LncDisAP: a computation model for LncRNA-disease association prediction based on multiple biological datasets

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Abstract

Background: Over the past decades, a large number of long non-coding RNAs (lncRNAs) have been identified. Growing evidence has indicated that the mutation and dysregulation of lncRNAs play a critical role in the development of many complex human diseases. Consequently, identifying potential disease-related lncRNAs is an effective means to improve the quality of disease diagnostics and treatment, which is the motivation of this work. Here, we propose a computational model (LncDisAP) for potential disease-related lncRNA identification based on multiple biological datasets. First, the associations between lncRNA and different data sources are collected from different databases. With these data sources as dimensions, we calculate the functional associations between lncRNAs by the recommendation strategy of collaborative filtering. Subsequently, a disease-associated lncRNA functional network is built with functional similarities between lncRNAs as the weight. Ultimately, potential disease-related lncRNAs can be identified based on ranked scores derived by random walking with restart (RWR). Then, training sets and testing sets are extracted from two different versions of a disease-lncRNA dataset to assess the performance of LncDisAP on 54 diseases.

Results: A lncRNA functional network is built based on the proposed computational model, and it contains 66,060 associations among 364 lncRNAs associated with 182 diseases in total. We extract 218 known disease-lncRNA pairs associated with 54 diseases to assess the network. As a result, the average AUC (area under the receiver operating characteristic curve) of LncDisAP is 78.08%.

Conclusion: In this article, a computational model integrating multiple lncRNA-related biological datasets is proposed for identifying potential disease-related lncRNAs. The result shows that LncDisAP is successful in predicting novel disease-related lncRNA signatures. In addition, with several common cancers taken as case studies, we found some unknown lncRNAs that could be associated with these diseases through our network. These results suggest that this method can be helpful in improving the quality for disease diagnostics and treatment.

Keywords: Long non-coding RNAs, Disease, lncRNA network, Random walking with restart
Long non-coding RNAs (lncRNAs), which compose the largest portion of the mammalian non-coding transcriptome [1], are emerging as important regulators of tissue physiology and disease processes [2]. lncRNAs are expressed in a more tissue-specific fashion than mRNA genes [3] and are highly specific to cell type, organs, and species [4]. A large amount of lncRNAs have been demonstrated to have a close relationship with many complex human diseases [5–8]. Therefore, an increasing recognition of the roles of lncRNAs in human disease has created new diagnostic and therapeutic opportunities [9]. The identification of potential lncRNAs related to complex diseases is a hot topic in medicine.

LncRNAs are the key to explaining disease mechanisms. As analysing lncRNAs is very appealing to researchers, many researchers have devoted their work to lncRNAs for exploring complex human diseases at the molecular level. For example, BCYRN1 has been demonstrated to induce the proliferation and migration of non-small cell lung cancer (NSCLC) cells and play an important role in NSCLC progression [10]. LncRNA SNHG1 regulates NOB1 expression by sponging miR-326 and promotes tumourigenesis in osteosarcoma [11]. Ye et al. found that LINC00460 promotes the progression of lung adenocarcinoma by competitively binding miR-302c-5p and regulating the FOXA1 signalling pathway [12]. F. Aksoy et al. postulated that the overexpression of lncRNA DANC1 may be associated with poor outcomes in upper rectal cancer [13]. LncRNA HOTAIR plays a role as an oncogenic molecule in different cancers, including breast, gastric, colorectal and cervical cancer cells [14]. Similarly, lncRNA MALAT1 is considered a potential biomarker for the diagnosis and prediction of cancers and may also serve as a therapeutic target for the treatment of specific tumours [15]. In 2018, Chen C et al. deduced that the expression of lncRNA ZEB1-AS1 might be used as a promising prognostic biomarker for cancer [16]. The above studies show that lncRNAs have been recently regarded as possible biomarkers for disease.

Although a large number of lncRNAs have been recorded in public databases, such as GENCODE [17], NONCODE [18], LNCipedia [19], only a few lncRNAs have been characterized functionally [20]. Several methods have been developed to predict potential lncRNA-disease associations [21, 22]. However, they take into account only disease semantic similarity and ignore disease functional similarity. Improved knowledge has suggested that exploring both the semantic and functional associations of diseases, which are two types of significant associations, are beneficial in measuring disease similarity because not all associations between diseases are represented by the disease ontology, and many of them are reflected through the functional associations among disease-related genes [23]. Moreover, the lack of unified identifications for lncRNAs leads to an underutilization of information from different public lncRNA databases when lncRNA functional annotations are approached. Therefore, we aimed to identify more lncRNAs by efficiently analysing the lncRNA and disease data. First, we extracted and utilized functional information related to lncRNAs, including disease similarity, protein-protein interactions and lncRNA-mRNA associations. Subsequently, we established functional associations between lncRNAs and built a disease-related lncRNA network. Potential disease-related lncRNA signatures were predicted by a random walking with restart (RWR).

**Materials and methods**

**Workflow**

The workflow of LncDisAP is shown in Fig. 1. First, mappings between lncRNAs and lncRNA-related datasets are established, and these datasets are extracted from multiple biological datasets. Mappings between lncRNA and protein are provided by the Search Tool for the Retrieval of Interacting Genes/Proteins (STRING) [24] and starBase v2.0 [25] databases, while those between lncRNA and disease are from the Human Disease Ontology (DO) [26], MEDIC [27] and LncRNADisease [28] databases. The mappings of lncRNA-mRNA are from starBase v2.0 [25] and the Human Protein Reference Database (HPRD) [29]. Subsequently, different similarity measures can be defined considering that different data sources have different data characteristics. Given the associations between lncRNA and mRNA, the number of lncRNA-related mRNAs can be taken as a statistical indicator to calculate lncRNA similarity. In view of disease functional similarity and protein functional similarity, associations of lncRNA-disease and lncRNA-protein are used to make a multi-dimensional vector model for each lncRNA. Finally, a disease-related lncRNA functional network is built based on lncRNA functional similarity. We employ RWR in this network to calculate the ranking of candidate lncRNAs, which are related to certain diseases. Thus, the potential relationships between diseases and lncRNAs can be identified.

**Data source**

**Disease database**

DO [26] database is focused on representing a common and rare disease concept, which aims to provide an open source ontology for the integration of biomedical data associated with human disease. Each node in DO represents one disease term. All of these nodes are organized in a directed acyclic graph (DAG) with an ‘IS_A’ relationship. MEDIC [27], as a part of the Comparative Toxicogenomics Database (CTD) [30], integrates Online
Mendelian Inheritance in Man (OMIM) terms, synonyms and identifiers with MeSH [31] terms, synonyms, definitions, identifiers and hierarchical relationships. It is composed of 9700 unique diseases described by more than 67,000 terms. In this study, we map lncRNA-related diseases to DO, utilizing terms and synonyms from DO and MEDIC.

LncRNA database
RNAcentral [32] is a database of non-coding RNA (ncRNA) sequences that aggregates data from specialized ncRNA resources. It assigns unique identifiers to every distinct RNA sequence. Because there is no uniform identity number in the different lncRNA databases, we use identifiers from RNAcentral as unified labels of lncRNAs to ensure the smooth progress of this work.

Human lncRNA-disease association data
LncRNADisease [28] is a database that curated the experimentally supported lncRNA-disease association data. Presently, there are three versions available. The 2017 version of the LncRNADisease database integrated 2947 lncRNA-disease entries, including 888 lncRNAs and 328 diseases, while the 2015 version covered 1102 lncRNA-disease entries, including 373 lncRNAs and 252 diseases. The newest version [33] was released in 2018, containing 5714 lncRNAs and 423 diseases. Here, we extract associations between lncRNAs and diseases from this database and use the differences between its versions to validate the reliability of LncDisAP.

Human protein-protein interaction data
STRING [24] is a database of known and predicted protein-protein interactions. These interactions in STRING include direct (physical) interactions, as well as indirect (functional) interactions, which stem from computational prediction, knowledge transfer between organisms, and interactions aggregated from other databases. The STRING database currently covers 9,643,763 proteins from 2031 organisms. Here, protein-protein interactions from STRING are involved in the lncRNA similarity computation.

Human lncRNA interaction data
starBase v2.0 [25] systematically identified the RNA-RNA and protein-RNA interaction networks from 108 CLIP-Seq data sets generated by 37 independent studies, which provided 423,966 miRNA-mRNA, 10,212 miRNA-lncRNA and 17,609 protein-lncRNA experimentally confirmed interactions based on large scale CLIP-Seq data. The HPRD [29] represents an mRNA-mRNA interaction network for humans. All the information in HPRD has been manually extracted from the literature by expert biologists. Currently, HPRD covers 39,240 mRNA-mRNA interactions with 9465 mRNA.

LncRNA functional similarity calculation
Data pre-processing
The differences in different data sets bring some difficulties to the integration of lncRNA data. Two problems must be solved before constructing the lncRNA functional association network. One is the mapping of disease terms. MEDIC and DO are both comprehensive
disease corpuses and contain abundant disease terms, so we can annotate DO entries with the vocabulary from MEDIC and create a combined vocabulary of disease terms. Referring to this vocabulary, we build mappings between the DO terms and the disease terms of LncRNA-Disease. The other problem that must be addressed is the unification of lncRNA identifications. As mentioned above, the lncRNA naming rules of different LncRNA databases are different. Therefore, we employ the RNAcentral id as the unified identification system of LncRNAs considering that the RNAcentral database provides mapping data among various public LncRNA databases.

LncRNA-related disease similarity
The functional similarities between different diseases can be calculated. Therefore, in view of associations between LncRNAs and diseases, we can make a multi-dimensional vector model for each lncRNA with diseases as dimensions. The functional similarities between these LncRNA-related diseases can be taken as inputs to further calculate relevance scores for LncRNAs. Here, we employ FNSemSim [34] to calculate disease functional similarity. This method, which we previously developed, has good performance for calculating similarities between diseases. In this method, we first calculate disease functional similarity utilizing associations between diseases and genes. The functional similarity between disease $d_a$ and $d_b$ is defined as follows:

$$FNSim(G_a, G_b) = \frac{\sum_{g \in G_a} |R_G(g)| + \sum_{g \in G_b} |R_G(g)|}{\text{num}(G_a) + \text{num}(G_b)}$$ (1)

where the gene sets $G_a = \{g_{a1}, g_{a2}, \ldots\}$ and $G_b = \{g_{b1}, g_{b2}, \ldots\}$ are related to disease $d_a$ and $d_b$, respectively; $\text{num}(G)$ represents the numbers of genes related to one disease; and $R_G(g)$ represents the degree of association between a gene $g$ and a gene set $G$ (see details in [34]). Considering that semantic associations exist in DO, FNSemSim could be defined as follows:

$$FNSemSim(d_a, d_b) = FNSim(G_a, G_b) \cdot \frac{|G_a||G_b|}{GMICA}$$ (2)

where $|G|$ represents the size of a gene set $G$. $GMICA$ represents the genes related to the most informative common ancestor of disease $d_a$ and $d_b$. Finally, by min-max normalization, we normalize similarities between pair-wised diseases.

Vector model construction for lncRNAs
STRING provides human protein-protein interactions, and in the above section, the functional similarities between lncRNA-related diseases have been calculated. Therefore, we can obtain the relational degrees between one lncRNA and a certain disease or protein based on the similarities of lncRNA-related diseases or proteins. Then, these degrees can be used to make a multi-dimensional vector for each lncRNA. Hence, we can calculate lncRNA functional similarity by cosine similarity in a multi-dimensional space, which is defined by lncRNA-related diseases and proteins. The workflow of calculating lncRNA functional similarity based on the recommendation strategy of collaborative filtering is shown in Fig. 2.

In this multi-dimensional space, neither all diseases nor all proteins are directly related to one lncRNA. To predict the score of a disease that is not directly related to one lncRNA, we define $L$ as the set of lncRNAs, $D$ as the set of lncRNA-related diseases and $P$ as the set of lncRNA-related proteins. $DR_l$ is defined as the set of diseases directly related to lncRNA $l$. The predicted association score between disease $d$ and lncRNA $l$ is defined as follows:

$$AS(d, l) = \begin{cases} \text{MAX}(FNSemSim(d_i, d)) & d \in DR_l \text{ and } d \notin DR_l \\ 1 & d \in DR_l \end{cases}$$ (3)

where $l \in L$, $d \in D$, $DR_l \subseteq D$ and $1 \leq i \leq |DR_l|$; here, $|DR_l|$ represents the number of diseases in the set of $DR_l$. Similarly, for lncRNA-related proteins, $PR_l$ is defined as the set of proteins directly related to lncRNA $l$. The predicted association score between protein $p$ and lncRNA $l$ is defined as follows:

$$AS(p, l) = \begin{cases} \text{MAX}(SPscore(p_i, p)) & p \in PR_l \text{ and } p \notin PR_l \\ 1 & p \in PR_l \end{cases}$$ (4)

where $l \in L$, $p \in P$, $PR_l \subseteq P$ and $1 \leq i \leq |PR_l|$; here, $|PR_l|$ represents the number of proteins in the set of $PR_l$ and $SPscore(p_i, p)$ represents the relevance score between protein $p$ and $p_i$ from STRING.

Subsequently, we define a vector of each lncRNA with $|D| + |P|$ dimensions. $|D|$ and $|P|$ represent the size of the disease set $D$ and the protein set $P$, respectively. For each lncRNA, we can define its vector $\overrightarrow{I}$ as follows:

$$\overrightarrow{I} = (AS(d_1, l), \ldots, AS(d_k, l), AS(p_1, l), \ldots, AS(p_j, l))$$ (5)

where $\overrightarrow{I}$ represents the vector of lncRNA $l$ in this multi-dimensional space. $AS(d, l)$ and $AS(p, l)$ are the
scores of disease $d_k$ and protein $p_j$, respectively, for lncRNA $l$. Now, we can obtain $|L|$ vectors of lncRNAs.

**LncRNA functional similarity**

In this multi-dimensional space, each lncRNA can be depicted by a multi-dimensional vector. Therefore, we can measure the similarity between any two vectors of lncRNAs based on cosine similarity. The similarity between lncRNA $l_1$ and lncRNA $l_2$ is defined as follows:

$$CR(l_1, l_2) = \frac{\sum_{i=1}^{n} (AS_{1,i} \times AS_{2,i})}{\sqrt{\sum_{i=1}^{n} AS_{1,i}^2} \times \sqrt{\sum_{i=1}^{n} AS_{2,i}^2}}$$ \quad (6)

where $AS_{k,i}$ represents the association score in the $i$-th dimension of the vector $T_k$ for lncRNA $l_k$. The range of $CR(l_1, l_2)$ is $0$ to $1$ because these values of $AS_{k,i}$ are positive numbers.

In addition, mRNA can also be seen as a factor to calculate lncRNA functional similarity because of the existing links between lncRNAs and mRNAs. In view of the relationships between mRNAs, we can extract links between them from HPRD denoted as mRNALinkSet. First, the relevance between an mRNA $k$ and an mRNA set $M$ is defined as follows:

$$R(k, M) = \frac{1}{\text{links}(k, M)/|M|}$$ \quad (7)

where $\text{links}(k, M)$ represents the number of links between mRNA $k$ and members in the mRNA set $M$, and these links have to be included in mRNALinkSet. Let a pair of mRNA sets $M_1 = \{m_{11}, m_{12}, \ldots\}$ and $M_2 = \{m_{21}, m_{22}, \ldots\}$ be related to lncRNA $l_1$ and $l_2$, respectively. The similarity between lncRNA $l_1$ and $l_2$ based on mRNA is defined as follows:

$$MR(l_1, l_2) = \frac{\sum_{1 \leq i \leq |M_1|} R(m_{2i}, M_1) + \sum_{1 \leq i \leq |M_2|} R(m_{1j}, M_2)}{|M_1| + |M_2|}$$ \quad (8)

where $|M_1|$ and $|M_2|$ represent the numbers of mRNAs related to lncRNA $l_1$ and $l_2$, respectively. Finally, we
complete the calculation of lncRNA similarities based on different lncRNA-related knowledge.

Identifying novel candidate disease-related lncRNAs

We can take lncRNA similarities as weight to construct a lncRNA functional association network. In this network, the weight between lncRNA \( l_1 \) and \( l_2 \) is defined as follows:

\[
\text{LncFunNet}(l_1, l_2) = 1 - (1 - CR(l_1, l_2))(1 - MR(l_1, l_2))
\]

where the range of \( \text{LncFunNet}(l_1, l_2) \) is 0 to 1, as in \( CR(l_1, l_2) \) and \( MR(l_1, l_2) \). Utilizing this lncRNA network, we can identify novel candidate disease-related lncRNAs.

To identify novel candidate disease-related lncRNAs, we employ RWR to fully exploit the global functional associations between lncRNAs in this network. RWR, as a global optimization method, can reveal more information between one lncRNA and all the others in the network. The random walker in the network starts from the root node and moves to adjacent nodes with the probabilities from that node to the others. After enough iterations, the probabilities from the root node to all the other nodes will become stable, which can be used as scores for predicting novel disease-related lncRNAs (see [35] for RWR details). Finally, rankings for each lncRNA in this network can be listed by RWR.

Results

LncRNAs and diseases

We obtained 3,801,586 associations among 4703 disease terms from DO based on disease similarity calculations. Meanwhile, we found 1083 relationships between 184 diseases and 374 lncRNAs by mapping DO terms to the diseases in LncRNADisease (released in July 2017). There were 5,600,133 relationships between 13,716 mRNA and 1034 lncRNAs extracted from starBase v2.0. We found 15,622 associations between 33 proteins and 2750 lncRNAs from starBase v2.0 and STRING.

We calculated similarity among 374 lncRNAs and removed lncRNA pairs that had a similarity of 0. Finally, we built a lncRNA functional network, which contains 66,060 associations among 364 lncRNAs associated with 182 diseases.

Performance

To assess the performance of the lncRNA functional network, we compared two different versions of LncRNADisease and extracted 218 known disease-lncRNA pairs associated with 54 diseases from the newer version of LncRNADisease (released in June 2018). The detailed statistics for evaluating disease-related lncRNA networks are given in Additional file 1.
Fig. 3 The performance in predicting candidate IncRNAs associated with cholangiocarcinoma. a. ROC curves of these five IncRNAs based on the test set from the 2018 version, including AFAP1-AS1, SPRY4-IT1, PANDAR, CCAT1 and CCAT2. b. AUC of these five IncRNAs based on the test set from the 2018 version.

Fig. 4 Average AUC of 54 diseases based on the test set of the 2018 version. 22 diseases have average AUC values greater than 80%, while 27 diseases have average AUC values between 60 and 80%.
cholangiocarcinoma and cervical cancer as case studies. First, we built a lncRNA functional network based on the data source from LncRNADisease (released in 2017), and the unknown lncRNA-disease associations of each disease were ranked by RWR. We found that lncRNA H19 had a high score of 0.84 for acute myeloid leukaemia, which was ranked in the top 8% and not included in the latest version of LncRNADisease. Zhang et al. [37] and Zhao et al. [38] showed that lncRNA H19 is associated with acute myeloid leukaemia. For breast cancer, lncRNA Pvt1, which was ranked in top 5%, was validated to regulate triple-negative breast cancer through KLF5/beta-catenin signalling [39]. LncRNA AFAP1-AS1 and wrap53 were both ranked in top 5% for cholangiocarcinoma [40–42]. Furthermore, lncRNA XIST had a top ranking of 4% for cervical cancer, as shown in Fig. 5. Zhu et al. [43] explored the specific mechanism and biological function of lncRNA XIST in cervical cancer, and their experiments indicated that lncRNA XIST accelerates the progression of cervical cancer via upregulating Fus through competitively binding with miR-200a.

Discussion

The impact of data sources and test sets

The relationship between diseases and lncRNAs in the data source was extracted from LncRNADisease released in July 2017. Hence, we evaluated the impact of different data sources and different test sets on the performance of predicting disease-lncRNA pairs in the disease-related lncRNA functional network. First, with the LncRNADisease data set released in June 2018 as the test set, we compared the two lncRNA functional networks that were built based on data sources from the 2015 and 2017 versions. After the above-mentioned validation strategy was carried out, the lncRNA functional network based on the data source from LncRNADisease released in 2015 had an AUC value of 72.6%, while the AUC of the network based on the 2017 version reached 78.08%. Simultaneously, we assessed the performance of the lncRNA functional network based on the data source of the 2015 version with a test set extracted from the 2017 version, whose AUC reached 72.8%, as shown in Fig. 6. The test result of the lncRNA functional network based on the data source from LncRNADisease released in 2015 is given in Table 2. It can be seen that there is not much difference between the two test sets, which may be the reason why the AUCs of the network based on the two test sets do not have much difference. This suggests that the performance of predicting potential disease-lncRNA pairs in the disease-related lncRNA functional network can be noticeably impacted by different data sources and different test sets.

LncRNA expression similarity

The introduction of lncRNA expression similarity has been considered before. However, the results are not ideal. We obtained lncRNA expression profiles from NONCODE [18]. This database is an integrated knowledge database that provides expressed profiles from human lncRNAs. Spearman’s rank correlation coefficient is employed to calculate associations between lncRNA $l_1$ and $l_2$, denoted as $ER(l_1, l_2)$. The
similarity between lncRNA $l_1$ and $l_2$ is defined as follows:

$$LncFunNet(l_1, l_2) = 1 - (1 - CR(l_1, l_2))(1 - MR(l_1, l_2))(1 - ER(l_1, l_2))$$

(10)

Subsequently, we built two lncRNA functional networks based on data sources from the 2015 and 2017 versions with the lncRNA expression similarity operator introduced. LncRNADisease released in June 2018 was taken as the test set. The lncRNA functional network based on LncRNADisease released in 2015 had an AUC value of 68.3%, while the AUC of the network based on the 2017 version achieved 75.46%. As shown in Fig. 7, the original calculation model had a better performance than one with the lncRNA expression similarity operator introduced, regardless of whether the data source was extracted from the 2015 or 2017 version of LncRNADisease. We found that the number of mappings between RNAcentral and NONCODE was insufficient. This may have an impact on the performance because sufficient and reliable data can make a contribution to predicting potential disease-related lncRNAs, while a small amount of data may have a negative impact. Hence, this is the reason why the lncRNA expression similarity operator was not introduced.

Table 2 The test result based on different versions of the data source

| Disease Name               | Disease Ontology | 2017 Version | 2018 Version |
|----------------------------|------------------|--------------|--------------|
| lung benign neoplasm       | DOID:3683        | 0.6744       | 0.7479       |
| stomach cancer             | DOID:10534       | 0.7789       | 0.785        |
| nasopharynx carcinoma      | DOID:9261        | 0.6279       | 0.6607       |
| lung adenocarcinoma        | DOID:3910        | 0.6865       | 0.7433       |
| squamous cell carcinoma    | DOID:1749        | –            | 0.7354       |
| malignant glioma           | DOID:3070        | 0.8566       | 0.7659       |
| ovarian cancer             | DOID:2394        | 0.7364       | 0.5976       |
| cancer                     | DOID:162         | –            | 0.7192       |
| breast cancer              | DOID:1612        | 0.7534       | 0.7847       |
| melanoma                   | DOID:1909        | –            | 0.6667       |
| large intestine cancer     | DOID:5672        | 0.7859       | 0.7275       |
| non-small cell lung cancer | DOID:3908        | 0.6538       | 0.7782       |
| **Average value of AUC**   |                  | **0.7282**   | **0.726**    |
Conclusions

In this article, a computational model for potential disease-related lncRNA identification was proposed based on multiple biological datasets. The results showed that LncDisAP was proven to be successful in predicting novel disease-related lncRNA signatures with an average AUC value of 78.08% and can be an effective solution to improve the quality of disease diagnostics and treatments. To further evaluate the performance of our computational model, we used several common cancers as case studies. We found some unknown lncRNAs that could be associated with these diseases through our network. In addition, we discussed the impact of different data sources and different test sets on the performance of the disease-related lncRNA functional network in predicting disease-lncRNA pairs.

Supplementary information

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Additional file 1: Statistics of 54 diseases for evaluating disease-related lncRNA functional network.

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Authors’ contributions

WYT and JL did data collection and preprocessing. And with the guidance of ZT and WYD, WYT finished the algorithm design and validation. WYT and PJ was the major contributors in writing the manuscript. All authors have read and approved the final version of the manuscript.

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Availability of data and materials

All the datasets used in this paper could be downloaded from websites.

Ethics approval and consent to participate

Not Applicable.

Consent for publication

Not Applicable.

Competing interests

The authors declare that they have no competing interests.

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