Network medicine analysis of chondrocyte proteins towards new treatments of osteoarthritis

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Osteoarthritis (OA) is a progressive disorder with high incidence in the ageing human population that still has no treatment currently. This disorder induces the breakdown of articular cartilage, leading to the exposure and damage of bone surfaces. For a global understanding of OA development, the systematic integration of known OA-related proteins with protein–protein interaction (PPI) networks is required. In this work, the OA-related interactome was reconstructed using multiple data sources to have the most up-to-date information on OA-related proteins and their interactions. We then combined emergent concepts in network medicine to detect new unclassified OA-related proteins. The mapping of known OA-related proteins with PPI networks showed that these proteins are locally connected to each other and agglomerated in a large component. To expand this module, we applied a diffusion-based algorithm that probabilistically induces more searches in the vicinity of the seed OA-related proteins. As a result, the 10 topmost ranked proteins were connected to the OA disease module, supporting the local hypothesis. We computed structural modules and selected those that had the highest enrichment of OA-related proteins. The identified molecules show a link between structural topology and disease dysfunctionality. Interestingly, the protein Q6EEV6 was highlighted for OA association by both methods, reinforcing the potential involvement of this protein. These results suggest that similar disease-connected modules may exist in different human disorders, which could lead to systematic identification of genes or proteins that have a joint role in specific disease phenotypes.

1. Introduction

Osteoarthritis (OA) is a progressive disorder mainly characterized by the breakdown of articular cartilage, leading to the exposure and damage of bone surfaces [1]. OA is a widespread cause of joint degeneration, mostly affecting synovial joints, such as those found in the knees, hips and fingers, and is one of the leading disabling human conditions worldwide. Despite its high incidence, no treatment currently exists that can replace damaged cartilage or prevent cartilage degeneration [2]. A combination of various anti-inflammatory drugs that reduce joint inflammation is currently the best method to treat OA patients, although the most effective treatment appears to be weight loss to reduce the stress on joint cartilage. This lack of appropriate pharmaceutical intervention suggests that new approaches are needed to understand the functional perturbations linked to OA and to identify key proteins that can be targeted for therapeutic use.

Cartilage is a type of connective tissue found in the nose and ears, as well as in joints, such as the knees and fingers. Three major types are found in the human body, including articular (hyaline), elastic and fibrocartilage. Articular cartilage is the most prevalent and is found at the surface of synovial joints, primarily functioning to facilitate joint motion and distribute the forces of weight bearing [3]. Chondrocytes are the only cell population found within mature articular cartilage, constituting approximately 2–5% of the total tissue, with the rest consisting of a tough and flexible extracellular matrix (ECM). Chondrocytes control matrix turnover through the production and release of collagen, proteoglycans and enzymes involved in cartilage metabolism [4]. Chondrocytes have been the...
most studied component of the disease system to determine how changes in their function may be linked to OA. Whereas healthy articular cartilage is believed to be permanent, consisting of chondrocytes with low metabolic activity, in conditions, for example OA, they enter a stage of maturation becoming proliferative and then hypertrophic chondrocytes [5]. In this state, chondrocytes increase ECM synthesis but cannot compensate for the increased matrix degradation by proteolytic matrix enzymes. This leads to an overall decrease in matrix synthesis resulting in the breakdown of articular cartilage.

Biomedical knowledge combined with network science tools aim to identify genes or proteins that can be probably associated with specific disease phenotypes [6,7]. Several computation-based methods have been proposed to uncover disease-associated genes. Disease candidate molecules were prioritized using linkage methods [8–10], where macromolecules that interact directly with a disease protein were assumed to be associated with the same disorder. Another approach seeks to exploit the fact that molecules which belong to the same structural and functional module may have a joint role. The extension of this concept to disease phenotypes means that identifying an unclassified protein that belongs to a disease module with high enrichment over a random selection may indicate that the protein is likely to be involved in the same disorder [11,12].

Diffusion-based methods were also used to identify proteins that are related to specific disorders. The random walk particles navigate through the complex interactome structure, starting from those protein products originated by known disease genes. The nodes most often visited by the random walkers are considered to be more related to the known disorder genes. Several algorithmic versions of this concept have been applied to human diseases, from prostate cancer to Alzheimer’s disease [13,14]. One of the leading network-based diffusion algorithms is the PageRank, which is not only well known to rank web content but can also potentially classify any type of nodes given a suitable network structure [15,16]. The algorithm does not rank nodes using their intrinsic features, rather it uses the collective wisdom of the network, considering each link as a node that has a direct vote to increase the node’s ranking. Recently, the PageRank algorithm has been successfully used to identify disease-causing genes based on protein interactions, to detect cancer genes in protein networks and to classify the metabolic network of the tuberculosis bacterium [17–19]. It is expected that the advent of full-genome sequencing and genome-wide association studies can help to provide more accurate disease gene (seeds) as well as better validation methods.

Network and systems approaches have not been widely applied to complex skeletal diseases as arthritis so far. A molecular interaction map was presented for processes ongoing in patients affected by rheumatoid arthritis (RA) based on the results of functional genomic analyses and pathways available in the literature [20], which identified a new potential drug target for the treatment of RA [21]. Gene expression data from pooled joint tissue were overlaid with a protein interaction network to identify components that were significantly upregulated during the development of OA [22]. However, for a more global understanding of OA development, the systematic integration of known OA-related proteins with protein interaction networks is required.

In this study, we combine the recently introduced concepts in network medicine (structural proximity, modularity and diffusion algorithmic approach) to detect new unclassified OA-related proteins using protein interaction networks [6]. We first constructed a manually curated interactome network composed of all proteins linked to at least one chondrocyte protein. We then overlaid OA-related proteins onto this protein interaction network and showed that the non-random placement of the OA-related proteins in the interactome offers a new avenue for the prediction of OA relation. Next, the fully connected component composed of only OA-related proteins was extracted and the PageRank-based diffusion algorithm, with a personalized vector constructed using the known OA-related proteins, was applied to the interactome network. The personalized PageRank algorithm has already been used for biological applications in the context of protein interaction networks [18].

The results show that the 10 topmost ranked proteins, excluding natural overlaps with personalized vector proteins, are neighbours of the main OA-related connected component, which leads to an expansion of the OA disease module. Moreover, the computation of the modularity applied to the full network leads to the identification of a set of modules, in which the enrichment of OA-related proteins over a random selection was significantly higher. Among them, one module shared a protein already identified in the 10 topmost ranked proteins, leading to the same prediction using two independent methods.

2. Material and methods

(a) Datasets

To construct a suitable network for the investigation of OA, we first used the complete list of protein–protein interactions from the Human Protein Reference Database (HPRD) [23] as a baseline protein interaction network. We then used several large-scale proteomic studies to identify proteins that were detected in healthy and osteoarthritic chondrocytes. Seventeen studies were used in total (table 1), among which nine focused on healthy chondrocyte proteomics, whereas the remaining eight contained data about chondrocyte proteins that had their expression altered in OA. All 17 studies were used to build the chondrocyte protein interaction network, as it was assumed that proteins detected in OA cells were still present in healthy cells.

Each protein identified in these proteomic studies was queried within the HPRD interaction network to extract all proteins interacting with the query protein, assuming that if one interaction partner is present in chondrocytes, the second partner must be present too.

Table 1. List of proteomic studies used to construct the chondrocyte protein interactome.

<table>
<thead>
<tr>
<th>healthy chondrocyte proteome</th>
<th>altered proteins in OA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Akiyama et al. [24]</td>
<td>Gobeze et al. [25]</td>
</tr>
<tr>
<td>Ciller-Pasto et al. [26]</td>
<td>Guo et al. [27]</td>
</tr>
<tr>
<td>Guo et al. [28]</td>
<td>Lambrecht et al. [29]</td>
</tr>
<tr>
<td>Jmeian &amp; El Rassi [30]</td>
<td>Ling et al. [31]</td>
</tr>
<tr>
<td>Peca et al. [32]</td>
<td>Ma et al. [33]</td>
</tr>
<tr>
<td>Ruiz-Romero et al. [34]</td>
<td>Rosenthal et al. [35]</td>
</tr>
<tr>
<td>Ruiz-Romero et al. [36]</td>
<td>Ruiz-Romero et al. [37]</td>
</tr>
<tr>
<td>Ruiz-Romerso et al. [38]</td>
<td>Wu et al. [39]</td>
</tr>
<tr>
<td>Vincourt et al. [40]</td>
<td></td>
</tr>
</tbody>
</table>
for the interaction to occur, even though the second partner may not necessarily have been detected by the proteomic analysis. This process resulted in a protein interaction network specific to chondrocyte cells. All proteins names were converted to UniProt accession identifiers in order to eliminate any duplication owing to different naming conventions. Proteins that have altered levels in OA were extracted from the eight OA studies and labelled by a specific attribute. The complete network comprised 2955 nodes, among which 168 were related to OA (figure 1). Background data for the full network is provided in the electronic supplementary material.

(b) PageRank algorithm with personalized vector

The PageRank algorithm describes a diffusion process, where a particle moves by following the links of the network and jumps to randomly selected nodes with given probabilities. These probabilities are parameters of the model. As a result, the frequency of visitation of a node by the particle at the stationary state of this diffusion process is the PageRank of the node and it is used to determine its ranking relative to other nodes of the network [15,16].

The fundamental PageRank algorithm can be described using the following equation:

\[ \pi^{(k+1)T} = \pi^{(k)T}G, \]

where \( \pi^{(k)T} \) is the PageRank row vector at the \( k \)th iteration. Each component of the vector is associated with a node, therefore \( \pi^{(k)T} \) gives the probability that the particle is at each node of the network at time step \( k \). The matrix \( G \) can be written as

\[ G = (1 - \alpha)S + \alpha \frac{ee^T}{n}, \]

where \( e^T \) is the row vector of all 1s and \( \alpha \) is a parameter that indicates the proportion of time the random surfer follows a random teleportation process, or conversely \( 1 - \alpha \) is the proportion of time it is guided through the links on the network structure. The stochastic matrix \( S \) reads as \( S = H + \alpha (1/n e^T) \), which is constructed using the row normalized probability transition matrix \( H \) and the dangling node vector \( a \). This column vector has 1 s in components associated with nodes with no-outgoing links and 0 s for the rest of the components.

Here, we can distinguish between random teleportation \( \alpha_T \) and personalized teleportation \( \alpha_P \) probabilities [18]. Each \( i \) component of the personalized vector takes the value \( 1/W \) if the node \( i \) is a known OA-related protein, and 0 otherwise. \( W \) is a normalizing weighted factor that represents the total number of OA-related proteins.

Then, equation (2.2) can read as

\[ G = (1 - \alpha_T - \alpha_P)S + (\alpha_T + \alpha_P)\frac{ee^T}{n}. \]

By substituting the expression of matrix \( S \) and by defining a personalized vector as \( v^T = e^T/n \), then equation (2.1) reads as

\[ \pi^{(k+1)T} = (1 - \alpha_P - \alpha_T)\pi^{(k)T}H + [(1 - \alpha_T - \alpha_P)\pi^{(k)T}a + \alpha_T e^T/n + \alpha_P v^T. \]

Our computations were performed using \( \alpha_T = \alpha_P = 0.15 \) and a value of \( \epsilon = 0.0001 \) for the error of convergence of the algorithm at the stationary state.

(c) Statistical significance of the observed size of the osteoarthritis-related connected component

The OA-related network was randomized using a shuffling algorithm that exchanges edges in the network but preserves the degree distribution. For each of the 100 network samples, we generated with the same size as the observed OA-related network, we performed 200 shuffling edges steps. Then, the mean value and standard deviation of the observed giant connected component were computed. The statistical significance of the analysis was examined using the two-tailed \( p \)-value, as shown in the Results section.

(d) Modularity algorithm

To identify topological modules, we used a multi-scale method in which modularity is optimized by means of a greedy local
The nodes that form a discovered partition are then merged and a supernode for each new partition is created. The process is repeated in the re-scaled network [41].

The algorithm starts with as many different modules as there are nodes. For each node \( i \), the gain of modularity is evaluated when removing node \( i \) from its module and by assigning it to another module \( j \). The node \( i \) will be finally assigned to the community that offers the highest gain and is positive. The process is applied for all nodes until no new gains can be achieved. The modularity of the network will increase in each iteration and tends to maximize the following modularity functional form:

\[
Q = \frac{1}{2m} \sum_{i,j} \left( A_{ij} - \frac{k_i k_j}{2m} \right) \delta(c_i, c_j),
\]

where \( m \) is the number of edges, \( A_{ij} \) is an element of the adjacency matrix of the network, \( k_i \) denotes the degree of node \( i \) and \( c_i \) is the community to which node \( i \) is assigned. The \( \delta(x, y) \) function is 1 if \( x = y \) and 0, otherwise.

(e) Community analysis
The detection of communities in networks is a common method to identify groups of closely related entities or functional modules. We applied a community detection algorithm by Blondel et al. [41] to detect groups of highly connected OA-related proteins in the protein–protein interaction (PPI) network. This algorithm constructs a hierarchical community structure for the network, where modularity is maximized at each level. We focused on the first level and tested the detected communities for OA enrichment. The probability of detecting a number \( n \) of OA-related proteins in a community of size \( N \), considering that the ratio of OA-related proteins in the full network was 0.0569 (168/2955), was calculated using the hypergeometric distribution. A community that was deemed to be significantly enriched in OA-related proteins if this probability was lower than 0.05. As there is generally no unique solution to decompose the network into communities maximizing modularity, the exact composition of communities may vary when the algorithm is run several times. We therefore repeated the analysis 10 times and retained for each significant community the composition observed the largest number of times.

3. Results
(a) The osteoarthritis-related proteins are locally connected to each other and agglomerated in a large component
The assembled interactome network is composed of 2955 nodes, of which 168 are known OA-related proteins. The visualization of the giant connected component of the network can be seen in figure 1. Note that 37 nodes (including six OA-related proteins) and 21 edges did not belong to the main component so they are absent in figure 1. The total number of links is 5477 with 151 links connecting two OA-related proteins (red), 3310 connecting OA-related to non-OA-related proteins (orange) and 2016 connecting non-OA-related proteins to each other.

Evidence of the non-random localizations of disease genes is necessary in our network-based approach to identify unclassified disease-related molecules. To investigate whether the OA-related proteins were distributed randomly in the network, we extracted all the OA-related proteins by keeping only edges that were connecting two OA-related proteins. The resulting network is shown in figure 2. Out of 168 proteins, 162 are connected to the main component of the interactome and 89 nodes form an OA-related connected component. These results show that the OA-related proteins tend to be locally connected to each other and a large fraction of them are agglomerated in a large component (table 2).

The analysis of the degree distribution of the full interactome network reveals a non-random structure. The network follows a power law, a characteristic pattern of scale-free networks. On the other hand, the OA-related network, which is of smaller size, decays faster than a power law. The cumulative degree distributions of the networks are shown in figure 3. The fact that the full interactome network follows a scale-free distribution also allows the use of diffusion-based algorithms to analyse the network.

(b) The osteoarthritis-related connected component statistically emerges as disease module seed to be expanded
To evaluate the significance of the OA-related connected network, we performed two statistical analyses (see Material and methods for details). The network was randomized with 2000 shuffling edges for each of the 100 network samples. The component size distribution for randomized networks is shown to be normal distribution (figure 4, light blue curve), which has a mean value of 97 and a standard deviation of 2.45. The results show that the observed giant component (89) is over 3 s.d. from the mean of a distribution (figure 4, light blue curve), which has a mean value of 97 and a standard deviation of 2.45. The results show that the observed giant component (89) is over 3 s.d. from the mean of a randomized network sample (97). The statistical significance analysis leads to a two-tailed \( p \)-value of 8.70 \( \times 10^{-3} \). This indicates that the observed size of the OA-related connected component would be highly unlikely in a random network.

(c) Expansion of the osteoarthritis-related connected component
It has been hypothesized that molecular routes can overlap with the shortest paths between known disease-related molecules. This network parsimony principle has supported the idea of using diffusion-based algorithms. We therefore applied the PageRank algorithm with personalized vector to the interactome network in order to predict new disease candidate proteins that are likely to be associated with OA. As shown in a recent work, the success of the PageRank algorithm in ranking nodes in large networks is rooted in the fact that the degree distribution follows a power law, which
makes the top-ranked node easier to find and the ranking more reliable when the networks grow in size [42]. Here, the personalized vector is simply a vector that includes the known OA-related proteins. This vector takes a non-zero weighted probability for each protein that is associated with OA, and a zero score for the remaining proteins, where the sum of all scores is normalized to one. The PageRank results showed that several already known OA-related proteins appeared in the top ranks. Therefore, we selected the 10 top-ranked proteins excluding known OA-related proteins (table 3). These proteins were added to the OA-related network and are shown in figure 5 (green nodes). Interestingly, the results show that the identified proteins are in the neighbourhood of the OA-related connected component and always adjacent to at least one OA-related protein. This fact supports the idea of a disease module that can be expanded using the collective wisdom of the full interactome network, unveiling new potential disease-protein candidates.

(d) Module enrichment analysis

In network medicine, the disease module hypothesis assumes that the molecules related to a specific disorder tend to be located in the same network neighbourhood, forming a structural disease module. To support our analysis done on new identified OA-related proteins using a diffusion algorithmic method, we used a different approach based on maximizing modularity in complex networks that detects modules with the highest enrichment of OA-related proteins.

### Table 2. Main statistical network features of the analysed networks.

<table>
<thead>
<tr>
<th></th>
<th>full protein network (giant connected component)</th>
<th>OA-related extracted network</th>
<th>OA-related giant connected component</th>
</tr>
</thead>
<tbody>
<tr>
<td>nodes</td>
<td>2955 (2918)</td>
<td>168 (89)</td>
<td>89</td>
</tr>
<tr>
<td>average degree</td>
<td>3.7</td>
<td>1.79</td>
<td>3.2</td>
</tr>
<tr>
<td>clustering degree</td>
<td>0.069</td>
<td>0.092</td>
<td>0.174</td>
</tr>
<tr>
<td>average path length</td>
<td>4.3</td>
<td>4.8</td>
<td>4.8</td>
</tr>
</tbody>
</table>

Figure 3. (a) The cumulative degree distributions for both the fully connected network and the extracted OA-related connected network.

![Figure 3](image)

Figure 4. Statistical significance analysis of the giant connected component. The expected value obtained in a randomized network (blue curve) is over 3 s.d. from the observed value. The statistical significance analysis gives a two-tailed *p*-value of $8.70 \times 10^{-3}$ (blue curve).

Figure 4. (b) The expected degree distribution for the OA giant component.

The detection of communities in networks is a common method to identify groups of closely related entities or functional modules. We applied a community detection algorithm by Blondel et al. [41] to detect groups of highly connected OA-related proteins in the PPI network. This method optimizes modularity using a greedy local algorithm, where nodes are iteratively exchanged between modules as long as the modularity can be increased and is positive. This method is widely used owing to its efficiency to deal with large networks as...
well as the high quality of results it provides. It scored the highest rank when evaluated in computer-generated benchmark problems [43,44].

We focused on those modules that showed highest enrichment of OA-related proteins. Overall, four significant communities enriched in OA-related proteins were detected (figure 6). Ranked from the highest to lowest significance, the first community contains three OA-related proteins out of seven ($p = 0.005$) (figure 6b), the next contains three OA-related proteins out of 11 ($p = 0.021$) (figure 6c), the next contains two OA-related proteins out of five ($p = 0.029$) (figure 6d) and the last contains two OA-related proteins out of six ($p = 0.041$) (figure 6e). All non-OA-related proteins belonging to these communities are therefore likely to be subjected to disease perturbations owing to their high interconnectivity with disease-associated proteins.

We then filtered the proteins found in these modules to investigate whether there is an overlap between them and the 10 top-ranked proteins detected by the diffusion algorithm (table 3). Interestingly, one protein was predicted by these two independent methods, namely Q6EEV6 (figure 7). The probability for this simultaneous detection to occur randomly is only $2 \times 10^{-5}$. Q6EEV6 is the small ubiquitin-related modifier 4 (SUMO-4).

### Table 3. List of top 10 candidate proteins predicted to be OA-related by the PageRank algorithm, excluding known OA-related proteins.

<table>
<thead>
<tr>
<th>UnitProt</th>
<th>standard name</th>
<th>main biological activity</th>
</tr>
</thead>
<tbody>
<tr>
<td>P26641</td>
<td>elongation factor 1-γ (EEF1G)</td>
<td>protein biosynthesis</td>
</tr>
<tr>
<td>Q6EEV6</td>
<td>small ubiquitin-related modifier 4 (SUMO-4)</td>
<td>ubiquitin conjugation pathway</td>
</tr>
<tr>
<td>P17252</td>
<td>protein kinase C-α (PKCA)</td>
<td>angiogenesis, regulation of apoptosis</td>
</tr>
<tr>
<td>P61981</td>
<td>14-3-3 protein gamma (YWHAH)</td>
<td>cellular membrane regulation</td>
</tr>
<tr>
<td>P68104</td>
<td>elongation factor 1-α (EEF1A1)</td>
<td>protein synthesis</td>
</tr>
<tr>
<td>P28482</td>
<td>mitogen-activated protein kinase 1 (MAPK1)</td>
<td>transcription regulation</td>
</tr>
<tr>
<td>P63104</td>
<td>14-3-3 protein zeta/delta (YWHAZ)</td>
<td>cellular membrane organization</td>
</tr>
<tr>
<td>P35222</td>
<td>catenin β-1 (CTNNB1)</td>
<td>cell adhesion, transcription regulation</td>
</tr>
<tr>
<td>P31946</td>
<td>14-3-3 protein β/α (YWHAH)</td>
<td>cellular membrane organization</td>
</tr>
<tr>
<td>P62736</td>
<td>actin, aortic smooth muscle (ACTA2)</td>
<td>cell motility</td>
</tr>
</tbody>
</table>

Figure 5. Expansion of the OA-related giant component by adding the 10 top-ranked proteins by PageRank analysis excluding known OA-related proteins (green nodes). Links between newly identified proteins are shown in green, whereas links between new proteins and known OA-related proteins are shown in yellow.
associated with susceptibility to type I diabetes and cellular response to oxidative stress, but no explicit link to cartilage degradation or skeletal diseases was found. In total, SUMO-4 was found to interact with 15 OA-related proteins, including several proteins involved in glycolysis and redox regulation (table 4). The potential involvement of this protein in OA is hereby reinforced.

4. Discussion and conclusion

Network medicine is based on several widely used principles related to network structure and functionality. The local hypothesis or structural proximity suggests that proteins, which belong to the same disorder, are assumed to interact with each other. Next, the disease module hypothesis assumes that the molecules related to a specific disorder tend to be located in the same network neighbourhood, leading to a structural disease module. Moreover, molecular routes often overlap with the shortest paths between known disease-related molecules. This network parsimony principle has encouraged the usage of diffusion-based algorithms. While the structural proximity has been very well experimentally supported, the existence of disease modules overlapping with structural modules and the parsimony principles have been less well quantified or verified.

In this work, we have combined these three organizing principles to analyse the OA-related network. The interactome was reconstructed using multiple data sources and manual curation to have the most up-to-date information on OA-related proteins and their interactions. First, the integration of known OA-related proteins with protein interaction networks has shown that the OA-related proteins are locally connected to each other and are furthermore agglomerated in a large component. This component may represent the seed of the OA
disease module, which can be expanded by identifying OA-related proteins in the neighbourhood. To this end, we have applied a diffusion-based algorithm with a personalized vector that probabilistically induces more searches in the structural vicinity of the seed OA-related proteins. As shown in Ivan & Grolmusz [18], this feature is essential to successfully identify new candidate disease proteins in protein–protein interaction networks. In addition, we also allowed random teleportation to explore distant locations on the network with the same probability. As a result, the 10 topmost ranked proteins were connected to the OA disease module, supporting the local hypothesis.

To investigate the overlap between the structural and disease module, we computed structural modules and selected those ones that had the highest enrichment of OA-related proteins. The identified molecules show a link between structural topology and disease dysfunctionality. In addition, they confirm the local hypothesis because they are also connected to the main OA-related connected component. Interestingly, the protein Q6EEV6 (SUMO-4) was identified among the top 10 candidates for OA association by the PageRank analysis and also by the modularity analysis. Note the complex interconnectivity in the protein’s neighbourhood.

Table 4. List of OA-related interacting partners of the Q6EEV6 (SUMO-4) protein.

<table>
<thead>
<tr>
<th>UnitProt</th>
<th>standard name</th>
<th>main biological activity</th>
</tr>
</thead>
<tbody>
<tr>
<td>P00558</td>
<td>phosphoglycerate kinase 1 (PGK1)</td>
<td>glycolysis</td>
</tr>
<tr>
<td>P11142</td>
<td>heat shock cognate 71 kDa protein (HSP8)</td>
<td>transcription regulation stress response</td>
</tr>
<tr>
<td>P04406</td>
<td>glyceraldehyde-3-phosphate dehydrogenase (GAPDH)</td>
<td>glycolysis translation regulation</td>
</tr>
<tr>
<td>P08107</td>
<td>heat shock 70 kDa protein 1A/1B (HSPA1A/1B)</td>
<td>protein folding stress response</td>
</tr>
<tr>
<td>P30048</td>
<td>thioredoxin-dependent peroxide reductase, mitochondrial (PRDX3)</td>
<td>redox regulation</td>
</tr>
<tr>
<td>P30041</td>
<td>peroxiredoxin-6 (PRDX6)</td>
<td>redox regulation</td>
</tr>
<tr>
<td>P07195</td>
<td>l-lactate dehydrogenase B chain (LDHB)</td>
<td>glycolysis</td>
</tr>
<tr>
<td>P21796</td>
<td>voltage-dependent anion-selective channel protein 1 (VDAC1)</td>
<td>ion transport</td>
</tr>
<tr>
<td>P10809</td>
<td>60 kDa heat shock protein, mitochondrial (HSPD1)</td>
<td>protein folding</td>
</tr>
<tr>
<td>P06733</td>
<td>α-enolase (ENO1)</td>
<td>glycolysis transcription regulation</td>
</tr>
<tr>
<td>Q0UZ1</td>
<td>stromatin-like protein 2, mitochondrial (STOML2)</td>
<td>mitochondrial regulation</td>
</tr>
<tr>
<td>P11021</td>
<td>78 kDa glucose-regulated protein (HSP85)</td>
<td>protein complex assembly</td>
</tr>
<tr>
<td>P02545</td>
<td>prelamin-A/C (LMNA)</td>
<td>nuclear assembly</td>
</tr>
<tr>
<td>00299</td>
<td>chloride intracellular channel protein 1 (CLIC1)</td>
<td>ion transport</td>
</tr>
<tr>
<td>P30101</td>
<td>protein disulfide-isomerase A3 (PDIA3)</td>
<td>redox regulation</td>
</tr>
</tbody>
</table>
top 10 candidates for OA association by the PageRank analysis and was also detected in the modular analysis. SUMO-4 mutations were previously associated with other diseases. A study by Guo et al. [45] suggested that a substitution in the SUMO-4 gene is associated with increased risk of type I diabetes mellitus. While subsequent studies confirmed these findings in Asian populations [46,47], these observations were not consistent in European Caucasians [48], suggesting that further study is required to fully elucidate the relationship between SUMO-4 mutations and type I diabetes mellitus. The effect of SUMO-4 mutations in RA and systemic lupus erythematosus has been investigated, showing no consistent sign of predisposition to these diseases [49–51].

Several glycolytic enzymes were found among the top candidate proteins predicted by our analysis to be involved in OA. The importance of glycolysis in the prevention of cartilage breakdown and responses to apoptotic processes has already been suggested [29,37]. Chondrocytes require a large amount of energy to regulate the cartilage ECM through the release of proteins, such as collagens and proteoglycans. Perturbation of this cellular function may lead to a decrease in ATP production, thereby preventing the normal control of matrix turnover and contributing to the gradual breakdown of cartilage, as the decreased ECM is no longer as effective in distributing the forces of weight bearing.

Other proteins are involved in the stress response and regulation of apoptosis. The link between cartilage degeneration and chondrocyte apoptosis has already been described and is thought to play an important role in the pathogenesis of OA [52]. In particular, the level of protein kinase C-β (P17252) was found to be significantly increased in human knee OA cartilage by a recent study [53] in correlation with an increase in chondrocyte apoptosis. This observation confirms that our network analysis is a valid approach to identify candidate proteins for disease association, as the link between P17252 and OA is too recent to have been included in databases and did not appear in our original dataset.

The third main group of proteins is related to cellular adhesion and mechanotransduction. Actins are ubiquitously expressed in eukaryotic cells and changes in the cytoskeleton are believed to be associated with the onset of the disease [54]. Protein kinase C-α itself is known to be involved in the regulation of cell adhesion and motility [55]. Catenin β-1 (P35222) is another detected protein that is involved in the regulation of cell adhesion through the Wnt signalling pathway. Although a direct link between catenin β-1 and OA was not established, the activation of catenin β-1 signalling was observed to lead to premature chondrocyte differentiation in mice, which is a symptom of OA [56,57].

To summarize, we have reconstructed a large-scale OA-related interactome, in which the module of OA-related proteins emerged as potential disease seed. The presence of a main connected component of OA-related proteins enables the discovery of newly identified proteins using network diffusion algorithms. The promising results shown here suggest that similar disease-connected modules may also exist in different human disorders. The curation of a high volume of data for other disorders and its overlapping with the background interactome network could also lead to highly connected disorder modules (seeds) that could be expanded using a rich variety of computational methods [43]. This network medicine approach could lead to systematic identification of genes or proteins that can have a joint role in a specific disease phenotype.

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References

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