process of autophagy are organized into interacting complexes, having different functions in the autophagic process (e.g., initiation, membrane sequestration, and in targeting the materials to degrade in the forming phagophore). Most of the interactions within the core machinery of autophagy are well known, however, there are some unanswered questions that are needed to be resolved in order to better understand the mechanism.⁵ Interestingly, direct connections between the initiation and execution complexes were only found in the past year: it has been proved that ULK1/2, the major initiator could activate autophagy by phosphorylating another key autophagic protein, BECN1/Beclin 1.⁶ A similar important finding was obtained from yeast, where Atg1 (yeast ortholog of ULK1/2) phosphorylates Atg9 and Atg2, enhancing the membrane trafficking to the phagophore assembly site.⁷ These recent and key findings indicate that post-translational regulation of autophagy could still provide unexpected and undiscovered connection with high evolutionary or biomedical relevance. To facilitate such discoveries in silico, structure-based predictions could guide experimental researchers to validate and identify such connections.

There is no doubt that post-translational regulation of autophagy is only one part of the story. Autophagic activity also depends on the expression of autophagy-related genes and is regulated by certain transcription factors (TFs) and microRNAs (miRNAs).^{1,2} These regulatory influences can be realized on different time scales, be driven by external signals, and constitute feedback loops. Considering the transcriptional regulation of autophagy, some elements have already been highlighted in the literature, such as the transcription factors TFEB, FOXO, and SREBFs/SREBPs.⁸⁻¹⁰ By modulating autophagy, these TFs take part in the cellular response to starvation, stress, or lipid depletion, and are also involved in the pathomechanism of several diseases.^{1,11} TFEB is activated upon starvation, and facilitates the transcription of many autophagy and lysosome related genes and maintains the regeneration of lysosomes.¹ FOXO1 and FOXO3 act as effectors of the insulin signaling pathway, to regulate autophagic activity.^{1,11} Analogously, SREBF2/SREBP2 activates autophagy in case of sterol depletion.¹⁰ Beyond the role of the few TFs extensively examined and highlighted in the literature, further transcriptional regulatory components are expected to regulate autophagy in certain context. Given the advances of novel high-throughput techniques in protein-DNA interaction discovery, such as ChIP-Seq, PBA, and SELEX,¹²⁻¹⁴ numerous candidate TFs have been discovered. In addition, with resources containing TF binding site information, like JASPAR,¹⁵ potential target genes for a given TF can be predicted on a genome-wide scale. One may think that the current limitation in the search for autophagy regulators is the available data and computational expertise to evaluate and analyze data sets.

Several miRNAs downregulate mRNAs of autophagy-related genes by specific binding. However, little is known about their systems-level role. A recent review listed more than 16 miRNAs regulating autophagy genes post-transcriptionally.² These miRNAs are able to block specific steps of autophagy (e.g., *MIR376B* acts on *ATG4* and *BECN1*, while *MIR630* acts on *ATG12* and *UVRAG*). Remarkably, most of these miRNAs affect

the early stage of autophagic vacuole formation, possibly because this way miRNAs could prevent the accumulation of autophagosomes.² The growing number of experimental data on miRNA-driven regulation necessitates repositories for the post-transcriptional regulation of autophagy. Such resources could facilitate our understanding on the context-dependent role of these regulators.

The importance of identifying such context-dependent regulators is also supported by the fact that autophagy is a promising therapeutic target in several pathologies, especially in cancer and neurodegenerative diseases.¹⁶ Because autophagy has an ambiguous role in cancer, described by the 'double-edged sword' metaphor, therapies targeting the process need to be specific and context-dependent.¹⁷ Considering the complexity of autophagy and its regulation, searching for therapeutic targets without a systems-level analysis is like looking for needle in a haystack. The first step on the way to investigate the regulation of autophagy as a system is to collect all the available knowledge, including all levels of regulation. Currently elements of this knowledge are scattered in huge number of articles and bioinformatics resources, like databases of protein-protein interactions, transcriptional regulation, or post-transcriptional regulation.¹⁸ An integrated and precisely compiled interaction network could allow mapping feedback loops at all levels of regulation; to investigate differences by tissue, physiological or pathological state, drug effect, or gender; to build models using different mathematical formalisms, and thus simulate different conditions, and verify the models experimentally.

Until now few systems-level resources about autophagy have been published. The Human Autophagy Database¹⁹ (HADb) is a collection of 234 autophagy-related genes, containing references to major genome and protein databases.¹⁹ It does not intend to provide an interaction network, so it completely lacks interaction data. Another database named Autophagy Database²⁰ (ADB) contains orthologs from 40 species, and gives a comparative list of them, including a total of 206 proteins in human. For some proteins, it also collects a list of interactions—641 interactions in human—but the sources of those data and the scope of the collection is not clearly defined. A large-scale LC-MS (liquid chromatography and mass spectrometry) study provided a network of 751 interactions between 409 autophagy-related proteins.²¹ The advantage of this dataset is the uniform methodology and the relatively wide range of proteins involved in the study. However, this resource contains only the interactions detectable by the LC-MS method, and omits other interactions described in the literature. The 2 mentioned autophagy-focused resources lack data on transcriptional and post-transcriptional regulation.

Prompted by the lack of a proper bioinformatics database that extensively collects available data from the literature, from protein-protein interaction databases, and prediction methods, and contains data on several levels of regulation, we developed Autophagy Regulatory Network (ARN; <http://autophagy-regulation.org>), a novel resource to help both in silico and wet lab researchers in their investigation of the human autophagic process.

Results

The Autophagy Regulatory Network (ARN) database

The ARN database (<http://autophagy-regulation.org>) contains proteins involved in the mechanisms of autophagy, their regulators, and their TF and miRNA regulators as well as connections between all these components and signaling pathways (Fig. 1). Six main layers build up the structure of ARN: (1) autophagy proteins, (2) their direct regulators from autophagy specific resources, (3) post-translational regulators that directly regulate proteins in the first 2 layers, (4) transcriptional regulators of the first 3 layers, (5) post-transcriptional regulators of the first 4 layers, (6) signaling pathways and protein-protein interactions connecting pathways to autophagy regulators. ARN contains interactions from manual curation, 19 external databases, and 4 prediction methods (listed in Table 1). For basic statistics, please see Figure 2. Users are able to filter interactions by sources, and use resources in a comparative way, according to their requirements. Interactions may have confidence scores, users can filter by the data set, setting preferable level of confidence, using the customizable download module.

The ARN website

ARN's website is available at <http://autophagy-regulation.org>. The website is designed to give a comfortable way to browse interactions, providing hyperlinks to original sources and PubMed references of each interaction. The download section of the website gives an opportunity to customize the data to download: select between layers, and filter interactions by source, or by confidence score.

The search field on the main page autocompletes the search term, and understands several different database IDs and accession numbers. If the search is successful, the page navigates to the datasheet of the selected protein. The protein datasheet shown in Figure 3 illustrating the interactions of a key autophagy protein, BECN1, contains 4 main sections. At the top of the page, in a box the full name, gene name, UniProt ID and Ensembl ID of the protein are available. Below the names, a list of related diseases and cancer types can be found. On the left side, a list of interactions enumerates all the first neighbors of the protein, grouped by layers. The lists of the layers are expandable, and within these lists, detailed information (e.g., sources, references, confidence scores) can be obtained about an individual interaction. Below the list of the interactions, the connections between

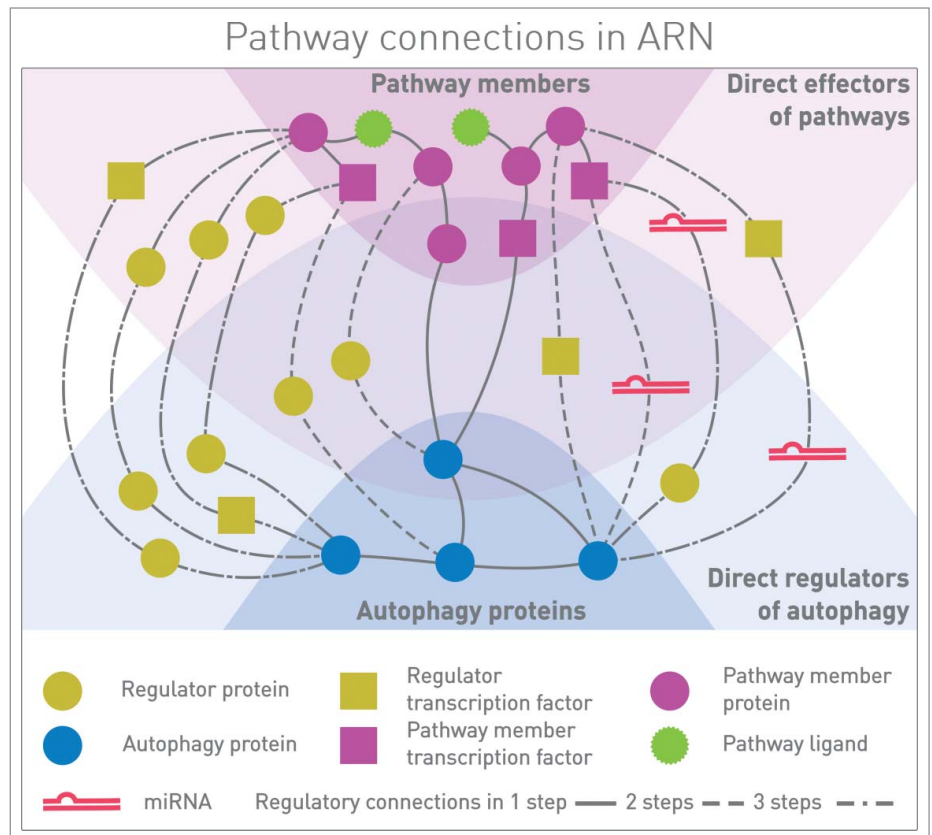


Figure 1. Connections between autophagy components and signaling proteins in one, 2 or 3 steps. One-step connections are direct protein-protein interactions (PPIs), or a pathway member TF regulates the transcription of an autophagy protein. Two-step connections also can include PPIs and TF-gene interactions, but TF-miRNA-mRNA interactions as well. Three-step interactions are combinations of all these types of interactions, involving 4 molecular species. In this representation, signal is coming from the signaling pathway receptors binding ligands, toward the proteins executing autophagy. By analyzing the whole network, feedback circuits and network motifs can be identified along the paths.

signaling pathways and the autophagy system are listed. We defined this pathway connection either in one or 2 steps, where one of the proteins is a member of a given pathway, and the other one is a present in the ARN database. On the right side of the protein datasheet, an interactive view of the first neighbors' network is presented. In this view, interactors can be filtered by layers of ARN, and users are able to get more information on proteins and interactions by clicking on them (Fig. 3).

Comparison with other resources

Compared with general protein-protein interaction (PPI) databases, BioGRID⁴⁴ contains 76 interactions between 30 autophagy proteins, while in IntAct⁴⁶ 136 interactions between 34 autophagy proteins can be found. ADB²⁰ contains 114 interactions between 31 autophagy proteins. ARN as an integrated resource contains 238 interactions between 38 autophagy proteins. Note that nearly all of the PPIs in ARN are present in other sources but it is ARN that contains them together in a single resource. Thanks to our manual curation we could increase the number of well-referenced interactions with 18, which are not present in the other sources.

Table 1. The data sources of the Autophagy Regulatory Network

Type of interactions	Data sources
Protein-protein interactions	ADB ²⁰
	ARN manual curation
	Behrends et al. ²¹
	BioGRID ⁴⁴
	HPRD ⁴⁷
	InnateDB ⁴⁵
	IntAct ⁴⁶
	ELM based prediction ³⁸
	Prediction based on domain-domain interaction
	Signalink 2.0 manual curation ⁵⁵
Transcriptional regulation	ABS ⁴⁰
	ARN manual curation
	ENCODE distal ⁴¹
	ENCODE proximal filtered ⁴¹
	HTRI ²⁹
	JASPAR ¹⁵
	ORegAnno ⁴²
	PAZAR ⁴³
	miR2Disease ⁴⁸
	miRDeathDB ⁴⁹
miRNA-mRNA interactions	miRecords ⁵⁰
	miRTarBase ⁵¹
	TarBase ⁵²
	ENCODE ⁴¹
	PuTmiR 1.1 ⁵³
	PuTmiR 2.0 ⁵³
Transcriptional regulation of miRNAs	TransmiR 1.2 ⁵⁴

ARN contains data from manual curation and from 23 external resources. From SignaLink we included 3,287 manually curated interactions; we used 4 prediction methods in ARN (domain-domain based prediction (using data from Pfam,³⁹ DOMINE⁶⁰ and Negatome,⁶¹) domain-motif based prediction using the structure filter for ELM,³⁸ TF-promoter binding prediction using the JASPAR¹⁵ algorithm, and TF-miRNA gene regulation from PuTmiR⁵³); the remaining 18 databases, including all the miRNA-mRNA data sets, contains data mainly from high-throughput experiments.

Similar comparison with transcriptional and post-transcriptional resources is shown in **Table 2**. Note that many of these connections might be false positives or highly context specific.

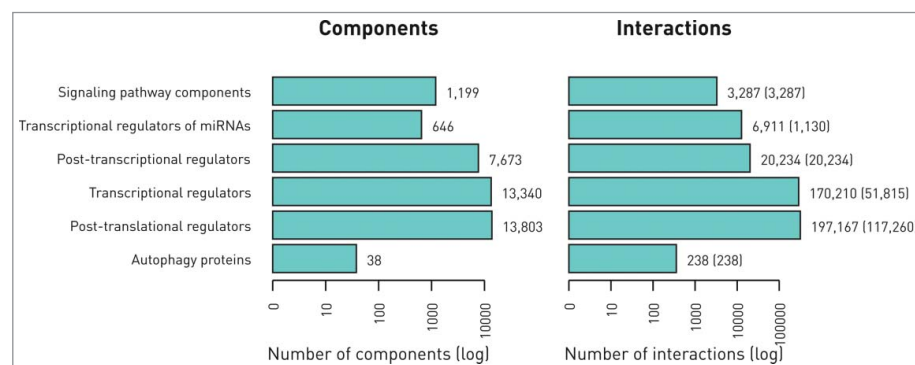


Figure 2. Basic statistics of ARN. Number of components (A) and interactions (B) in different layers of ARN is shown. The numbers of experimentally verified interactions are indicated in parenthesis next to the total number of interactions, which also includes predicted ones.

However, similarly to PPI predictions, these potential connections could also serve as a pool of possible autophagy-related regulatory mechanisms that should be examined and confirmed experimentally. The ARN resource contains 98 known and predicted TFs for 37 autophagy genes with 557 TF-gene connections; 35 of them are manually curated, and cannot be found in other resources. Of note, we found only a few TFs present in multiple bioinformatics resources, indicating the importance of different approaches to discover TFs capable to regulate autophagy, and the usefulness of ARN as an integrated single resource.

We extracted the interactions relevant in the regulation of autophagy from all constituting databases, while we have integrated different types of molecular interactions (protein-protein, TF-gene, miRNA-mRNA) into a uniform data scheme. Overall, ARN contains more regulatory interactions for the autophagy proteins than any of the constituting databases. Interactions from 23 sources have been integrated into one comprehensive database, giving the opportunity for comparison and selection between the data sources. Note that the total numbers in each ARN layer at **Table 2** are higher than any of the sources. This indicates the increased amount of data in ARN, compared to other resources.

Application

ARN can be used to examine the autophagy system in humans for both a global analysis or for gene-specific studies. For both cases, different levels of the regulation can be examined, validated or experiments can be evaluated. Here, we highlight another key feature of ARN that is its immersive connection with signaling pathways: ARN connects autophagy proteins directly and indirectly with 7 major signaling pathways taken from SignaLink 2. We included all connections up to 3 steps (4 elements) length, considering PPIs, TF-gene, and miRNA-mRNA interactions as well. There are 357 direct connections between pathway member proteins and autophagy proteins, indicating the robust and context specific regulation of autophagy by signaling pathways. On the **Figure 4** one- and 2-step long connections between pathways and autophagy components are shown. This is a global map that could be specifically analyzed or zoomed in by users who download ARN.

In the following, we illustrate the power of multilayered connection between autophagy and signaling pathways with the example of the nuclear hormone receptor (NHR) pathway. Most of the transcription factors regulating autophagy proteins belong to the NHR pathway. Using ARN data, we found potential androgen or estrogen receptor binding sites in the promoters of 2-thirds of the autophagy proteins (32). Though gender differences at the level of autophagy are observed in many diseases, little is

Table 2. Basic statistics of the Autophagy Regulatory Network

Data sources and layers	Identical nodes	Identical edges
Core autophagy proteins	38	238
ADB ²⁰	31	114
ARN manual curation	20	26
Behrends et al. ²¹	26	38
BioGRID ⁴⁴	30	76
ELM-based prediction ³⁸	16	64
HPRD ⁴⁷	11	11
InnateDB ⁴⁵	28	40
IntAct ⁴⁶	34	136
Post-translational regulators	13,803	197,167
ADB	206	381
ARN manual curation	47	46
Behrends et al.	398	441
BioGRID	12,051	71,496
Domain-domain based prediction	166	1,138
ELM-based prediction	937	78,824
HPRD	7,290	29,617
InnateDB	2,816	6,125
IntAct	9,862	45,745
Transcriptional regulators	13,340	170,245
ABS ⁴⁰	23	14
ARN manual curation	31	35
ENCODE ⁴¹	2,209	9,217
HTRI ²⁹	12,209	39,477
JASPAR ¹⁵	12,813	119,873
ORegAnno ⁴²	908	932
PAZAR ⁴³	1,940	3,018
Post-transcriptional regulators	7,633	20,186
miR2Disease ⁴⁸	171	124
miRDeathDB ⁴⁹	126	108
miRecords ⁵⁰	664	760
miRTarBase ⁵¹	7,203	19,177
TarBase ⁵²	1,798	2,584
Transcriptional regulation of miRNAs	646	6,911
ENCODE	195	590
PuTmiR 1.1 ⁵³	413	3,034
PuTmiR 2.0	288	3,095
TransmiR ⁵⁴	291	542
Signaling pathways and interactions	1,199	3,287
SignaLink 2 manual curation ⁵⁵	1,199	3,287
BioGRID	535	697
HPRD	481	611
InnateDB	9	51
IntAct	12	69

Data sources of each layer are listed with the corresponding number of nodes (i.e., proteins or miRNAs) and edges (i.e., protein-protein interactions, TF-gene, miRNA-mRNA, or TF-miRNA regulatory connections). The number of identical nodes shows both connecting component pairs (i.e., TFs and target genes as well). For each major layer we highlighted the total number of nodes and edges in ARN that is generally less than the sum of the components due to the overlap among the resources. Note that the highlighted numbers in each layer are higher than those in any of the sources.

known about the mechanisms underlying this phenomenon.²² For example, in cardiomyocytes and neurons, following ischemia and reperfusion, autophagy mediates in part the cytoprotective effect of estrogen,²³⁻²⁵ resulting in a higher level of apoptosis in males.^{26,27} Also in neurodegenerative diseases, gender differences in autophagy have been described.²⁸ In addition, almost all neurodegenerative diseases have higher incidence in females.²² At certain prostate cancer cell types, androgen signaling plays a

cardinal role in the choice between autophagy and apoptosis, former helping survival and metastasis formation, while latter delaying tumor growth.⁴ ARN could help to find the connection between sex steroid signaling and autophagy. In Figure 5, the transcriptional regulation of autophagy proteins by the androgen and estrogen receptors is presented. Regulation of ULK1/2 and UVRAG by ESR1 is experimentally verified,²⁹ according to the HTRI database. All the other connections were predicted in ARN using the JASPAR algorithm.¹⁵ Another autophagy protein, WIPI1 can also be transcriptionally regulated by sex steroid receptors. In addition, WIPI1 contains an LXXLL motif, which enables it to bind to ESR1, ESR2, and AR in a hormone independent way.³⁰ This connection is important, because localization of WIPI1 depends on autophagic activity, and at the same time it regulates sex steroid signaling, affecting the transcription of several autophagy proteins, including WIPI1 itself. As it is shown in Figure 5, 84% of the core autophagy proteins are transcriptionally regulated by sex steroid receptors. ESR1, ESR2, and AR regulate different but overlapping sets of autophagy proteins (23, 12, and 12 proteins, respectively). AR is also able to heterodimerize and activate ESR1, as well as ESR1 and ESR2 each other. Further research studies might reveal the role of these mechanisms in a gender-specific regulation of the autophagic activity.

Discussion

Here we present a novel resource on the regulation of autophagy in human. Autophagy Regulatory Network (ARN; <http://autophagy-regulation.org>) is a comprehensive interaction database featuring a manually curated core dataset, integrated and predicted data from numerous sources, and direct connection to literature curated interactions of 7 major signaling pathways. Directions, signs, confidence scores, and references are available for each interaction. ARN is accessible through a user-friendly webpage, and the data can be downloaded in all major bioinformatics standard formats, including simple text/table files and visualized Cytoscape networks.

To achieve a better understanding of context-dependent regulation of autophagic activity, a systems-level analysis of regulatory mechanisms is necessary. External stimuli processed by the signaling network can modulate autophagy at post-translational, transcriptional, and post-transcriptional level. Applying this approach in research studies can lead to the identification of key regulatory circuits, which are responsible for specificities in the regulation of autophagy, in different tissues, and under pathologic or therapeutic conditions. We created ARN with the aim to support the systems-level analysis of context-dependent regulation of autophagy, and also to facilitate the large-scale examination of a single autophagy-related protein.

Primary data on post-translational, transcriptional, and post-transcriptional regulation of autophagy proteins can be obtained from various resources. However, to use data from multiple resources in one analysis can be tedious because of the different data formats and molecular database IDs. Furthermore, many

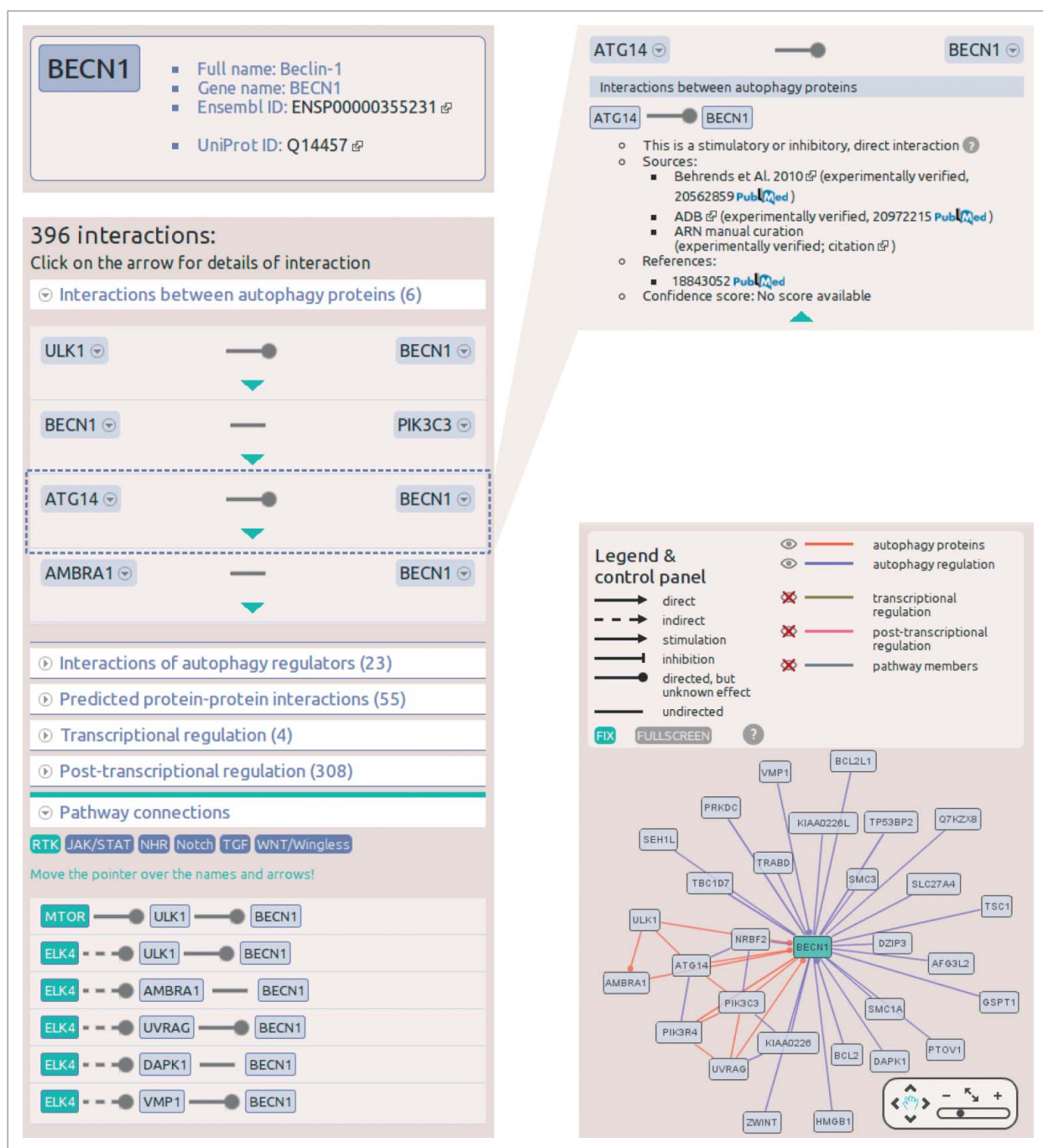


Figure 3. Screenshots from the protein datasheet of BECN1 from the ARN webpage. (A) At the top of the datasheet the name, gene name, UniProt ID, and Ensembl protein ID of the selected protein is shown, with hyperlinks to the UniProt and Ensembl webpages. Below this box, the potential signaling properties and disease related information with a special highlight on cancer types is listed. (B) The interactions of the selected protein are listed, grouped by layers. In addition, at the bottom of the list, the pathway connections can be browsed by pathways. (C) Information on sources, references and confidence scores of each interaction can be obtained by clicking on the green triangles. (D) On the right side of the datasheet, an interactive network image of the first neighbors of the selected proteins is available. Note that unlike ULK1, ULK2 is not present in the BECN1 network as ARN contains only those interactions that were specifically identified between exact proteins, and no publications were curated that experimentally verified the likely connection between ULK2 and BECN1.

resources often contain erroneous interactions between proteins, derived from high-throughput methods or predictions. To address this problem, ARN involves manually curated interactions between autophagy proteins, their post-translational

regulators, and between signaling components. Most of the interactions in ARN have confidence values allowing the user to set an own cut-off value (or use the default value calculated by ROC analysis, using manually curated interactions as gold standard

set). For all protein-protein interactions we offer the Gene Ontology semantic similarity score.³¹ This score is based on the assumption, that proteins involved in similar biological processes are more likely to interact *in vivo*. This way we can decrease the ratio of erroneous interactions from high-throughput screenings or predictions.

Before ARN, 2 autophagy-focused resources have been published. The Human Autophagy Database¹⁹ (HADb) contains only sequence data of genes from an autophagy-dedicated microarray. Autophagy Database²⁰ (ADB) provides orthology data from 41 species, and for some proteins also a list of interactions. However, the source of these interactions and the scope of the curation are not clearly defined. Indeed, the main aim of ADB is to serve a comprehensive collection of orthologs of autophagy-related genes. The interaction data are not available for download in a single file, but can be browsed only on the webpage. Compared to HADb and ADB, in ARN the data sources are well defined, and the size of the network is determined by the principles of its design. ARN provides data not only on post-translational regulators, but also on transcriptional and post-transcriptional regulators. In addition, beside the direct regulators of the proteins involved in autophagy initiation and execution, ARN makes a connection between the cellular signaling network and the regulation of autophagy. The directions, signs, and confidence scores of the interactions are supplied in format ready for computational processing.

ARN serves as a good basis for various kinds of bioinformatics approaches, while it also effectively supports wet lab research work. Using the ARN website, researchers are able to search for potential interactors or regulators affecting their subject of interest. ARN database contains many potential regulators of the entire autophagic process and even for a single component that allows researchers to combine expression or mutation data sets and analyze autophagy in context-specific states. For example, ARN data can be used to point out important alterations in autophagy regulation upon a disease.¹⁷ Therefore, ARN can support experiment design and evaluation for both basic and translational research works.

Furthermore, network data of ARN can be analyzed with graph topological methods, modularization methods,

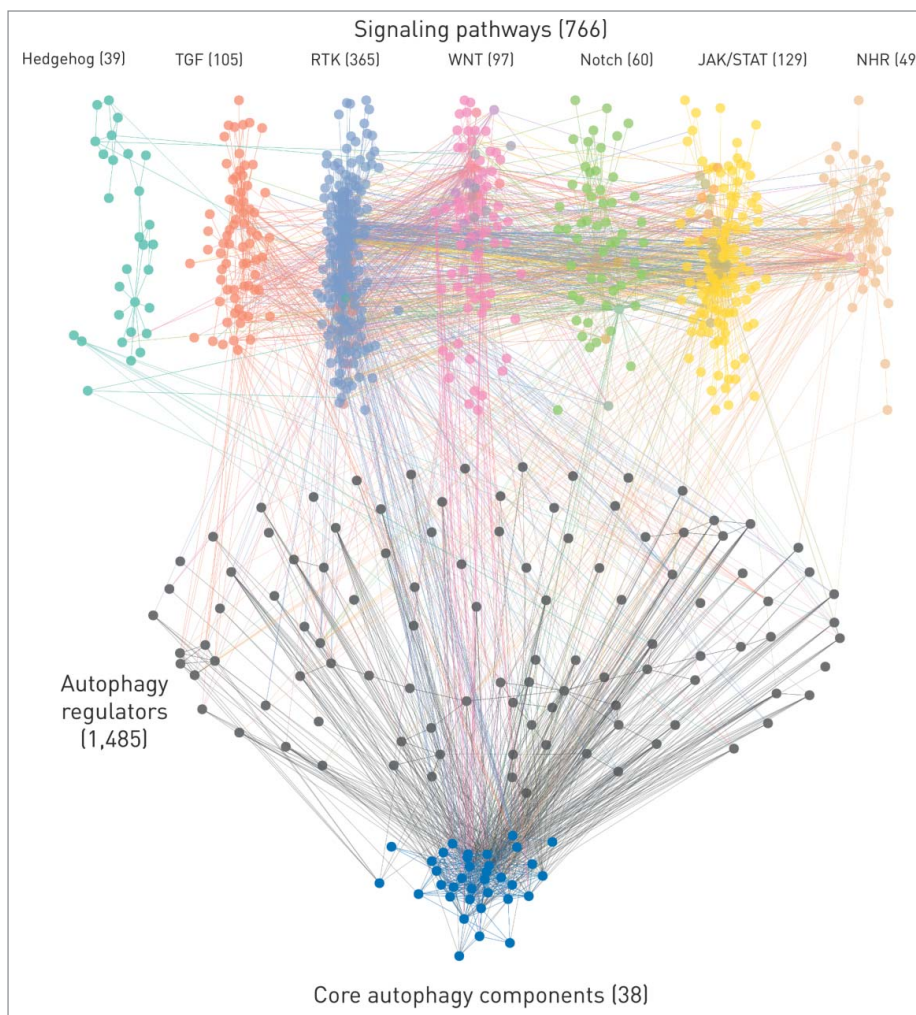


Figure 4. The network of 7 signaling pathways with direct autophagy regulators and core autophagy proteins. The numbers represent the total number of components in each section but for clarity, only components with the highest confidence, one- or 2- step long connections are shown on this figure. We also omitted the connections through transcription factors or miRNAs. Edges between autophagy proteins are blue. Intermediate components (i.e., direct autophagy regulators) in the 2-step connections and their edges are colored with black. Pathways are color-coded, multipathway proteins and edges between different pathways have the colors of the involved pathways mixed. Edges directly connecting pathways and autophagy proteins have the color of the source pathway.

perturbation simulations, and models can be built using different mathematical formalisms. Having an appropriate, good quality *a priori* knowledge as a starting point is a crucial requirement of successful modeling.³² ARN aims to support modeling approaches by serving as a good basis for a variety of methods, such as Boolean and rule-based modeling.^{33,34} Combining with gene expression or mutation data, comparative analyses can be carried out to investigate differences in autophagy regulation by tissue, physiological or pathological conditions, gender, and many other aspects. With the inclusion of drug compound and target interaction data, ARN is suitable to support network-based pharmacological attempts, such as multi-target and allo-network drug design.³⁵

Knowing that the list of components and interactions in each layer is not complete, we will include further experimentally

reaction or protein binding to them. For this purpose, we used the ELM server³⁸ and searched for enzymes (i.e., phosphatases, ubiquitin-ligases, peptidases, etc.) that can directly or indirectly modify autophagy components. We also used protein domain information from PFAM³⁹ to predict a protein-protein interaction (PPI) based on domain-domain interactions.

The next 3 layers contain information on the transcriptional and post-transcriptional regulators of the above described inner-layers (i.e., autophagy components, their known and predicted protein regulators). The transcriptional regulatory layer contains transcription factors that are known or predicted to transcriptionally regulate the inner layers. These regulatory connections were integrated from databases such as ABS,⁴⁰ ENCODE,⁴¹ HTRIdb,²⁹ ORegAnno⁴² and PAZAR,⁴³ or predicted with JASPAR.¹⁵ We also performed manual curation to collect TFs directly regulating autophagy proteins. In addition, to add the known complexity of transcriptional regulation, this layer also contains PPIs between the TFs from BioGRID,⁴⁴ InnateDB,⁴⁵ IntAct⁴⁶ and HPRD⁴⁷ databases. In the next layer, we integrated miRNAs as post-transcriptional regulators of the inner-layers (autophagy components and their direct regulators, including enzymes and TFs) from experimentally verified miRNA-mRNA interaction databases: miR2Disease,⁴⁸ miR-DeathDB,⁴⁹ miRecords,⁵⁰ miRTarBase,⁵¹ and Tarbase.⁵² The third regulatory layer contains the transcriptional regulators of these miRNAs (i.e., TFs known to regulate the expression of the miRNAs known to downregulate autophagy component or regulators). We used ENCODE,⁴¹ PuTmiR⁵³ 1.1 and 2.0 versions and TransmiR v1.2⁵⁴ to integrate this information. Data from the integrated resources were downloaded in the spring of 2013.

In the last step of the compilation, we connected signaling pathways from Signalink 2 (<http://signalink.org>),⁵⁵ a resource we recently developed, containing manually curated data of signaling pathways. Signalink 2 contains 7 major signaling pathways: RTK (receptor tyrosine kinase), TGF β /TGF- β (transforming growth factor β), WNT, Hedgehog, JAK-STAT, NOTCH and NHR. Connections between signaling pathways and autophagy were derived in 3 different ways: (a) predicted or experimentally verified direct PPIs between a signaling protein and an autophagy protein; (b) via the transcriptional regulation of a signaling pathway related TFs and its autophagy-related target; and (c) through post-transcriptional regulation, where a signaling pathway affects a TF of a miRNA, which regulates a protein involved in autophagy or its regulation. Note that we also added further protein-protein interactions from BioGRID,⁴⁴ InnateDB,⁴⁵ IntAct,⁴⁶ HPRD,⁴⁷ and predictions between all the already included protein components.

For every integrated data source containing interactions collected with different methods, quality control is highly important. From each source databases we included the available confidence scores, maintaining the possibility for the users to exclude low confidence interactions from their analysis. However, these scores are only available for the subset of interactions derived from the specific source. To obtain a general confidence

score for all protein-protein interactions, we calculated semantic similarity score³¹ between the Gene Ontology Biological Process⁵⁶ properties of the interacting pairs of proteins. In case of PPIs inferred from domain-domain based prediction, we performed a ROC analysis to minimize the false positive rate. With the domain-motif based prediction, we used the cut-off value suggested by the authors of the ELM Structure Filter³⁸ algorithm.

For each protein in ARN we included disease and cancer type annotations. We obtained diseases from GAD (The Genetic Associations Database),⁵⁷ and OMIM (Online Mendelian Inheritance in Man),⁵⁸ and cancer-type mutation patterns from COSMIC (Catalog of Somatic Mutations in Cancer).⁵⁹

Database implementation and structure

Data storage is based on MySQL, which serves data to the webpage by a PHP interface. The webpage uses jQuery on the client side to offer a high interactivity. Information can be loaded asynchronously by small http requests, giving an efficient and comfortable browsing experience through hundreds of interactions. We wrote a separate data export module in Python language that offers various choices to download data in CSV, BioPAX, PSI-MI TAB, PSI-MI XML, SBML, and Cytoscape's CYS format. Several options are available to customize the network to download: users are able to filter by interaction types (e.g., PPIs, transcriptional regulation), as well as by sources. There is also an option to separate experimentally verified and predicted interactions. The customized network files are generated according to the selected options by the export module running in the background. This process can take few minutes. Then, for each download, we generate a URL, where users can access the data for 14 days. Optionally, users can provide their e-mail addresses to which files smaller than 10 MB will be emailed. The whole dataset is also available as a standard SQL dump, so any complex query or modification can be applied using SQL statements.

The core of the ARN database is the interaction table. In the interaction table source and target fields are integers pointing to the primary keys of protein or "mirna" tables. The layer field denotes the type of the interaction, and its value determines if the source or the target refers to a protein or miRNA. The meanings of the values in the layer field are the followings: 0: interactions between autophagy executor proteins; 1: PPIs between autophagy proteins and their direct regulators from our manual curation, ADB²⁰ and the ChIP-Seq study of Behrends et al.;²¹ 2: direct and indirect regulators of autophagy proteins from general PPI resources⁴⁴⁻⁴⁷ and from predictions based on domain-domain³⁹ and domain-motif³⁸ interactions; 3: value not used due to technical reasons; 4: TF-target connections; 5: miRNA-mRNA connections; 6: PPIs in the signaling pathways, imported from Signalink 2;⁵⁵ 7: TF-miRNA connections; 8: PPIs between TFs, signaling pathways and autophagy regulators, from the same sources as layer 2. Each interaction has 3 main attributes: is_directed (0: undirected; 1: directed; 2: direction is predicted), is_direct (0: indirect; 1: direct) and is_stimulation (0: unknown; 1: stimulation, -1: inhibition). In addition, interactions have one or

more sources. Sources are listed in the source table, and the interaction_source table contains their assignment to the interaction table. Manually curated interactions have literature references, contained by the interaction_reference table. In the interaction_reference table, articles are identified by their Pubmed IDs. Most of the interactions have confidence scores. These are stored as float values in the interaction_weight table, the different types of scores are listed in weight table. Components of ARN are listed in the protein and "mirna" tables. The protein table contains the uniprot_name field, which is unique, and it contains the UniProt accession number of proteins. All records imported from other databases, as well protein names from articles are mapped to their primary UniProtKB ID. Proteins may have signaling topological properties and pathway assignments, available in protein_topology and protein_pathway tables. In the "mirna" table we used miRBase AC and miRNA name to identify miRNAs.

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Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

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