OGEE v3: Online GEne Essentiality database with increased coverage of organisms and human cell lines

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ABSTRACT

OGEE is an Online GEne Essentiality database. Gene essentiality is not a static and binary property, rather a context-dependent and evolvable property in all forms of life. In OGEE we collect not only experimentally tested essential and non-essential genes, but also associated gene properties that contributes to gene essentiality. We tagged conditionally essential genes that show variable essentiality statuses across datasets to highlight complex interplays between gene functions and environmental/experimental perturbations. OGEE v3 contains gene essentiality datasets for 91 species; almost doubled from 48 species in previous version. To accommodate recent advances on human cancer essential genes (as known as tumor dependency genes) that could serve as targets for cancer treatment and/or drug development, we expanded the collection of human essential genes from 16 cell lines in previous to 581. These human cancer cell lines were tested with high-throughput experiments such as CRISPR-Cas9 and RNAi; in total, 150 of which were tested by both techniques. We also included factors known to contribute to gene essentiality for these cell lines, such as genomic mutation, methylation and gene expression, along with extensive graphical visualizations for ease of understanding of these factors. OGEE v3 can be accessible freely at https://v3.ogee.info.

INTRODUCTION

Gene essentiality is a core concept of genetics, and has important relevance in relation to fundamental areas such as evolutionary, and systems and synthetic biology, and applicability in drug development. Essential genes (EGs) are often responsible for important biological processes and cellular fitness in an organism, and are critical for its survival. EGs are of particular importance in mechanistic biological studies and therapeutic applications, such as studying behavior of a biological system under perturbation (1), studying sequential gene deletions to define a minimal genome/organism (2,3) and identifying or prioritizing of therapeutic targets in pathogens (4–6) and human cancers (7–11). EGs complements have been identified in well-characterized small model organisms such as Saccharomyces cerevisiae and Caenorhabditis elegans (12,13) and

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some large and complex organisms, such as human (14), using various technologies.

The essentiality of a gene is highly dependent on various factors including the genetic context, genetic background of the host and environment (15). Hence, gene essentiality is not a static, but rather a context-dependent property of a gene. Since version 1, OGEE has been promoting the context-dependent concept and its importance for understanding on gene essentiality (16). In our last update (17), we focused on the importance of ‘conditionally essential genes’ (CEGs) or ‘differentially essential genes’ (DEGs) in cancer cell lines. The current update provides: (a) an increased coverage of essentiality tested genes and species, (b) up-to-date essentiality status of existing genes, (c) an increased coverage of human cell lines, (d) an extensive graphical visualization for various factors that influence gene essentiality and (e) a dedicated domain with secure HTTPS protocol, https://v3.ogee.info.

DATA GENERATION

Collection and organization of essentiality datasets
To find large-scale gene essentiality experiments, we examined publications from the NCBI PubMed database (18) identified using the key words ‘essential genes’, ‘gene essentiality’, ‘tumor dependency genes’, ‘cancer vulnerable genes’, and then downloaded all classified essential or non-essential genes. For model organisms such as Drosophila melanogaster, Caenorhabditis elegans, Saccharomyces cerevisiae and Arabidopsis thaliana, we also obtained their tested genes from online public databases (19–30). The full list of reference databases and publication records are provided in Supplementary Table S1. The tested results (i.e. essential and non-essential statuses) were organized into datasets, according to their experimental conditions. In total, we collected 127 datasets for 91 non-human species, containing 213,608 tested genes. This collection represents a significant increase from our previous version, OGEE v2 (17), which contained 99 datasets from 48 non-human species, with in total 167,799 tested genes.

Essentiality datasets from human cancer cell lines
Tumor essential genes, also known as tumor dependency genes, are potential targets for treatment and/or drug development (31,32). During the last decade, cancer cell lines have been used extensively for tumor essential gene identification (32,33). We included results for 16 human cell lines in our last version. In OGEE v3, we expanded this collection to include a total of 581 cell lines which were tested by genome-wide CRISPR-Cas9 and RNAi screening. The 581 cell lines were related to 25 different tissues based on their histological origins, as shown in Figure 1A.

In total, 150 cell lines were tested by both RNAi- and CRISPR-Cas9-based methods (Supplementary Table S2). As shown in Figure 1B, the CRISPR-Cas9-based method identified more essential genes than RNAi in most cell lines, likely because of higher efficiency of CRISPR-Cas9 in gene knockdown/out. Essential genes identified by RNAi are often not a subset of those identified by CRISPR–Cas9 in the same cell line, thus the combined data from two distinct methods could offer more robust results and coverage over human essential genes, as a previous study suggested (34).

In summary, the current version (v3) of OGEE include 16 datasets from 16 non-human eukaryotes and 111 datasets from 75 prokaryotes. Of the 213,608 collected genes, 31,177 were tested in multiple datasets, accounting up to 14.6% of all collected genes. Of the tested genes from multiple datasets, 15,440 genes were conditionally essential genes (CEGs) or context-dependent, representing around 49.52% of genes covered by multiple datasets. In addition, OGEE v3 also includes experimental essentiality results for 581 human cancer cell lines.

Collection of known factors influencing gene essentiality
The essentiality of a gene at any given time point depends on various factors. In OGEE v3, we included several factors known to influence gene essentiality, including paralogs and orthologs or duplication status (35), functional annotation of a gene (36), connectivity of protein-protein interaction (PPI) network (37) and early expression during embryonic development. OrthoFinder (38) was used to identify paralogs and orthologs for all species included here. The PPI data were downloaded from latest version of STRING v11.0 database (39); interactions with score ≥900 were used to compute the connectivity score for all the genes. As described in previous studies (40,41), the EGs had more interaction partners at the protein level. Protein sequences were obtained from NCBI protein database (18), while their functional annotations were obtained from the Gene Ontology (GO) database (42). For more information, please refer to the ‘Help’ page of the OGEE database.

For human cell lines, factors known to influence gene essentiality included mutation, methylation and gene expression (43); in fact, factors including these have been used to predict gene essentiality in human (43,44). Thus, for each cell line, we collected data on gene mutations from cancer (45), on methylation data from NCBI GEO database (18) and/or on gene expression data from the CCLE database (46).

Built-in tools for analysis
OGEE v3 also has integrated tools to analyze the impact of gene properties on the page ‘Analyze’. The available options include developmental vs. non-developmental and duplicates vs. singlets, calculated in terms of proportion of essentiality ($P_E$). For more detail on built-in tools, please refer to our previous publication (17).

EXTENSIVE ANNOTATIONS OF HUMAN ESSENTIAL GENES
The identification and characterization of human EGs provides a better understanding of human diversity, has practical medical applications for disease genetics and clinical interpretation of disease-associated genetic variants; EGs are associated with the core developmental, metabolic and
AB Figure 1. Statistics on 581 human cell lines collected in OGEE v3. (A) The 581 cell lines were related to 25 different tissues by their histological origins. (B) CRISPR-based method often identified more essential genes than RNAi-based method. Each point indicates a cell line, and the x-axis and y-axis mean the number of identified essential genes of the cell line in CRISPR–Cas9 and RNAi datasets respectively. The colors (blue, green, red, yellow) illustrate the number of overlapping essential genes between CRISPR–Cas9 and RNAi data. The red line denotes the fitted regression line between numbers of essential genes in RNAi and CRISPR-Cas9 data. In total 150 cell lines were tested by both techniques.

signaling pathways; and they are less likely to be tolerant missense variation, and more prone to pathogenicity (41). While effort has been put into identifying targets for cancer therapeutics, several research groups (32,33) have identified cancer dependencies genes and lineage-specific EGs in human cell lines using RNAi screening or CRISPR–Cas9 technology. Since both RNAi and CRISPR are promising complementary methods for identifying EGs, we annotated human EGs extensively by considering both of these techniques as well as factors influencing gene essentiality.

As shown in Figure 1A, the 581 human cell lines collected in OGEE v3 were assigned into 25 groups according to their tissue of origin. The largest number of cell lines were for lung (n = 123), followed by central nervous system (n = 59), haematopoietic and lymphoid tissue (n = 53), ovary (n = 40) and breast (n = 38). We selected the RKO cell line set to exemplify the annotation of essential gene in OGEE v3 (https://v3.ogee.info/#/cellline/large%20intestine/RKO/summary). The cell line RKO is inferred to have 672 essential genes based on RNAi screening, and 1251 essential genes based on CRISPR–Cas9 dataset (Figure 2A). To understand more about the characteristics of essential genes and the relationship with the influencing factors in human cell lines, we compared tested genes of RKO against the known influencing factors data that we had collected (Figure 2C–E). Mutations, methylation and gene expression all contributed significantly to gene essentiality in this cell line. For example, genes harboring pathogenic mutations are more likely to be essential regardless of the experimental methods used (Figure 2C); genes with less methylation at their transcription start site (TSS200) are also more likely to be essential (Figure 2D); in general, lowly-expressed genes are less likely to be essential (Figure 2E).

To better appraise the pattern of essentiality across human cell lines and identify potentially meaningful biological relationships between genes (47), we selected data of those tissues with at least ten cell lines from a conditioned experiment, and assessed gene-gene essentiality relationships among individual tissues by Spearman and Pearson correlation. Raw fitness scores instead of binary essentiality assignments (i.e. essential or non-essential genes) were used for such calculations. After filtering the genes with essentiality in at least 50% cell lines of a specific tissue, we then selected gene pairs with significant correlation results (P-value < 0.001). The results of this analysis are available via ‘Correlation co-essentiality’ in OGEE v3 for each human gene tested that met the cut-off (see Figure 2B for an example).

Orthology and comparisons with mouse essential genes
To date, mouse is the only mammalian species for which gene essentiality has been tested at organismal level. About 30% of tested mouse genes are essential for survival, which is markedly higher than that of human cell lines (21). Thus, unique essential genes in mouse experiments may be important during growth and development, while those uniquely essential in human cell lines may represent suitable treatment and/or drug targets with putatively less side-effects. To compare gene essentiality between mouse and human cells, we first selected tissues for which at least six cell lines were available, and defined 1607 core essential human genes in >50% of the cell lines (n ≥ 3). We downloaded the mouse knock-out and associated phenotype data from databases IMPC (21) and MGI (22) to deeply understand the functions of essential genes in mice; these databases contained information for 5799 and 9693 knockout genes, respectively.
Figure 2. Extensive annotations on human essential genes provided by OGEE v3. Shown here are graphical visualizations taken from the OGEE v3 (https://v3.ogee.info/#/cellline/large%20intestine/RKO/summary). (A) gene essentiality determined using different experimental methods. (B) Correlational co-essentiality chart of AAMP gene from central nervous system tissue tested by CRISPR-Cas9 technique. The orange edges denote negative correlation while the blue edges denote positive correlation among the connected genes. (C) Gene essentiality as function of mutation: mutation outcomes predicted using FATHMM were used to group all tested genes into three categories, the percentage of essential in each group was then calculated (please consult ‘Help’ page for more details); left panel: gene essentiality tested using RNAi, right panel: gene essentiality tested using CRISPR–Cas9. (D) Gene essentiality as function of methylation: genes are grouped based on methylation sites and calculate percentage of essential for each sites; red color gradient is used to denote the methylation score, increasing gradually from 0 to 1; users can select any methylations from the drop down menu. (E) Gene essentiality as function of gene expression: all 0 rpkm genes are in the first bin and rest of the genes are assigned into equal size nine bins; lastly, percentage of essential is calculated for each bin.
37.6% (4463 by OrthoFinder (38) (see ‘Data generation’), we identified the mouse genome. From the orthogroup data generated models, and building a complete functional catalogue for phenotypes of mutants and controls, identifying the disease Both MGI and IMPC are generating a knockout mouse line for each protein-coding gene, collecting the corresponding phenotypes of mutants and controls, identifying the disease models, and building a complete functional catalogue for the mouse genome. From the orthogroup data generated by OrthoFinder (38) (see ‘Data generation’), we identified 3522 and 6214 knock-out genes for mice in the IMPC (21) and MGI (22) dataset, respectively. We also identified that 37.6% (4463/11 868) mouse genes were identified as lethal or sub-viable, and there existed 4192 genes corresponding to human orthologs, 13.5% (565/4192) of which are consistently essential in human cells (84.7%, 565/667) (shown in Figure 3). These results showed that the mice exhibit similar survival behavior with human cell line screened in their corresponding orthologous genes.

**CONCLUSION**

Here, we introduce OGEE v3, an updated version of the Online GEne Essentiality database. This update almost doubles the number of species for which EGs have been inferred, and a marked increase in the number of genes that have been tested for essentiality. Furthermore, we thoroughly revisited information on conditionally essential genes in both existing and new datasets. We also included 581 datasets for human cell lines and genes tested using CRISPR–Cas9 and/or RNAi methods. In addition, we also included information on factors that influence gene essentiality to assist biological interpretation. Thus, OGEE v3 is expected to be a useful and an important database for biologists, bioinformatician and other colleagues from a range of scientific communities who are working on, or are interested in, exploring gene essentiality.

**OUTLOOK**

Over the coming years, the OGEE team aims to continue tracking all the available genes data sets and gene properties with the latest experimental records. Specific foci of the OGEE team will be to include a BLAST function (48) in the database and to improve the user-interface, visualizations and crosslinking of resources.

**DATA AVAILABILITY**

All data are freely accessible to all academic users. This work is licensed under a Creative Commons Attribution 3.0 Unported License (CC BY 3.0). Users can download datasets from the ‘Download’ page. Individual datasets or combined datasets of individual species can be downloaded via the ‘Browse’ page.

**SUPPLEMENTARY DATA**

Supplementary Data are available at NAR Online.

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