GRASP: Graph Alignment through Spectral Signatures

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Abstract. What is the best way to match the nodes of two graphs? This graph alignment problem generalizes graph isomorphism and arises in applications from social network analysis to bioinformatics. Some solutions assume that auxiliary information on known matches or node or edge attributes is available, or utilize arbitrary graph features. Such methods fare poorly in the pure form of the problem, in which only graph structures are given. Other proposals translate the problem to one of aligning node embeddings, yet, by doing so, provide only a single-scale view of the graph.

In this paper, we transfer the shape-analysis concept of functional maps from the continuous to the discrete case, and treat the graph alignment problem as a special case of the problem of finding a mapping between functions on graphs. We present GRASP, a method that first establishes a correspondence between functions derived from Laplacian matrix eigenvectors, which capture multiscale structural characteristics, and then exploits this correspondence to align nodes. Our experimental study, featuring noise levels higher than anything used in previous studies, shows that GRASP outperforms state-of-the-art methods for graph alignment across noise levels and graph types.

1 Introduction

Graphs model relationships between entities in several domains, e.g., social networks, protein interaction networks, email communication or chemical molecules. The structure of such graphs captures rich information on how people are connected, how molecules function, or how proteins interact.

At the same time, the expressive nature of graphs also implies complexity, which renders some fundamental problems hard. For instance, the graph isomorphism problem, which is to determine whether two graphs share the same structure is neither known to be polynomially solvable nor NP-complete, and
has been used to define the GI complexity class [17]. Problems that generalize graph isomorphism occur frequently in the field of graph analytics. One of those is the NP-complete subgraph isomorphism problem; another is graph alignment, which aims to find the best (exact or inexact) matching among the nodes of a pair of graphs; a solution to this problem is sine qua non in tasks such as identifying users in different social networks [14], matching objects in images by establishing feature correspondences and comprehending protein response in the body [15].

In case additional background information is available, such as node and edge attributes in the two graphs to be aligned, or existing valid seed matches, then the problem is solvable via supervised methods [24,6]. However, in case only graph structures are given, then the problem of aligning two graphs by matching structures, is at least as hard as graph isomorphism even in its approximate version [1].

Existing approaches to the graph alignment problem are oriented toward using a few heuristic graph features, such as landmarks, in order to detect a good alignment [12], exploiting additional information such as node attributes [35] or bipartite networks [20], or optimizing objectives based only on local connections among nodes [10,27,22]. On the other hand, the spectra of Laplacian matrices have been successfully employed to devise a similarity measure among graphs [31]. Laplacian spectra capture important multiscale properties, such as local-scale ego-nets and global-scale communities. Previous approaches rooted in spectral characteristics decompose large matrices expressing all alignments among edges in two graphs [10,27,22] and formulate the solution as finding the leading eigenvector of such matrices. These approaches disregard most eigenvectors and consider only local edge variations. To our knowledge, the spectral properties of Laplacian matrices have not yet been utilized to any significant extent for an end-to-end graph alignment method.

We propose GRASP, short for GRaph Alignment through SPectral Signatures, a principled approach towards detecting a good alignment among graphs, grounded on their spectral characteristics, i.e., eigenvalues and eigenvectors of their Laplacian matrices [7]. We transfer the methodology of matching among
shapes based on corresponding functions \[28\] to the domain of graphs: we first extract a mapping of node-evaluated functions grounded on the graph’s heat kernel, and then apply this mapping to the matching on nodes. Figure 1 shows an example alignment of the Karate club with a deteriorated version obtained by removing some edges; GRASP correctly aligns most of the nodes, while REGAL \[12\] based on local descriptors fails to do so.

In short, we propose GRASP, a graph alignment method based on spectral graph characteristics and show its effectiveness in recovering real-graph alignments, with higher accuracy and similar runtime as the state of the art.

2 Related Work

We discuss related work in two main categories: restricted alignment, which requires ground-truth mapping or other additional information, and unrestricted alignment, which requires neither supervision nor additional information. Table 1 gathers together previous works’ characteristics.

2.1 Restricted Alignment

Restricted methods incorporate non-structural information; a restricted method can be supervised or assisted.

Supervised methods exploit pre-aligned pairs of seed nodes to construct a first alignment. Percolation graph matching (PGM) \[14,33\] propagates ground-truth alignments across the network. Representation learning approaches, such as IONE \[24\], PALE \[26\], and DeepLink \[36\], learn a low-dimensional embedding of the graph nodes and map the node embeddings of one graph to another. A similar method aligns multiple networks at once \[6\]. Active network alignment \[25\] applies active learning to elicit expert guidance on alignments. Overall, supervised methods rely on prior knowledge, which may not be available.

Assisted methods utilize auxiliary information or structural constraints. Bi-Align \[20\] focuses on bipartite graphs; however, most graphs are not bipartite. FINAL \[35\] aligns nodes based on similarity of topology and attributes. GSANA \[34\] lets pairwise distances to seed nodes guide the matching. Another variant matches weighted matrices using their spectra \[32\]; that is inapplicable to the unweighted case. Overall, restricted methods cannot handle cases where the only given information is graph structure.

2.2 Unrestricted Alignment

Unrestricted methods require neither prior knowledge of ground-truth pairs nor other information on the input graph.

Integer-programming methods. Klau \[15\] presents a Lagrangian relaxation for the integer programming problem posed by network alignment; the resulting algorithm is polynomial, yet still impracticable for large networks.
Table 1: Related work in terms of present (✔) and absent (✘) properties. **Supervised** methods [24,6,26,36] require aligned nodes as input. **Spectral** methods [27] use spectral properties of alignment matrices. **FINAL** [35] does not work on plain graph structures as it requires node attributes. **REGAL** [12] and IsoRank [22,30] are flexible in allowing different algorithms for alignment (e.g. bipartite matching, nearest neighbors). Among unrestricted methods (rows 4–7), **LREA** [27] cannot benefit from offline precomputation (results in Figure 6). **GRASP** explicitly captures multiscale properties through the heat kernel.

**Embedding-based methods.** **REGAL** [12] constructs node embeddings based on the connectivity structure and node attributes, and uses the similarity between these features for node alignment; we classify REGAL as an unrestricted method since it can work without attributes. **CONE-Align** [5] realigns node representations, without prejudice to the representation used.

**Matrix decomposition methods.** **IsoRank** [30] aligns multiple protein-protein interaction networks aiming to maximize the overall quality across all input networks; it constructs an eigenvalue problem for every pair of input networks and extracts a global alignment across a set of networks by a \( k \)-partite matching; it uses structural properties (PageRank), but also relies on a similarity measure between nodes which in a biology-specific case builds on the similarity of the proteins; it is improved with greedy approaches [18]. Another improvement on IsoRank, **IsoRankN** [22], performs spectral clustering on the induced graph of pairwise alignment scores; as it is based on spectral methods, IsoRankN is claimed to be both error-tolerant and computationally efficient. **EigenAlign** [10] formulates the problem as a Quadratic Assignment Problem that considers both matches and mismatches and solves it by **spectral decomposition** of matrices. Building thereupon, **Low-Rank EigenAlign** [27] solves a maximum weight bipartite matching problem on a low-rank version of a node-similarity matrix, hence requires memory linear in the size of the graphs. However, EigenAlign variants use the first eigenvector of a joint adjacency matrix between the two graphs to be aligned, rather than the eigenvectors of graph Laplacians, which provides richer information.

**Belief propagation methods.** **NetAlign** [3] solves a sparse graph alignment variant by message passing.
2.3 Shape Matching

Our work is inspired by shape matching methods that employ spectral properties [23,28,21]. Functional maps [28] generalize the matching of points to the matching of corresponding functions among shapes, by revealing a common decomposition of such functions using the eigenvectors of the Laplace-Beltrami operator; the graph equivalent of that operator is a graph’s Laplacian matrix. Extensions of this methods match non-isometric shapes by aligning their Laplace-Beltrami operators’ eigenbases [21], and match a part of a shape to another full shape in the spectral domain [23] without requiring spatially modeling the part of a shape.

2.4 Spectral Methods

Graph spectra [7] facilitate problem-solving in graph analysis, image partitioning, graph search, and machine learning [29]. NetLSD [31] uses Laplacian spectral signatures to detect graph similarity, but not to align graphs, in a multi-scale fashion. LaplMatch [16] derives a permutation matrix for shape matching from Laplacian eigenvectors, without considering multiscale properties. While calculating a graph’s spectrum is computationally challenging, recent work proposes an approximation via spectral moments estimated through random walks [8].

3 Background and Problem

Graph Alignment. Consider two undirected graphs, $G_1 = (V_1, E_1)$ and $G_2 = (V_2, E_2)$, where $V_*$ are node sets, $E_* \subseteq V_* \times V_*$ are edges, and $|V_1| = |V_2| = n$. A graph’s adjacency matrix $A \in \{0, 1\}^{n \times n}$ is a binary matrix where $A_{ij} = 1$ if there is an edge between nodes $i$ and $j$ and $A_{ij} = 0$ otherwise.

Definition 1. Given two graphs $G_1 = (V_1, E_1)$ and $G_2 = (V_2, E_2)$, a graph alignment $R : V_1 \rightarrow V_2$ is an injective function that maps nodes of $G_1$ to nodes of $G_2$.

The graph alignment problem is to find such a function, which, expressed as a permutation matrix $P$, minimizes the difference $\|PA_1P^T - A_2\|^2$. In case of isomorphic graphs, there exists a $P$ such that $PA_1P^T = A_2$, i.e., aligns the two graphs exactly. We are interested in the general, unrestricted problem case, in which there are no additional constraints on node attributes or matches known in advance. The problem is hard and not known to be in \textbf{NP}.

We may express graph alignment in terms of a ground truth function $\tau : V_1 \rightarrow V_2$ that returns the correct alignment between the nodes $V_1$ in $G_1$ and the nodes $V_2$ in $G_2$. In the case of isomorphic graphs, this ground truth function $\tau$ is a bijection that admits an inverse mapping $\tau^{-1} : V_2 \rightarrow V_1$. The composition of the indicator function $\delta_i : V_1 \rightarrow \{0, 1\}$ with $\tau^{-1}$, $\delta_i \circ \tau^{-1} : V_2 \rightarrow \{0, 1\}$ expresses the

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5 Solutions to the problem of aligning graphs with unequal numbers of nodes can rest on solutions to this basic problem form.
complete isomorphism among the two graphs, returning 1 if node \( u \in V_2 \) maps to node \( i \in V_1 \), 0 otherwise. By generalization, the composition \( g_i = f_i \circ \tau^{-1} \) maps functions in \( G_2 \) to functions in \( G_1 \) for any family of real-valued functions \( f_1, \ldots, f_q, f_i : V_1 \rightarrow \mathbb{R} \) and \( g_1, \ldots, g_q, g_i : V_2 \rightarrow \mathbb{R} \) that associate a real value to each node in \( G_1 \) and \( G_2 \). This transformation among functions is called a functional representation of the mapping \( \tau \). In effect, finding an alignment among the nodes of two graphs corresponds to finding an alignment among functions on those nodes. We use such functional alignments as a shortcut to node alignments. To get there, we extend the concept of a functional map \([28]\) from the continuous to the discrete case.

**Functional maps.** The operator \( T_f : (V_1 \times \mathbb{R}) \rightarrow (V_2 \times \mathbb{R}) \) maps functions \( f \) on the nodes in \( G_1 \) to functions \( g \) on the nodes in \( G_2 \), i.e. \( T_f(f)=f \circ \tau^{-1}=g \). This operator is linear in the function space, i.e., \( T_f(c_1 f_1 + c_2 f_2) = (c_1 f_1 + c_2 f_2) \circ \tau^{-1} = c_1 f_1 \circ \tau^{-1} + c_2 f_2 \circ \tau^{-1} = c_1 T_f(f_1) + c_2 T_f(f_2) \). In addition, let \( \phi_1, \ldots, \phi_n \) and \( \psi_1, \ldots, \psi_n \) denote orthogonal bases for the space of functions on \( G_1 \)'s nodes, \( V_1 \times \mathbb{R} \), and that on \( G_2 \)'s nodes, \( V_2 \times \mathbb{R} \), respectively. Since those functions produce \( n \)-dimensional vectors, we can represent them as linear combinations of their basis vectors, \( f = \sum_{i=1}^{n} a_i \phi_i \) and \( g = \sum_{j=1}^{n} b_j \psi_j \). Then, by the linearity of \( T_f \),

\[
T_f(f) = T_f \left( \sum_{i=1}^{n} a_i \phi_i \right) = \sum_{i=1}^{n} a_i T_f(\phi_i) = \sum_{i=1}^{n} a_i \sum_{j=1}^{n} c_{ij} \psi_j = \sum_{j=1}^{n} b_j \psi_j
\]

where \( T_f(\phi_i) = \sum_{j=1}^{n} c_{ij} \psi_j \). It follows that each coefficient \( b_j \) is the dot-product \( \sum_{i=1}^{n} a_i c_{ij} \) between the coefficients \( (a_1, \ldots, a_n) \) of functions in \( G_1 \) and the coefficients \( (c_{1j}, \ldots, c_{nj}) \) of the operator \( T_f \). In conclusion, in order to align real-valued functions on the nodes of two graphs, we need to find a mapping matrix \( C \in \mathbb{R}^{n \times n} \) of coefficients among those functions; such a mapping matrix \( C \) maps functions from \( G_1 \) to \( G_2 \), even when the ground-truth mapping \( \tau \) is unknown. In a nutshell, GRASP obtains such a mapping matrix \( C \) for a well-chosen function and applies that \( C \) to mapping the indicator function \( \delta \) from \( G_1 \) to \( G_2 \), thereby constructing a node alignment. The main question we need to answer is what orthogonal basis and functions we should use to construct our mapping matrix \( C \). The next section answers this question and builds on that answer to devise a solution.

### 4 Solution

Here, we choose an orthonormal basis and a function, which are, in our judgement, appropriate for node alignment purposes, and define the complete pipeline of our solution.

#### 4.1 Choice of basis: Normalized Laplacian

As a basis for representing functions as linear combinations of base functions, we use the eigenvectors of the graph’s normalized Laplacian, i.e., the matrix \( L = I - \)
where $D$ a diagonal degree matrix of node degrees $D_{ii} = \sum_{j=1}^{n} A_{ij}$ and $A$ is the graph adjacency matrix; its eigendecomposition is $L = \Phi \Lambda \Phi^T$, where $\Lambda$ is a diagonal matrix of eigenvalues, $\{\lambda_1, \ldots, \lambda_n\}$, i.e., the graph’s spectrum, which encodes information about communities, degree distribution, and diameter, and $\Phi$ is a matrix of eigenvectors, $\Phi_L = [\phi_1 \phi_2 \ldots \phi_n]$. The eigenvectors form an orthogonal basis, which we use a standard basis. We use $\phi (\psi)$ to indicate the eigenvectors of the Laplacian of graph $G_1$ ($G_2$).

We consider this basis to be suitable, since the eigenvectors of the normalized Laplacian converge to the eigenfunctions of the Laplace-Berltrami operator [4], which measures the smoothness of continuous surfaces.

### 4.2 Choice of function: Heat Kernel

The choice of functions $f_i : V_1 \rightarrow \mathbb{R}$, $g_i : V_2 \rightarrow \mathbb{R}$, is critical for our method. A poor choice would be detrimental. A good choice should have the following properties:

**Expressiveness.** The function should express the graph’s structure. For instance, a constant function returning the same value for all nodes would not yield a meaningful alignment.

**Permutation-invariance.** The function should not depend on the node index $i$; the indicator function lacks this property.

**Multiscale robustness.** The function should robustly capture both local and global structures (e.g., edges and communities), insensitively to small perturbations.

A function fulfilling these requirements is the time-parameterized heat kernel [31]:

$$H_t = \Phi e^{-t\Lambda} \Phi^T = \sum_{j=1}^{n} e^{-t\lambda_j} \phi_j \phi_j^T$$

where $H_t[i,j]$ measures the flow of heat from node $i$ to node $j$ at time $t$, as it diffuses from each node’s neighborhood to the whole graph. We build our model functions over a sequence of time steps $t$ using the diagonal of the heat kernel, which measures the heat flowing back to each node at time $t$.

The heat kernel expresses multiscale graph structure in a permutation-invariant manner and is robust to small changes. In the beginning of the diffusion, Equation (1) emphasizes large $\lambda$, which correspond to local edge and ego-net properties. As time progresses, smaller eigenvalues get emphasized, reflecting global graph properties, such as communities.

We build our corresponding functions $f_i, g_i$, from the heat kernel at different time steps $t$, as linear combinations of the graph’s Laplacian orthogonal eigenvectors. Let $F \in \mathbb{R}^{n \times q}$, $F = [f_1, \ldots, f_q]$ be the matrix containing the diagonals of the heat kernel of $G_1$, $H_t^{G_1}$, over $q$ time steps, $f_i = \sum_{j=1}^{n} e^{-t\lambda_j} \phi_j \phi_j^T$ where $\odot$ denotes the element-wise vector product. Likewise, the matrix $G \in \mathbb{R}^{n \times q}$, $G = $
contains the diagonals of \( H_{G_2}^t \), the heat kernel of \( G_2 \). While the \( q \) columns of \( F \) and \( G \) contain the same time-dependent heat-kernel-diagonal functions on the nodes of two graphs, their \( n \) rows (i.e., nodes) are not aligned. We need to obtain such a node alignment.

**4.3 Mapping matrix**

We approximate each function \( f_i \) using only the first \( k \) eigenvectors, as done, by analogy, on shapes analysis [4], and thereby calculate the corresponding function matrices \( F \) and \( G \). \( F \) and \( G \) can be thought as coefficient matrices used to produce linear combinations, \( F^\top \Phi \) and \( G^\top \Psi \), of the Laplacian eigenvectors of \( G_1 \) and \( G_2 \), respectively. With a slight abuse of notation, we denote with \( \Phi \) and \( \Psi \) the first \( k \) eigenvectors, hence \( F^\top \Phi \) and \( G^\top \Psi \) are in \( \mathbb{R}^{q \times k} \). In the projection of the functions on the first \( k \) eigenvectors, we would like the corresponding functions to be equal up to a coefficient matrix \( C \in \mathbb{R}^{k \times k} \). In the case of isomorphic graphs, it holds that

\[
\begin{bmatrix}
\text{diag}(g_1^\top \Psi) \\
\vdots \\
\text{diag}(g_q^\top \Psi)
\end{bmatrix}
\begin{bmatrix}
c_{11} \\
\vdots \\
c_{kk}
\end{bmatrix}
= 
\begin{bmatrix}
\Phi^\top f_1 \\
\vdots \\
\Phi^\top f_q
\end{bmatrix}
\tag{2}
\]

Matrix \( C \) is diagonal in the case of isomorphic graphs and deviates from a diagonal form as graphs diverge from isomorphism; for simplicity, we assume a diagonal \( C \), and obtain the diagonal entries that minimize the \( L_2 \)-norm difference \( \| \cdot \|_2 \) between the left and right side of Eq. (2) using the ordinary least squares method, as in [21]. In Section 4.5 we delve into the case of non-isomorphic graphs.

**4.4 Node-to-node correspondence**

We consider the delta function \( \delta_i(\cdot) \) as corresponding function; these functions yield an \( n \times n \) identity matrix. We express such a function as a vector of coefficients, since the vector of \( \delta_i \) is the \( i \)th row of the heat kernel at \( t = 0 \):

\[
\delta_i = H_{G_1}^{t=0} = \sum_{j=1}^{n} \phi_{ij} \phi_j
\]

The computation for delta functions on \( G_2 \) follows equivalently using \( \Psi \) in place of \( \Phi \). We may match the coefficient vectors of these corresponding indicator functions, as, ideally, for two matching nodes \( v_i \in V_1 \) and \( v'_j \in V_2 \), the coefficients of \( \delta_i \) and \( \delta_j \) for \( \Phi \) and \( \Psi \) should be identical. In particular, the coefficients expressing \( \delta_i \) as a linear combination of the first \( k \) eigenvectors are \( \phi_{i1}, \ldots, \phi_{ik} \). We set \( \Phi^\top \) and \( C \Psi^\top \) in \( \mathbb{R}^{k \times n} \) as coefficient matrices of the delta functions, aligned by \( C \). Rows correspond to the first \( k \) Laplacian eigenvectors, while columns stand for graph nodes, rather than for time steps of heat diffusion. We need to match coefficient vectors, i.e., columns of \( \Phi^\top \) and \( C \Psi^\top \), to each other. This problem amounts to a linear assignment problem; we apply an
off-the-shelf algorithm therefore, such as nearest neighbor search or Jonker-Volgenant (JV) [13], to obtain a one-to-one matching between the columns of $\Phi^\top$ and $C\Psi^\top$, and hence an alignment of nodes. GRASP is flexible in that we may choose any matching method.

### 4.5 Base Alignment

We have hitherto assumed that the graphs to be aligned are isomorphic, hence their eigenvectors correspond to each other with possible sign changes and an orthogonal diagonal mapping matrix $C$ exists. Still, if the graphs are not isomorphic, then their eigenvectors diverge and the diagonal matrix $C$, which we enforce, cannot capture their relationship well. Figure 2 highlights this issue: at a high level the eigenvectors underline common structures, but they differ at the node level. In this case, we need to align the two eigenvector bases before we consider aligning corresponding vectors and, eventually, nodes. We express this base alignment [21] in terms of an alignment matrix $M$.

**Diagonalization.** We align the eigenvectors $\Psi$ by a rotation matrix $M$ so as transform $\Psi$ into $\Phi$: $\hat{\Psi} = \Psi M$. Since $L\Psi = \Psi A$, finding $\Psi$ is equivalent to the solution of the quadratic minimization problem $\min_{\Psi} \text{off}(\Psi^\top L_2 \Psi)$ s.t. $\Psi^\top \Psi = I$ which penalizes elements off($\cdot$) outside of the diagonal, in order to preserve orthogonality of the basis.

Since the eigenvectors are orthonormal, $\Psi^\top \Psi = I$ and for $G_2$’s graph Laplacian eigenvectors $A_2$, $\Psi^\top L_2 \Psi = \Psi^\top \Psi A_2 = A_2$, and $M^\top \Psi^\top L_2 \Psi M = M^\top A_2 M$. Putting the above together, our diagonalizing term is:

$$\min_{M} \text{off}(M^\top A_2 M) \text{ s.t. } M^\top M = I$$

As we are minimizing over orthogonal matrices we can equivalently express the objective above as a minimization over orthogonal matrices of size $n \times n$, $S(n,n)$:

$$\min_{M \in S(n,n)} \text{off}(M^\top A_2 M)$$

**Coupling.** In addition, the correspondence $\tau : G_1 \to G_2$ so that $\phi_i \approx \tau \circ \psi$ translates to
\[
\min_{\Phi} \| F^T \Phi - G^T \Psi M \|_F^2
\]

where \( F \) and \( G \) contain each graphs’s corresponding functions. We combine the minimization terms for diagonalization and coupling, to get the following minimization problem, with regularization factor \( \mu^7 \):

\[
\min_{M \in S(n,n)} \text{off}(M^T A_2 M) + \mu \| F^T \Phi - G^T \Psi M \|_F^2.
\]  

Eq. (3) leads to a manifold optimization problem, which we solve by trust-region methods [2].

**Scalability.** We avoid computing all eigenvectors \( n \times n \), exploiting the fact that we only need the first \( k \) eigenvectors for calculating \( C \) (see Section 4.3). So we only align the first \( k \) eigenvectors of \( \Psi \) to the first \( k \) eigenvectors of \( \Phi \), i.e \( \tilde{\Phi} = \Psi M \) with \( \tilde{\Phi} = [\phi_1, \ldots, \phi_k] \) and \( \tilde{\Psi} = [\psi_1, \ldots, \psi_k] \). Let \( A_2 = \text{diag}(\lambda_1, \ldots, \lambda_k) \), the problem in Eq. (3) becomes

\[
\min_{M \in S(k,k)} \text{off}(M^T A_2 M) + \mu \| F^T \tilde{\Phi} - G^T \tilde{\Psi} M \|_F^2.
\]  

After obtaining \( M \), we use the eigenvectors in \( \tilde{\Phi} \) and the aligned eigenvectors \( \tilde{\Psi} = \Psi M \) in the next step for the final alignment of nodes. Our approach effectively trades off graph alignment with a proxy problem of manifold optimization, which we solve with reasonable accuracy and scalability.

### 4.6 Our algorithm: GRASP

Putting it all together, GRASP consists of five steps; the pseudocode is given in the supplementary material.

**Steps 1:** Compute eigenvectors. In the first step, calculate the Laplacians \( L_1, L_2 \) of the two graphs \( G_1 \) and \( G_2 \). Then compute the eigenvectors \( \Phi, \Psi \) and eigenvalues \( \lambda_1, \lambda_2 \) by the eigendecomposition \( L_1 = \Phi \Lambda_1 \Phi^T \) and \( L_2 = \Psi \Lambda_2 \Psi^T \).

**Step 2:** Compute corresponding functions. In the second step, calculate the matrices of corresponding functions \( F = [f_1, \ldots, f_q] \) and \( G = [g_1, \ldots, g_q] \) as diagonals of the heat kernel at time steps \( [t_1, \ldots, t_q] \) with \( f_i = \sum_{j=1}^n e^{-t_i \lambda_j} \phi_j \odot \phi_j \) and \( g_i \), equivalently using \( \Psi \).

**Step 3:** Base alignment. After the corresponding functions are calculated, obtain the base alignment matrix \( M \) by minimizing Eq. 3. Then align the first \( k \) columns of \( \Psi \), denoted by \( \tilde{\Psi} \) to the corresponding first \( k \) columns \( \tilde{\Phi} \) of \( \Phi \) as \( \tilde{\Psi} = \Psi M \).

\( ^7 \mu = 0.132 \) in our experiments.
Step 4: Calculate mapping matrix. Under the assumption that $C$ is a diagonal matrix, calculate its diagonal elements $c_{11}, \ldots, c_{kk}$ by solving the least squares problem:

\[
\min_{[c_{11}, \ldots, c_{kk}]^T} \left\| \begin{bmatrix} \text{diag}(g_1^T \tilde{\Psi}) & \vdots & \text{diag}(g_q^T \tilde{\Psi}) \end{bmatrix} \begin{bmatrix} c_{11} \\ \vdots \\ c_{kk} \end{bmatrix} - \begin{bmatrix} \tilde{\Phi}^T f_1 \\ \vdots \\ \tilde{\Phi}^T f_q \end{bmatrix} \right\|_2^2
\]  

(5)

We then set $C = \text{diag}(c_{11}, \ldots, c_{kk})$.

Step 5: Node alignment. To get the final node alignment, we apply a linear assignment algorithm on the rows of $\tilde{\Phi}$ and $C^T \tilde{\Psi}$, which hold the indicator function coefficients.

**Complexity analysis** The computation of the first $k$ Laplacian eigenvectors takes $O(k \max\{|E_1|, |E_2|\})$ by fast methods for diagonally dominant matrices [19]. Base alignment needs $O(k^3)$ to solve the orthogonality constraint through trust-region methods. The least-squares method runs in $O(qk)$. The matching step by JV runs in $O(n^3)$. Overall, the $O(n^3)$ time factor is dominant.

**Connection to Differential Geometry** Our work rests on the theory on Riemannian manifolds [11] and builds on the analogy between a graph’s Laplacian and the continuous Laplace-Beltrami operator [31].

5 Experiments

We experiment on three different real-world networks; Table 1 in the Supplementary Material lists their properties. As in [12], we randomly permute the node order and inject noise by randomly deleting edges with probability $p$. We generate a graph for $p = 0.01$, which we align to 5 noisy graphs, one for each $p$ value, ranging from 0.05 to 0.25; we measure alignment accuracy as the average ratio of correctly aligned nodes; note that none of the noisy graphs in a pair is a subset of the other.

**Baselines.** We compare against the following established state-of-the art baselines for unrestricted graph alignment.

- **REGAL** [12]: A method based on embeddings utilizing local structural features. REGAL allows one-to-many matchings. For the sake of fairness, we let REGAL provide one-to-one matchings using the JV linear assignment algorithm, as GRASP does; we confirmed that, doing so, it fares better than using nearest neighbors.

- **Low Rank EigenAlign (LREA)** [27]: A spectral method that yields one-to-one matchings via the minimization of edge mismatches.

We eschew a comparison with IsoRank [30, 22] and other methods for the alignment of biological networks [9], since REGAL and LREA significantly outperform those methods.
Parameter tuning. After experimenting with different settings, we settled for using $k = 20$ eigenvalues and $q = 100$ corresponding functions, which yield best accuracy. We report the tuning experiments in the supplementary material.

![Graph showing accuracy comparison](image)

**Fig. 3:** Accuracy of nearest neighbor and JV matching algorithms.

Justifying algorithmic choices We evaluate the impact of (i) the choice of algorithm for node-to-node assignment (Section 4.4) and (ii) base alignment (Section 4.5). Figure 3 shows that both the JV linear assignment algorithm and base alignment bring a substantial advantage over their rudimentary counterparts, consistently across datasets. In the following experiments, unless otherwise stated, we settle on the variant of GRASP equipped with base alignment.

![Accuracy comparison graph](image)

**Fig. 4:** Accuracy compared to REGAL and LREA

Comparison to previous methods Figure 4 shows that GRASP outperforms others by a large margin, achieving 62% accuracy in Arenas and 43% in Facebook with 5% noise, and fares at comparably well as REGAL on the CA-AstroPH graph.
Real world networks  We also try matching among real world networks. Multi-Magna is a collection of graphs consisting of a base yeast network and five variations thereof. We match these five variations to the original; Figure 5 presents our results. HighSchool and Voles are two evolving proximity networks. We match their latest version to versions at time steps with 80%, 85%, 90%, and 99% of all edges. Figure 5 shows the results. The advantage of GRASP observed with synthetic noise transfers to real-world alignment problems.

Efficiency  Steps 1–3 of GRASP (Section 4.6) can be performed offline, while REGAL also allows for precomputation of representations. Figure 6 (left) shows the time to compute the alignments after precomputation. GRASP outperforms REGAL and LREA in the largest CA-AstroPh data. Figure 6 (right) shows the time with online precomputation; REGAL does not exhibit any substantial advantage even in the smaller Arenas and Facebook graphs, while GRASP attains more accurate results with a negligible increase in time. We obtained similar results on real-world network matching tasks.
6 Conclusion

We proposed a graph alignment method using their Laplacian eigenvectors. We establish a functional correspondence among pre-aligned eigenvectors, capturing multiscale graph properties and extract a linear assignment among matrix columns, attaining superior alignment quality over the state of the art. To our knowledge, this is the first work to apply a functional alignment primitive to graph alignment. In the future, we plan to extend our method to partial correspondences and examine to what extent our representations can be employed within the framework of [5].

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