Graph Matching with Adaptive and Branching Path Following

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Abstract—Graph matching aims at establishing correspondences between graph elements, and is widely used in many computer vision tasks. Among recently proposed graph matching algorithms, those utilizing the path following strategy have attracted special research attentions due to their exhibition of state-of-the-art performances. However, the paths computed in these algorithms often contain singular points, which could hurt the matching performance if not dealt properly. To deal with this issue, we propose a novel path following strategy, named branching path following (BPF), to improve graph matching accuracy. In particular, we first propose a singular point detector by solving a KKT system, and then design a branch switching method to seek for better paths at singular points. Moreover, to reduce the computational burden of the BPF strategy, an adaptive path estimation (APE) strategy is integrated into BPF to accelerate the convergence of searching along each path. A new graph matching algorithm named ABPF-G is developed by applying APE and BPF to a recently proposed path following algorithm named GNCCP (Liu & Qiao 2014). Experimental results reveal how our approach consistently outperforms state-of-the-art algorithms for graph matching on five public benchmark datasets.

Index Terms—Graph Matching, Path Following, Singular Point, Branch Switching, Adaptive Path Estimation.

1 INTRODUCTION

Given two graphs, graph matching involves establishing correspondences between their vertex sets while taking into consideration the consistency between their edge sets as well. It is a fundamental problem in computer science and closely relates to many computer vision problems including feature registration [46], [49], [50], shape matching [5], [41], [53], object recognition [19], [55], visual tracking [8], [52], activity analysis [9], etc. Despite decades of research effort devoted to graph matching, it remains challenging to acquire a global optimum solution in acceptable computational time due to the combinatorial nature over the solutions. A typical way is to utilize relaxation to harness the solution searching, aiming to find approximate optimization methods at the cost of acceptable sub-optimal solutions. Popular approaches include, but not limited to, bipartite matching [43], [47], spectral relaxation [12], [14], [32], continuous optimization [24], [36], [37], [57], [60] and probabilistic modeling [20], [58], to name just a few.

Among recently proposed graph matching algorithms, the ones utilizing the path following strategy have attracted much research attention due to their exhibition of state-of-the-art performances [36], [37], [57], [60]. These algorithms reformulate graph matching as a convex-concave relaxation procedure (CCRP) problem, which is solved iteratively by interpolating between two relaxed formulations that are simpler than the original one. The solutions of all iterations form a path that recast smoothly the bistochastic matrix solution in the discrete domain. The path following algorithms can be viewed as special cases of the numerical continuation method (NCM) [3], which computes approximate solutions of parameterized nonlinear equation systems. These algorithms succeed at regular points but may fail at singular points (details in Sec. 4). It therefore demands research attention on how to address this issue to improve matching performance.

Motivated by above discussion, we propose a novel path following strategy, named branching path following (BPF), to improve path following graph matching algorithms. Roughly speaking, BPF extends the traditional path following strategy by branching new paths at singular points, with three key components. First, it discovers singular points on the original path by analyzing the Jacobian of the associated KKT system. Second, it branches a new path at each singular point using the pseudo-arclength continuation method [17], [28]. Third, an adaptive path estimation (APE) algorithm is devised that dynamically estimate the iteration steps when exploring each path, and brings further improvement in efficiency. Finally, after searching along these branching paths, BPF chooses the best one in terms of the objective function as the final solution. Note that the original path, which leads to the original path following solution, is always exploited the same as other paths in BPF. As a result, BPF is guaranteed to achieve solutions that are better than, or occasionally as good as, the solutions by the original path following strategy.

Using the proposed strategy, we develop a new graph matching algorithm, named ABPF-G, by applying APE and BPF to the GNCCP (graduated nonconvexity and concavity procedure) algorithm [36]. Note that GNCCP is chosen since it is one of the latest path following algorithms, while APE and BPF are by no means limited to working with GNCCP. For a thorough evaluation, we test the proposed ABPF-G algorithm on four popular benchmarks and a synthetic dataset. Experimental results show that, the proposed algorithm significantly improves the path following procedure and outperforms state-of-the-art graph matching algorithms in comparison. Regarding computational cost, ABPF-G achieves similar or even better computational efficiency than GNCCP.

In summary, with the proposed path following strategy for graph matching, this paper makes contribution in four-fold: (1)
we discuss the pitfalls of path following algorithms at singular points, and propose an efficient singular point discovery method; (2) we design a novel branching path following strategy to explore potentially better paths at detected singular points and thus improve matching performance; (3) we devise an adaptive path estimation method for efficiency improvement; and (4) we develop a new graph matching algorithm by applying the proposed strategies to the GNCCP algorithm, and demonstrate the effectiveness of the algorithm in a thorough evaluation. The source code of the proposed algorithm is made available for research usage (http://www.dabi.temple.edu/~hbling/code/BPF/bpf-2017.htm).

In the rest of this paper, Sec. 2 summarizes related work; then Sec. 3 reviews path following algorithms and Sec. 4 discusses the numerical continuation interpretation; after that, Sec. 5 introduces the proposed APE strategy, and Sec. 6 presents the BPF strategy that employs APE as a component, and further applies them to GNCCP to drive a new algorithm. We finally present experimental validation in Sec. 7 and conclusion in Sec. 8.

2 Related work

Graph matching has been investigated for decades and many algorithms have been proposed. In general, its combinatorial nature makes the global optimum solution hardly available. As a result, approximate solutions are commonly applied to graph matching. In this section we review some sampled classical studies or those closely relate to ours, and leave general graph matching research to two comprehensive surveys [13], [22].

2.1 Classical and recent studies in graph matching

The graph edit distance (GED) [45] is one of the most flexible and versatile graph distance metric that arises naturally in the context of inexact graph matching. The idea is to define graph edit operations and use the amount of operations between two graphs to measure their similarity. The exponential complexity, however, restricts the applicability of the edit distance to small graphs. To reduce the complexity, extended algorithms either find approximate solutions to the problem with acceptable sub-optimal solutions [27], or find efficient algorithms for restricted classes of graphs such as ordered graph [26]. Instead of acquiring global consistency between graphs, bipartite graph matching (BGM) considers only local edge structure during the optimization process [43], and convert the difficult GED problem to a linear assignment problem by encoding the local edge structure into individual nodes.

Aiming to generate a solution obeying the discrete constraints, some recent studies [2], [25], [44] employ an $\ell_1$-norm constraint in the nonnegative domain to induce a sparse (approximate discrete) solution to the matching problem. However, the sparsity of solution acquired by these algorithms is usually uncontrollable, and a post discretization step is also usually required to guarantee the desired discrete solution. Recently, some pure discrete methods [1], [56] are proposed to search the solution directly in the discrete space. These methods can generate a solution obeying the discrete affine mapping constraints, and thus does not require any post-optimization step. However, the optimality of the discrete search method is usually limited which is dependent largely on the initialization [1]. Other sampled recent studies also include [59] that solves the matching between attributed relational graphs (ARGs) via mining a hierarchical and-or graph (AoG) model, and [40] that analyzes in depth the relaxation in graph matching.

2.2 Path following and related algorithms

Since graph matching is inherently a discrete optimization problem, a typical strategy is to relax it to continuous domain for an initial solution, and then project it into the discrete domain in a graduated and iterative process in which the objective function is employed. Many important algorithms have been designed on top of the relaxation and discretization strategy. For example, Gold and Rangarajan [24] employ the graduated assignment technique to iteratively solve a series of linear approximations of the cost function using Taylor expansion. Leordeanu and Hebert [33] propose an integer projected fixed-point algorithm to optimize the objective function in the integer domain, which takes any continuous or discrete solution, possibly given by some other graph matching method, as input. Subsequently, Lu et al. [39] develop a more efficient projected fixed-point algorithm by utilizing a doubly stochastic projection method.
Along the line there comes the group of path following algorithms. In particular, Zaslavskiy et al. [57] reformulate graph matching as a convex-concave relaxation procedure (CCR) problem and then solve it by interpolating between two simpler relaxed formulations, convex relaxation and concave relaxation. More specifically, the path following algorithm proposed by them iteratively searches a solution by tracing a path of local minima of a series of functions that linearly interpolate between the two relaxations. Following the similar strategy, Zhou and De la Torre [60], [61] factorize an affinity matrix into a Kronecker product of smaller matrices that encodes the structure of the graphs and the affinities between nodes and between edges, and provide specified convex relaxation and concave relaxation based on the factorized matrices. Later, Liu and Qiao [36] propose the graduated nonconvexity and concavity procedure (GNCCP), and prove that it equivalently realizes CCRP on partial permutation matrix. GNCCP provides a much simpler way for CCR without explicitly involving the convex or concave relaxation.

Our work falls into the group of path following algorithms. Compared with previous arts, our method focuses on improving the path following strategy itself, which has not been fully explored before. Preliminary versions of some parts of this work has been published in two conference papers [51], [54], with [54] focusing on the idea of branching path and [51] on adaptive steps. The studies in the two papers are roughly independent, while in this work, we combined their ideas for a further improved graph matching algorithm. As shown in our experiments, such combination benefits not only the computational efficiency but also the matching performance. Details can be found in Sec. 7.

3 PATH FOLLOWING FOR GRAPH MATCHING

3.1 Problem formulation

An undirected graph of \( n \) nodes can be represented by \( G = (V, E) \), where \( V = \{v_1, \ldots, v_n\} \) and \( E \subseteq V \times V \) denote the node set and the edge set, respectively. A graph is often conveniently represented by a symmetric adjacency matrix \( A \in \mathbb{R}^{n \times n} \), such that \( A_{ij} = 1 \) if and only if there is an edge between \( v_i \) and \( v_j \). The adjacency matrix is usually generalized to a weighted graph by associating a nonnegative real-valued weight \( A_{ij} = w_{ij} \) to all pair of nodes. This generalization is important for many applications to capture structural relations among nodes. In the rest of this paper, all adjacency matrices mentioned are weighted with real values unless otherwise stated.

For graph matching, given two graphs \( G^{(i)} = (V^{(i)}, E^{(i)}) \) of size \( n_i, i = 1, 2 \), we assume \( n_1 \leq n_2 \) without loss of generality. The problem is to find a node correspondence \( X \in \{0, 1\}^{n_1 \times n_2} \) between \( G^{(1)} \) and \( G^{(2)} \) in favor of the following global consistency:

\[
E_1(X) = \sum_{i_1, i_2} c_{i_1 i_2} X_{i_1 i_2} + \sum_{i_1, j_1, j_2} d_{i_1 j_1 i_2 j_2} X_{i_1 i_2} X_{j_1 j_2}, \quad (1)
\]

where \( c_{i_1 i_2} \) measures the consistency between the \( i_1 \)-th node in \( G^{(1)} \) and the \( i_2 \)-th node in \( G^{(2)} \), and \( d_{i_1 j_1 i_2 j_2} \) the the consistency between edge \((i_1, j_1)\) in \( G^{(1)} \) and edge \((i_2, j_2)\) in \( G^{(2)} \). The corresponding matrix \( X \) denotes matching result, i.e., \( X_{i_1 i_2} = 1 \) if and only if \( v_{i_1} \in V^{(1)} \) corresponds to \( v_{i_2} \in V^{(2)} \). In practice, the matching is often restricted to be one-to-one function, which requires \( X^T 1_{n_2} = 1_{n_1} \) and \( X^T 1_{n_1} \leq 1_{n_2} \), where \( 1_n \) denotes a vector of \( n \) ones and \( < \) the element-wise \( \leq \).

Let \( A^{(i)} \) be the adjacency matrix for \( G^{(i)}, i = 1, 2 \), a more commonly used formulation for weighted graph matching [4], [57] is defined as

\[
E_2(X) = \text{tr}(C^T X) + \alpha \| A^{(1)} - X A^{(2)} X^T \|_F^2, \quad (2)
\]

where \( C = (c_{i_1 i_2}) \in \mathbb{R}^{n_1 \times n_2} \) is the node dissimilarity matrix, \( \alpha \geq 0 \) the weight balancing between the node and edge comparisons, and \( \| \cdot \|_F \) the Frobenius norm.

Eq. (2) for weighted graph matching is limited in practice because only a scalar attribute is associated to each node and the edge consistency function is restricted to the difference of edge weights. In recent studies [12], [20], [32], [36], [61], a more general formulation of Eq. (1) is formulated in a pairwise compatibility form

\[
E_3(x) = x^T K x, \quad (3)
\]

where \( x = \text{vec}(X) \in \{0, 1\}^{n_1 n_2} \) is the vectorized version of matrix \( X \) and \( K \in \mathbb{R}^{n_1 n_2 \times n_1 n_2} \) is the corresponding affinity matrix defined as:

\[
K_{\text{ind}(i_1, i_2)\text{ind}(j_1, j_2)} = \begin{cases} c_{i_1 i_2} & \text{if } i_1 = j_1 \text{ and } i_2 = j_2, \\ d_{i_1 j_1 i_2 j_2} & \text{if } A^{(1)}_{i_1 j_1} A^{(2)}_{i_2 j_2} > 0, \\ 0 & \text{otherwise}, \end{cases} \quad (4)
\]

where \( \text{ind}(\cdot, \cdot) \) is a bijection mapping a node correspondence to an integer index.

In this paper, we mainly study graph matching algorithms for Eq. (3), since it can encode not only the difference of edge weights but also many complex graph compatibility functions.

3.2 The path following algorithm

In [57], Zaslavskiy et al. introduce the convex-concave relaxation procedure (CCR) into the graph matching problem defined in Eq. (2) by reformulating it as interpolation between two relaxed problems that are simpler than the original one. The first relaxation is obtained by expanding the convex quadratic function \( E_2(X) \) from the set of permutation matrices \( P \) to the set of doubly stochastic matrices \( D \):

\[
\min_{X \in D} E_2(X) = \text{tr}(C^T X) + \alpha \| A^{(1)} - X A^{(2)} X^T \|_F^2. \quad (5)
\]

The second relaxation is a concave function

\[
\min_{X \in D} E_4(X) = - \text{tr}(\Delta X) - 2 \text{vec}(X)^T ((L^{(1)})^T \otimes (L^{(2)})^T) \text{vec}(X), \quad (6)
\]

where \( \Delta \) is a matrix with element \( \Delta_{ij} = (D_{i_1 j_1}^{(1)} - D_{i_2 j_2}^{(2)})^2 \); \( D^{(i)} \) and \( L^{(i)} \) represent respectively the diagonal degree matrix and the Laplacian matrix of an adjacency matrix \( A^{(i)}, i = 1, 2 \); and \( \otimes \) denotes the Kronecker product. A key property of formulation (6) is that, its optimum solution over \( D \) leads to the optimization over \( P \); more importantly, its optimum solution over \( D \) is exactly the solution of the original graph matching problem.

Thus inspired, the path following strategy proposed in [57] can be interpreted as an iterative procedure that smoothly projects an initial solution of \( E_2 \) in the continuous space \( D \) to the discrete space \( P \) by tracking a path of local minima of a series of functionals \( E_\lambda \) over \( D \):

\[
E_\lambda = (1 - \lambda) E_2 + \lambda E_4, \quad (7)
\]

for \( 0 \leq \lambda \leq 1 \). Each local minimum of \( E_\lambda + \delta_\lambda \) is gained by the Frank-Wolfe algorithm [23] given the local minimum of \( E_\lambda \) as the
start point. The convex relaxation guarantees the path starts from the unique local minimum of $\mathcal{E}_2$ for $\lambda = 0$. Increasing $\lambda$ from 0 to 1, this approach searches toward a local minimum of $\mathcal{E}_1$, and takes it as the final solution. The concave property of $\mathcal{E}_4(X)$ guarantees that the final solution lies in the discrete space $\mathcal{P}$. More details about the path following algorithm can be found in [57].

4 Numerical continuation interpretation

In this section, we interpret the path following algorithms in a numerical continuation view, and then discuss their pitfalls due to singular points. This guides the subsequent improvement on these algorithms. For simplicity, we assume the two graphs to be matched have the same size $n$, and formulations can be easily extended to handle varied sizes, e.g., by adding dummy nodes.

4.1 KKT system

According to the path following strategy described above and by converting $\mathcal{D}$ into constraints, we need to solve a series of optimization problems with equality and inequality constraints parameterized by $\lambda$:

$$\begin{align*}
\mathbf{x}^* &= \arg\max_x \mathcal{E}_4(x), \\
&\text{s.t. } \begin{cases}
Bx = 1_{2n}, \\
x \succeq 0_{n^2}.
\end{cases}
\end{align*}$$

(8)

where $Bx = 1_{2n}$ encodes the one-to-one matching constraints, and $\succeq$ is the element-wise $\geq$. Specifically, $B \in \{0, 1\}^{2n \times n^2}$ can be decomposed into two components $B = [B^{(1)}; B^{(2)}]$, where $B^{(1)}, B^{(2)} \in \{0, 1\}^{n \times n^2}$ encode the one-to-one matching constraints on nodes in graphs $\mathcal{G}^{(1)}$ and $\mathcal{G}^{(2)}$ respectively. In particular, $B^{(1)}$ and $B^{(2)}$ are defined as

$$
B_{1k}^{(1)} = \begin{cases}
1, & \text{if } \exists j \text{ s.t. } k = \text{ind}(i, j), \\
0, & \text{otherwise},
\end{cases}
$$

(9)

$$
B_{2k}^{(2)} = \begin{cases}
1, & \text{if } \exists j \text{ s.t. } k = \text{ind}(j, i), \\
0, & \text{otherwise}.
\end{cases}
$$

(10)

where $\text{ind}(\cdot, \cdot)$ is the same bijection mapping as in Eq. (4).

Using Lagrange multipliers $\alpha_i$, $i = 1, 2, \ldots, 2n$, and KKT multipliers $\mu_j$, $j = 1, 2, \ldots, n^2$, the above constrained problem can be converted to the following unconstrained one

$$\begin{align*}
\mathbf{x}^* &= \arg\max_{\mathbf{x}} \left( \mathcal{E}_4(\mathbf{x}) + \sum_{i=1}^{2n} \alpha_i h_i(\mathbf{x}) - \sum_{j=1}^{n^2} \mu_j g_j(\mathbf{x}) \right),
\end{align*}$$

(11)

where

$$h_i(\mathbf{x}) = B(i, :)\mathbf{x} - 1, \quad \text{and } g_j(\mathbf{x}) = -x_j.$$  

(12)

This results in the following system of Karush-Kuhn-Tucker (KKT) equations [30]:

$$
\begin{align*}
\nabla \mathcal{E}_4(\mathbf{x}) + B^\top \alpha + \mu &= 0_n, \\
h_i(\mathbf{x}) &= 0, \quad 1 \leq i \leq 2n, \\
\mu_j g_j(\mathbf{x}) &= 0, \quad 1 \leq j \leq n^2, \\
\mu_j &\geq 0, \quad 1 \leq j \leq n^2,
\end{align*}
$$

(13)

where $\alpha = (\alpha_1, \alpha_2, \ldots, \alpha_{2n})^\top \in \mathbb{R}^{2n}$ and $\mu = (\mu_1, \mu_2, \ldots, \mu_{n^2})^\top \in \mathbb{R}^{n^2}$ are the multiplier vectors.

4.2 Numerical continuation method

The KKT system in Eq. (13) can be reformulated as a constrained nonlinear system

$$F(\lambda, \mathbf{x}, \alpha, \mu) = 0_m, \quad \text{s.t. } \mu \succeq 0_{n^2}$$

(14)

where $m = 2n^2 + 2n$ is the number of equations in the KKT system. We further introduce vector notation $\mathbf{u} = (u_1, u_2, \ldots, u_m)^\top \in \mathbb{R}^m$ such that $\mathbf{u} = [x^\top, \alpha^\top, \mu^\top]^\top$, then we can have a parameterized nonlinear equations system in the following implicit form:

$$F(\lambda, \mathbf{u}) = 0_m,$$

(15)

such that $F$ is a mapping: $\mathbb{R}^{m+1} \rightarrow \mathbb{R}^m$. The parameter $\lambda$ is usually a real scalar, and the solution an $m$-dimensional vector, i.e., $\mathbf{u}$. For a fixed $\lambda$, $F(\cdot, \lambda)$ maps $m$-dimensional Euclidean space into itself. Depending on applications, some or all of the variables may be subjected to inequality constraints, e.g., nonnegative constraints in Eq. (14). The feasible set of inequality constraints is usually defined as $\Omega = \{\mathbf{u} \in \mathbb{R}^m | f(\mathbf{u}) \leq 0_m\}$. In path following it can be assumed that the problem is well-defined, i.e., $F(\lambda, \mathbf{u})$ has at least one solution over $\Omega$.

Most solutions of nonlinear equation systems are iterative methods. The existing path following algorithms (PATH [57], FGM [60], GNCCP [36]) can be viewed as special cases of the numerical continuation method (NCM) [3]. In general, NCM is a method to compute approximate solutions of a system of parameterized nonlinear equations, by discretizing iteratively trajectories of the flow of $F = 0$. This method works as well in the presence of constraints on some or all of the variables [15], [48].

In fact, the PATH, FGM and GNCCP algorithms correspond to a particular implementation of the so-called generic predictor corrector (GPC) approach [3]. GPC is a simple adaptation of the iterative solver to a parameterized problem. As shown in Fig. 1(a), an initial guess $\lambda_0$ ($\lambda_0 = 0$ for path following) is acquired to achieve an initial solution $\mathbf{u}_0$, then GPC traces out a solution path by repeatedly incrementing $\lambda$ until the desired value of $\lambda$ is reached ($\lambda = 1$ for path following). The solution at a specific $\lambda$ is used as the initial guess for the solution at $\lambda + \delta\lambda$, where $\delta\lambda$ is the step length for each iteration. With $\delta\lambda$ sufficiently small the iteration applied to the initial guess should converge [3].

4.3 Pitfalls at singular points

An advantage of the GPC approach is that it takes the solution of the original problem as a black box, i.e., all it requires is an initial solution. Simple and intuitive, GPC has been demonstrated to be a reasonable method for solving nonlinear equation systems. This method, however, may fail at points that violate the assumptions of the implicit function theorem [17]. Such a point of potential failure is called a singular point when the Jacobian of $F$ (with respect to $\mathbf{u}$) at the point is rank deficient. By contrast, a regular point of $F$ is associated with a Jacobian of full rank. Next, we give an intuitive explanation of why and how GPC fails at singular points, and detailed discussion can be found in [17].

A solution component $\Gamma(\lambda_0, \mathbf{u}_0)$ of the nonlinear system $F$ is a set of points $(\lambda, \mathbf{u})$ such that $F(\lambda, \mathbf{u}) = 0$ and these points are connected to the initial solution $(\lambda_0, \mathbf{u}_0)$ by paths of solutions. As discussed in [17], near a regular point the solution component $\Gamma$ is an isolated curve passing through the regular point. By contrast, for a singular point, there may be multiple curves passing through it. The local structure of a point in $\Gamma$ is determined by high-order derivatives of $F$. In general, a solution component $\Gamma(\lambda_0, \mathbf{u}_0)$ is a connected component that may contain multiple paths, and these paths intersect at singular points. It is theoretically possible to search all paths in $\Gamma(\lambda_0, \mathbf{u}_0)$ from the given initial point $(\lambda_0, \mathbf{u}_0)$. However, GPC searches along only one path and ignores all
4.4 Computational inefficiency of GPC

In addition to the pitfall at singular points, GPC suffers from high computational burden for two reasons. First, as illustrated in Fig. 1(a), the predictor of GPC is very rough that directly uses the solution at a specific $\lambda$ as the initial guess for the solution at $\lambda + \delta_\lambda$. The guess is often far away from the actual solution, so the corrector needs to spend much time to search the local optimum at each iteration. Second, the step length $\delta_\lambda$ is fixed and always set to be very small to guarantee reasonable solutions. This naive strategy often leads to unnecessarily many iterations for convergence at desired $\lambda$. It therefore demands research attention on how to reduce the computational burden of this approach.

Inspired by the discussion above, we develop a novel adaptive path estimation (APE) approach for the parameterized nonlinear equation problem. This approach utilizes a gradient predictor to estimate the solution at each iteration using previous solutions, and dynamically adjust the step length to further accelerate the convergence. We integrate it to the BPF method for graph matching aiming to improve the computational efficiency, and the excellent experimental results validate its effectiveness.

5 Adaptive Path Estimation

In this section we address the efficiency problem in GPC, as pointed out in Sec. 4.4. At each iteration, we develop a gradient predictor to estimate the solution using previous solutions, instead of using directly last solution, and then dynamically adjust the step length to further accelerate the convergence. Since we are only interested in solution $x$ and it is independent of Lagrange multipliers $\alpha_i$ and KKT multipliers $\mu_i$ in path estimation and step length adjustment, we quote $x$, rather than $u$, for description simplicity.

5.1 Path prediction

From the implicit function theorem [29], the solutions of the parameterized nonlinear equation system $F(\lambda, x, \alpha, \mu) = 0$ form a smooth path parameterized by $\lambda$. We thus are able to predict the $t$-th solution using previous solutions. In particular, we propose to predict current solution using the differences of the previous solutions.

Denote $x_i$ the solution at a previous iteration $i$ ($i < t$), the solution difference is computed as $d_i = x_i - x_{i-1}$. We predict the difference $d_t$ and the solution $x_t'$ at iteration $t$ as

$$d_t = \frac{\sum_{i=t-k}^{t-1} w_i d_i}{\sum_{i=t-k}^{t-1} w_i},$$

$$x_t' = x_{t-1} + d_t,$$

where $k$ controls the size of the window for approximation, and $w_i$ the weight of the differences at previous solutions. We prefer Gaussian weights as $w_i = \exp(-(t-i)^2/\gamma)$ and set $\gamma = 10k$ throughout our experiments. The motivation behind this prediction is the smoothness of the path, which implies the similarity between solution differences of consecutive iterations.

Once the prediction $x_t'$ is ready, it is used as the start point to search for the local optimum $x_t$ via the Frank-Wolfe algorithm. Compared with $x_{t-1}$, which is used in the original GPC approach as the start point, $x_t'$ is in general closer to $x_t$ and thus the search procedure is more efficient.
As described in Alg. 2, BPF contains three main components: framework for parameterized nonlinear equation systems. The full BPF algorithm is summarized in Alg. 2.

1. Initialization: \( \delta_{\lambda_0} \leftarrow \delta_{\text{min}}, \lambda \leftarrow \lambda_0, t \leftarrow 0 \)
2. Initialization: \( \mathcal{X} \leftarrow \{x_0\}, \Lambda \leftarrow \{\delta_{\lambda_0}\} \)
3. while \( x_i \) not discretized do
   4. \( t \leftarrow t + 1 \)
   5. \( x'_t \leftarrow x_{t-1} + \delta_{\lambda_{t-1}} \left( \sum_{i=t-k}^{t-1} (w_i/d_i) \delta_{\lambda_{t-1}} \right) \) % Prediction
   6. \( p \leftarrow x'_t \) % start point to search local optimum
   7. while \( p \) not converged do
      8. \( y = \arg \min_y \text{tr}(\nabla \mathcal{E}(\lambda)(p) \mathcal{Y} y), \ s.t. \ y \in \mathcal{D} \)
      9. \( \beta = \arg \min_{\beta} \mathcal{E}(\lambda + \beta(y - p)), \ s.t. \ 0 \leq \beta \leq 1 \)
     10. \( p \leftarrow p + \beta (y - p) \)
   11. end while % Frank-Wolfe algorithm
   12. \( x_t \leftarrow p, \ d_t \leftarrow x_t - x_{t-1} \)
   13. if \( \text{tr}(x_t - x'_t)^T (x_t - x'_t) \leq \theta \) then
      14. \( \delta_{\lambda_t} \leftarrow \rho \delta_{\lambda_{t-1}} \)
      15. else
      16. \( \delta_{\lambda_t} \leftarrow \max(\delta_{\text{min}}, \delta_{\lambda_{t-1}} - \rho \) % Adaptive step length
   17. end if
   18. \( \lambda \leftarrow \lambda + \delta_{\lambda_t} \)
   19. \( \mathcal{X} \leftarrow \mathcal{X} \cup \{x_t\} \)
   20. \( \Lambda \leftarrow \Lambda \cup \{\delta_{\lambda_t}\} \)
   21. end while
   22. return \( \mathcal{X} \)

Algorithm 2 Branching Path Following.

% \( T \): the set of candidate solutions, i.e., end points of branching paths.
% \( P_i \{u_{00}, \ldots, u_{Nt}\} \): the \( i \)-th branching path with \( N_i + 1 \) solution points.
1. Initialize a start point \( (\lambda_0, u_0) \)
2. Initial path \( P_0 \leftarrow \text{APE}(\lambda_0, u_0) \) % Adapative path estimation
3. \( T \leftarrow \{\{0\}_i\} \)
4. Discover the set of singular points \( S \) from \( P_0 \)
5. \( i \leftarrow 0 \)
6. for each point \( (\lambda, u) \) in \( S \) do
   7. \( i \leftarrow i + 1 \)
   8. Compute the solution \( (\lambda^*, u^*) \) of Eq. (23)
   9. \( P_i \leftarrow \text{APE}(\lambda^*, u^*) \) % Adapative path estimation
10. \( T \leftarrow T \cup \{u_{N_i}\} \)
11. end for
12. Select the best \( u^* \) from \( T \) in line with the objectives
13. Return \( u^* \) as the final solution

6 Branching Path Following

Although APE gains much higher computational efficiency than the original GPC approach as discussed above, it still meets the problem at singular points as GPC. In this section, we propose the branching path following (BPF) strategy that branches new paths at singular points toward potentially better solutions for nonlinear equation systems. The full BPF algorithm is summarized in Alg. 2.

Based on the numerical continuation method, BPF is a general framework for parameterized nonlinear equation systems. Specifically, we apply BPF to the graph matching problem in this paper. As described in Alg. 2, BPF contains three main components:

- **Singular point discovery** (line 4): Given an initial solution path, it finds potential singular points by determining the signs of determinants of Jacobian on consecutive sampled points.
- **Branch switching** (line 8): It branches new paths to be searched at discovered singular points by reparameterizing the problem with an introduced arclength parameter.
- **Adaptive path estimation** (lines 2 and 9): Starting from a given initial point, it searches along the path with adaptive steps based on path estimation.

We first discuss each component of BPF in detail in the following subsections, and subsequently apply BPF to GNCCP to develop a new graph matching algorithm, followed by analysis of optimality and computational complexity.

### 6.1 Singular points discovery

It is an essential step for BPF to discover singular points. Theoretically, these points can be detected by checking whether the Jacobian of \( F \) is of full rank. However, it is impractical to check discrete samples by sampling \( \lambda_t \), since these samples are rarely able to cover the exact singular points.

Denote the Jacobian of \( F \) at specific solution point \( (\lambda, u) \) as

\[
J_\lambda = F_u(\lambda, u) \in \mathbb{R}^{m \times m}.
\]  

A singular point \( (\lambda, u) \) should have determinant \( |J_\lambda| = 0 \). A reasonable assumption is that the curve formed by \( (\lambda, |J_\lambda|) \) over \( \lambda \) is continuous. This implies there is at least one singular point between two points \( (\lambda_1, u_1) \) and \( (\lambda_2, u_2) \) if \( |J_{\lambda_1}| \) and \( |J_{\lambda_2}| \) have different signs.

Thus inspired, we design a simple yet effective way for singular point discovery by checking the signs of determinants of Jacobian on consecutively sampled points. Specifically, denote \( (\lambda_t, u_t) \) the point at iteration \( t \) in the path, we have at least one singular points between \( (\lambda_t, u_t) \) and \( (\lambda_t+1, u_{t+1}) \) if \( |J_{\lambda_t}| \) and \( |J_{\lambda_{t+1}}| \) have different signs.

The determinant of a Jacobian matrix \( J_\lambda \) is computed through LU decomposition \( J_\lambda = LU \), where \( L \) is a lower unitriangular matrix and \( U \) an upper triangular matrix. Since we are only interested in the sign of \( |J_\lambda| \), it can be decided by accumulating the signs of the diagonal elements of \( U \) as \( \prod_i \text{sign}(U_{ii}) \), where \( \text{sign}(\cdot) \) is the sign function defined as

\[
\text{sign}(a) = \begin{cases} 
1, & \text{if } a > 0, \\
-1, & \text{if } a < 0, \\
0, & \text{if } a = 0.
\end{cases}
\]  

### 6.2 Branch switching

Finding the solution path passing a singular point is called branch switching. Our goal is to branch new paths at singular points and then search the best solution along these branched paths.

Once a singular point \( (\lambda_t, u_t) \) is discovered, we branch a new path using the pseudo-arclength continuation (PAC) algorithm [17], [28]. PAC is inspired by the observation that an ideal parameterization of a curve is arclength-based. It provides a
standard strategy to remedy the failure of GPC (and the proposed APE) at simple folds, which happened typically at singular points (for details about PAC and simple fold, please see [17]). More specifically, PAC reparameterizes path following by introducing the arclength parameter \( l \), so that both \( \mathbf{u} \) and \( \lambda \) depend on \( l \). With the parameterization, we extend equations (13) to the following

\[
G(\lambda, \mathbf{u}, l) = \begin{cases} 
F(\lambda, \mathbf{u}) \\
N(\lambda, \mathbf{u}, l) 
\end{cases} = \begin{bmatrix} 0_m \\
0 
\end{bmatrix},
\]

where the normalization equation \( N(\cdot) = 0 \) approximates the statement that \( l \) represents arclength. Denote \((\lambda, \mathbf{u})\) the tangent vector at point \((\lambda_i, \mathbf{u}_i)\), we have

\[
N(\lambda, \mathbf{u}, l) = \mathbf{u}^T(u - \mathbf{u}_i) + \lambda^T(\lambda - \lambda_i) - l = 0.
\]

The resulting modified system makes a step in pseudo-arclength \( l \) (rather than \( \lambda \)), and the step is usually set to be a sufficiently small nonnegative scalar. In this study we set

\[
l = \|\mathbf{u}_{j+1} - \mathbf{u}_j\|_2,
\]

where \((\lambda_j, \mathbf{u}_j)\) is the last regular point such that \( j \leq t - 1 \). Since the path is always smooth at regular points, \( l \) is guaranteed to be a small nonnegative scalar.

According to the implicit function theorem [29], the tangent vector \((\lambda, \mathbf{u})\) can be computed as

\[
(\dot{\lambda}, \dot{\mathbf{u}}) = (\delta_x, \delta_{\lambda} \frac{\partial \mathbf{u}}{\partial \lambda}(\lambda_t, \mathbf{u}_t)) = (\delta_x, \delta_{\lambda}(F_{\mathbf{u}}(\lambda_t, \mathbf{u}_t))^{-1}F_{\lambda}(\lambda_t, \mathbf{u}_t)).
\]

where \( \delta_x \) is the step length of each iteration. Such calculation, however, is inapplicable to the bifurcation point and computationally expensive. We therefore propose approximate of the tangent vector using previous points for computational efficiency. The tangent vector at a previous iteration \( i (i < t) \) is approximated as \((\delta_x, \mathbf{u}_{i+1} - \mathbf{u}_i)\), and the tangent vector \((\lambda, \mathbf{u})\) at \( t \) is estimated as

\[
(\lambda, \mathbf{u}) = (\delta_x, \frac{\sum_{i=1}^{k} (k - i + 1)(\mathbf{u}_{t-i+1} - \mathbf{u}_{t-i})}{\sum_{i=1}^{k} i}),
\]

where \( k \) controls the size of the window used for approximation. The motivation behind this approximation is the smoothness of the path, which implies the similarity between tangent vectors of consecutive iterations. After the tangent vector \((\lambda, \mathbf{u})\) is computed, we solve Eq. (23) using the trust-region-reflective algorithm [6], and then branch a new path using the solution \((\lambda^*, \mathbf{u}^*)\) as the initial solution.

The Jacobian of the pseudo-arclength system (Eq. (23)) is the bordered matrix \[
\begin{bmatrix}
F_{\mathbf{u}} & F_{\lambda}
\end{bmatrix}
\]

Appending the tangent vector as

\[
\begin{bmatrix}
\frac{\partial \mathbf{u}}{\partial \lambda}
\end{bmatrix}
\]

the last row can be seen as determining the coefficient of the null vector in the general solution of the Newton system [16] (particular solution plus an arbitrary multiplication of the null vector). It is known that the bordered Jacobian matrix of the pseudo-arclength system is nonsingular under the assumption of simple folds [17], [28].

Considering the multiple paths around a singular point of the system \(F(\lambda, \mathbf{u}) = 0\), both APE and PAC choose to search along one of them, but the selection mechanisms of APE and PAC are quite different. The selected path of APE is uncontrollable, which is largely dependent on the start point (it is generated via path prediction using the solutions at previous iterations). In contrast, PAC always choose a smooth path from the multiple paths, where the smoothness is achieved by the appended constraint \(N(\lambda, \mathbf{u}, l) = 0\). It is very occasional that our branch switching strategy cannot branch new paths other than the original path computed by APE. In that case, APE coincidentally chooses the same path with PAC.

While PAC can fully replace APE in path following to create generally better solutions, it runs much slower than APE. Fortunately, PAC and APE are equivalent at a regular point where only one path pass through, thus PAC is needed only at singular points which are usually sparsely distributed. Consequently, we utilize APE, rather than PAC, at regular points to reduce the computational burden.

### 6.3 Apply APE and BPF to GNCCP

As one of the latest work following the path following strategy, the \textit{graduated nonconvexity and concavity procedure} (GNCCP) [36] algorithm has demonstrated state-of-the-arts performance in graph matching. GNCCP realizes CPRP on partial permutation matrix without explicitly involving the convex or concave relaxation. It provides a general optimization framework for the combinatorial optimization problems defined on the set of partial permutation matrices, which is not restricted to weighted graph matching of Eq. (2) but is also suitable for the pairwise compatibility form of Eq. (3). Now we describe how to improve GNCCP by integrating the proposed branching path following strategy.

In Alg. 1, line 5 corresponds to the predictor and lines 7 to 11 the corrector via the Frank-Wolfe algorithm. Function \( \mathcal{E}_\lambda \) in lines 8 and 9 is the linear interpolation between the convex relaxation and the concave relaxation of the original objective function, which is the main difference between different path following algorithms (more details in [36], [57], [60]). By specifying \( \mathcal{E}_\lambda \) to be different objective functions, the proposed strategies can be integrated into various path following algorithms. In particular, we integrate the proposed APE and BPF strategies into GNCCP, and denote the enhanced algorithm as ABPF-G.

Note that, in both steps of singular point discovery and branch switching in BPF, we need to compute the Jacobian of \( F \) in advance, which includes a parameter-dependent sub-matrix, the Jacobian of \( \nabla \mathcal{E}_\lambda (x) \) (denoted as \( J_\lambda (x) \)). In GNCCP

\[
\mathcal{E}_\lambda (x) = (1 - |\lambda|)x^T K x + \lambda \text{tr}(x^T x).
\]

Applying BPF here, we have

\[
J_\lambda (x) = (1 - |\lambda|)(K^T + K) + 2\lambda I.
\]

The approach introduces several parameters including the window size \( k \), the minimum step length \( \delta_{\text{min}} \), the step length adaption rate \( \rho \) and the threshold for path estimation \( \theta \). In all the experiments we set \( k = 5, \delta_{\text{min}} = 0.001, \rho = 2 \) and \( \theta = 0.01n \).

### 6.4 Optimality and computational efficiency

BPF computes an initial path given an initial point, and branches new paths at discovered singular points in the initial path. Since the solution of the original algorithm is always in set \( T \), the new algorithm is guaranteed to achieve a better, or very occasionally the same, solution in terms of optimization objectives. It is notable that there may also exist singular points on the newly explored paths, and theoretically these singular points may lead to even better solutions if we branch paths along them recursively. However,
the computational burden of such exhaustive strategy increases exponentially, which makes it impractical in real applications. As a result, we consider only the singular points in the initial path, ignoring possible singular points in newly explored paths.

The computational complexity of GNCCP is $O(n^3)$ [36] for graphs of $n$ nodes. Thus, the computational complexity of the new ABPF-G algorithm is $O(n_p n^3)$, where $n_p$ is the number of explored additional branches. Theoretically, $n_p$ is bounded by the number of samples along the original path, that is, $n_p \leq 1/\delta_{\min}$ because there are at most $1/\delta_{\min}$ singular points. In practice, singular points are very sparse along the path, and in fact we have observed that $n_p \leq 10$ usually holds in all our experiments. As a result, the computational complexity of the proposed ABPF-G algorithm roughly equals to $O(n^3)$ by dropping $n_p$.

Moreover, the APE strategy significantly accelerates ABPF-G, which actually runs in a speed comparable to the original GNCCP. In our experiment, though ABPF-G needs to compute multiple paths to search better solutions, we find it achieves comparable and sometimes even better computational efficiency compared with the original GNCCP algorithm.1 One reason is that the embedded APE strategy accelerates significantly the convergence of searching along each path.

Interestingly, APE by itself brings benefits to not only computational efficiency but also matching results. In fact, ABPF-G in this work obtains better matching performance than that in our preliminary work [54] where APE is absent. The gain in matching performance mainly comes from the smaller step length which is made affordable by integrating APE to accelerate convergence of searching paths.2

Finally, as general strategies, APE and BPF can be applied to other path following algorithms. In such cases, we expect the similar analysis in the optimality and complexity as above.

7 Experiments

In this section, the proposed ABPF-G algorithm is evaluated thoroughly in comparison with six state-of-the-art graph matching algorithms on five datasets, including a synthetic dataset and four public benchmarks. The experimental results clearly support the benefit brought by the proposed APE and BPF strategies.

7.1 Experiment setup

We compare the proposed ABPF-G algorithm with six state-of-the-art graph matching algorithms:

IPFP [33]: The IPFP (integer projected fixed point method) algorithm can take any continuous or discrete solution as inputs and iteratively improve the solution. In our experiments, we start from the initial solution as in [33].

RRWM [12]: The RRWM (re-weighted random walk matching) algorithm reformulates graph matching as a node selection problem and uses an affinity-preserving random walk algorithm to solve the problem.

PSM [20]: The PSM (probabilistic spectral graph matching) algorithm extends the PM algorithm [58] by dropping some constraints, and presents a probabilistic interpretation of the spectral relaxation scheme proposed in [32].

GAGM [24]: The GAGM (graduated assignment graph matching) algorithm employ the graduated assignment technique to iteratively solve a series of linear approximations of the cost function using the Taylor expansion.

FGM [60]: FGM (factorized graph matching) follows the path following strategy and factorizes an affinity matrix into a Kronecker product of smaller matrices, each of them encodes the structure of the graphs and the affinities between nodes and between edges.

GNCCP [36]: GNCCP (graduated non-convexity and concavity procedure) algorithm is one of the latest path following graph matching algorithm. It equivalently realizes CCRP on a partial permutation matrix and provides a much simpler form without involving the convex or concave relaxation explicitly.

Two indicators, matching accuracy and objective ratio, are used to evaluate algorithms. Denote $H \in \{0, 1\}^{n_1 \times n_2}$ the ground-truth assignment matrix where $H_{ij} = 1$ if and only if there is a ground-truth correspondence between $v_i \in V(1)$ and $v_j \in V(2)$. For matching accuracy, denote $X$ the assignment matrix acquired by a matching algorithm, then its matching accuracy $r$ is defined as $r = \text{tr}(X^\top H)/\text{tr}(H^\top H)$.

For objective ratio, denote $\mathcal{E}(i)$ the objective achieved by the $i$-th algorithm, $i = 1, 2, \ldots, 7$, the objective ratio $r_i$ for algorithm $i$ is computed as $r_i = \mathcal{E}(i)/\max_j \mathcal{E}(j)$. Since these algorithms use the same objective function defined in Eq. (3), a higher objective ratio indicates a better solution in line with the objective.

7.2 Synthetic dataset

7.2.1 Matching random graphs

We first perform a comparative evaluation of the algorithms on graphs randomly synthesized following the experimental protocol in [12], [60]. For each trial, we construct two graphs, $G(1)$ and $G(2)$, with $n_{in} = 20$ inlier nodes, and later add $n_{out}$ outlier nodes to both graphs. The edges between nodes are randomly generated independently for each of two graphs to match, with respect to an edge density parameter $\nu \in [0, 1]$ following [60]. Each edge $(i, j)$ in $G(1)$ is assigned with an edge weight $A_{ij}^{(1)}$ sampled uniformly in [0,1]. For $(i, j)$’s corresponding edge $(a, b)$ in $G(2)$, its weight is assigned as $A_{ab}^{(2)} = A_{ij}^{(1)} + \epsilon$, such that $\epsilon \sim N(0, \sigma^2)$ is a Gaussian perturbation. The edge affinity is computed as $K_{\text{ind}(i,a)\text{ind}(j,b)} = \exp(-((A_{ij}^{(1)} - A_{ab}^{(2)})^2/0.15))$ and the node affinity is set to zero.

We compare the performance of the algorithms under three different settings by varying the number of outliers $n_{out}$, edge density $\nu$ and edge noise $\sigma$, respectively. For each setting, we construct 100 different pairs of graphs and evaluate the average matching accuracy and objective ratio. In the first setting (Fig. 2(a)), we fix edge density $\nu = 0.5$ and edge noise $\sigma = 0$, and increase the number of outliers $n_{out}$ from 0 to 10. In the second setting (Fig. 2(b)), we change the edge noise parameter $\sigma$ from 0 to 0.2 while fixing $n_{out} = 0$ and $\nu = 0.5$. In the last case (Fig. 2(c)), the edge density $\nu$ ranges from 0.3 to 1, and the other two parameters are fixed as $n_{out} = 0$ and $\sigma = 0.1$.

The comparison on both objective ratio and matching accuracy is summarized in Fig. 2. It can be observed that in almost all of cases under varying parameters, our approach achieves the best performance in terms of both objective ratio and matching accuracy. All algorithms are harmed by the increase of outliers. 3

1. For fairness, the step length of GNCCP is set to be equal to $\delta_{\min}$ in our experiments, and it is fixed across all iterations in GNCCP. 2. The step length in our preliminary work [54] is fixed to 0.01, comparing $\delta_{\min} = 0.001$ in this work.

3. Although a different formulation based on factorized matrices is adopted in FGM, it is equivalent to the objective function in Eq. (3).
and edge noise, but are benefited from the increase of edge density. From Fig. 2(c), the PSM, RRWM, FGM and GNCCP algorithms are comparable to our approach when the graphs are close to full connections (the density parameter $\rho$ near to 1). All algorithms fail to achieve reasonable solutions when graph pairs present extreme perturbation or sparsity.

7.2.2 Matching 2D points

To evaluate the algorithms in a more realistic scenario, we compare them on the task of finding correspondences between sets of 2D points generated following the experimental protocol in [32]. For each trial, we construct two sets, $\mathbb{P}^{(1)}$ and $\mathbb{P}^{(2)}$, with $n_\text{in} = 10$ inlier points, and later add $n_\text{out}$ outlier points to both sets. The inlier points in $\mathbb{P}^{(1)}$ are randomly selected in a given region of the plane. We then obtain the corresponding inliers in $\mathbb{P}^{(2)}$ by disturbing independently the points from $\mathbb{P}^{(1)}$ with white gaussian noise $\mathcal{N}(0, \sigma)$, and then rotating and translating the whole data set $\mathbb{P}^{(2)}$ with a random rotation and translation. The outlier points of each set are randomly selected from the same region as inliers. The range of the point coordinates in $\mathbb{P}^{(1)}$ is $25\sqrt{(n_\text{in} + n_\text{out})/10}$. Given two pairs of points $(i, j) \in \mathbb{P}^{(1)}$ and $(a, b) \in \mathbb{P}^{(2)}$, the edge affinity is computed as

$$K_{\text{ind}(i,a)\text{ind}(j,b)} = \begin{cases} 4.5 - \frac{(d_{ij} - d_{ab})^2}{2\sigma_d^2}, & \text{if } |d_{ij} - d_{ab}| < 3\sigma_d, \\ 0, & \text{otherwise}. \end{cases}$$

where $d_{ij}$ (respectively $d_{ab}$) denotes the Euclidean distance between points $i$ and $j$ (respectively $a$ and $b$), and the parameter $\sigma_d$ controls the sensitivity of the score on deformations. We set $\sigma_d = 5$ throughout our experiments.

We compare the performance of the algorithms under two different settings by varying the number of outliers $n_\text{out}$ and noise parameter $\sigma$ respectively. For each setting, we construct 100 different pairs of point sets and evaluate the average matching accuracy and objective ratio. In the first setting (Fig. 3(a)), we hold $\sigma = 5$ and increase $n_\text{out}$ from 0 to 20. In the second setting (Fig. 3(b)), we change $\sigma$ from 0 to 10 while fixing $n_\text{out} = 10$. The comparison on both objective ratio and matching accuracy is summarized in Fig. 3. It is shown that, our approach exhibits the best performance in terms of both objective ratio and matching accuracy, in all of cases under varying parameters.

7.3 CMU house dataset

The CMU house dataset includes 111 frames of image sequences, where all sequences contain the same house object with transformation cross sequence gaps. In order to assess the matching accuracy, following [7], 30 landmarks were manually tracked and labeled across all frames. We matched all possible image pairs, in total 560 pairs gapped by 10, 20, ..., 100 frames, where increasing sampling gaps implies the increase of deformation degree. To evaluate graph matching algorithms against noise, we use two different settings of nodes $(n_1, n_2) = (30, 30)$ and $(20, 30)$, where decreasing $n_1$ implies the increase of outlier. In the setting where $n_1 < 30$, $n_1$ points are randomly chosen out of the 30 landmark points.

We model each landmark as a graph node, and then build graph edges by Delaunay triangulation [31]. Each edge $(i, j)$ is associated with a weight $A_{ij}$, which is computed as the Euclidean distance between the connected nodes $v_i$ and $v_j$. The node affinity is set to zero, and the edge affinity between edges $(i, j)$ in $\mathbb{G}^{(1)}$ and $(a, b)$ in $\mathbb{G}^{(2)}$ is computed as

$$K_{\text{ind}(i,a)\text{ind}(j,b)} = \exp(-A_{ij}^2 - A_{ab}^2)/2500).$$
Fig. 5 presents an example for graph matching with 10 outliers under significant deformation. The matching accuracy and objective ratio of each algorithm with respect to the number of outliers are summarized in Fig. 7(a). It can be seen that our approach outperforms other algorithms remarkably in both matching accuracy and objective ratio. In the case when no outlier exists, our method achieves near 90% matching rate, while GNCCP and FGM obtain similar matching performance. With the increase of outlier level, our method shows stronger robustness than other algorithms.

7.5 Willow object dataset

In this experiment, following [10], we work on 500 image pairs using the Willow object class dataset. This dataset provides images...
of five classes, namely car, duck, face, motorbike and winebottle. Each class contains at least 40 images with different instances and 10 distinctive landmarks were manually labeled on the target object across all images in each class. We randomly select 100 pairs of images from each class respectively.

We use Hessian detector [42] to extract interesting points and SIFT descriptor [38] to represent the node attributes. To test the performance against noise, we randomly select 0 \sim 10 outlier nodes from the background. We utilize the Delaunay triangulation to connect nodes and compute the affinity between nodes via their appearance similarity. Edge affinity is computed following the method used in subsection 7.4.

Fig. 8 shows a representative example for graph matching selected from the duck class (10 outliers). The comparison on matching accuracy and objective ratio is summarized in Fig. 7(b). It is observed that our approach again performs better than other algorithms in both matching accuracy and objective ratio. The performance gaps are small when there are no outliers, but the gaps increase roughly in line with the number of outliers.

7.6 Caltech image dataset

The Caltech image dataset [12] contains 30 pairs of real images which are mainly collected from Caltech-101 dataset [35].

The authors provide detected MSER keypoints [18], initial matches, affinity matrix, and manually labeled ground-truth correspondences for each image pair. In [12], candidate matches were filtered according to the distance between SIFT features [38], and all the low-quality candidate matches were filtered if the feature pair has farer distance than a loose threshold $\delta = 0.6$. The affinity matrix is consequently computed by the mutual projection error function [11] considering the similarity between both the appearance-based feature descriptors and the geometric transformations. The FGM and GAGM algorithms are absent in this experiment because the provided affinity matrices cannot be directly utilized in these two algorithms due to its omission of many candidate matches.

Fig. 9 shows a representative example for graph matching with significant deformation and plenty of repeated patterns. The

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Fig. 6. A matching example of motorbike images in the Pascal dataset.

---

Fig. 7. Evaluation on (a) the Pascal dataset and (b) the Willow dataset.
matching accuracy and objective ratio of each algorithm are summarized in Table 1. Our approach gains remarkable improvement compared with the original GNCCP algorithm, and also outperforms other algorithms, in both matching accuracy and objective ratio. It is interesting to see that RRWM is comparative to our approach in terms of matching accuracy, but gains much lower objectives than our approach.

### 7.7 Comparison with base path following algorithms

Being an upgraded algorithm based on GNCCP, the proposed ABPF-G algorithm gains consistent improvement on matching results on all datasets, as described in the previous subsection and summarized in Table 2. It is notable that the better matching performance does not necessarily cost higher computational burden. Although ABPF-G needs to compute multiple paths, embedding of APE accelerates significantly the process of searching each path, and thus results in comparative and even better computational efficiency on some datasets.

To analyze the contributions of the two components (APE and BPF) of our method, we report detailed comparison between the original GNCCP and three variations of our method in Table 2, where APE-G (or BPF-G) denotes the downgraded version that applies the APE (or BPF) component only. With the assistance of the APE component, APE-G improves dramatically the computation efficiency, and achieves the similar matching accuracy compared with GNCCP. Applying the BPF component only, BPF-G gains remarkable improvement on matching results, at the cost of higher computational burden. Compared with the preliminary version of our algorithm in [54] (BPF-G), the new version in this paper (ABPF-G) improves not only in speed but also in accuracy. We observe that ABPF-G achieves higher computational efficiency than GNCCP on the CMU House and the Willow object datasets.

We also provide the average computational time of other com-

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**TABLE 1**

Comparison of graph matching on the Caltech dataset.

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Objective ratio</td>
<td>0.91</td>
<td>0.88</td>
<td>0.76</td>
<td>0.92</td>
<td>1</td>
</tr>
<tr>
<td>Accuracy (%)</td>
<td>68.71</td>
<td>75.49</td>
<td>67.40</td>
<td>68.72</td>
<td>75.51</td>
</tr>
</tbody>
</table>

**TABLE 2**

Comparison with baselines in matching accuracy (%) and computational time (seconds) on four image datasets.

<table>
<thead>
<tr>
<th>Criterion</th>
<th>Method</th>
<th>CMU House</th>
<th>Pascal</th>
<th>Willow</th>
<th>Caltech</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$n_1=30$</td>
<td>$n_1=20$</td>
<td>$n_1=30$</td>
<td>$n_1=20$</td>
<td>$n_1=30$</td>
</tr>
<tr>
<td>Accuracy</td>
<td>GNCCP</td>
<td>100</td>
<td>59.98</td>
<td>58.87</td>
<td>66.73</td>
</tr>
<tr>
<td></td>
<td>APE-G</td>
<td>100</td>
<td>60.88</td>
<td>61.64</td>
<td>65.26</td>
</tr>
<tr>
<td></td>
<td>BPF-G</td>
<td>100</td>
<td>71.76</td>
<td>67.08</td>
<td>68.27</td>
</tr>
<tr>
<td></td>
<td>ABPF-G</td>
<td>100</td>
<td>75.86</td>
<td>70.80</td>
<td>69.41</td>
</tr>
<tr>
<td>Speed</td>
<td>GNCCP</td>
<td>10.6</td>
<td>22.1</td>
<td>33.9</td>
<td>5.03</td>
</tr>
<tr>
<td></td>
<td>APE-G</td>
<td>0.38</td>
<td>0.06</td>
<td>0.29</td>
<td>0.02</td>
</tr>
<tr>
<td></td>
<td>BPF-G</td>
<td>14.3</td>
<td>58.9</td>
<td>23.3</td>
<td>18.6</td>
</tr>
<tr>
<td></td>
<td>ABPF-G</td>
<td>4.52</td>
<td>8.13</td>
<td>74.5</td>
<td>3.12</td>
</tr>
</tbody>
</table>

---

**TABLE 3**

Comparison in computational time (seconds) on four image datasets.

<table>
<thead>
<tr>
<th>Method</th>
<th>CMU House</th>
<th>Pascal</th>
<th>Willow</th>
<th>Caltech</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$n_1=30$</td>
<td>$n_1=20$</td>
<td>$n_1=30$</td>
<td>$n_1=20$</td>
</tr>
<tr>
<td>IPFP [33]</td>
<td>0.01</td>
<td>0.01</td>
<td>0.02</td>
<td>0.01</td>
</tr>
<tr>
<td>GAGM [24]</td>
<td>0.04</td>
<td>0.26</td>
<td>1.26</td>
<td>0.08</td>
</tr>
<tr>
<td>RRWM [12]</td>
<td>0.02</td>
<td>0.05</td>
<td>0.23</td>
<td>0.02</td>
</tr>
<tr>
<td>PSM [20]</td>
<td>0.28</td>
<td>3.36</td>
<td>9.76</td>
<td>1.22</td>
</tr>
<tr>
<td>FGM [60]</td>
<td>0.52</td>
<td>1.28</td>
<td>3.35</td>
<td>2.15</td>
</tr>
<tr>
<td>GNCCP [36]</td>
<td>10.6</td>
<td>22.1</td>
<td>33.9</td>
<td>5.03</td>
</tr>
<tr>
<td>ABPF-G</td>
<td>4.52</td>
<td>8.13</td>
<td>74.5</td>
<td>3.12</td>
</tr>
</tbody>
</table>
pared algorithms in Table 3, where all the compared algorithms are implemented in Matlab. It is shown that IPFP is the most efficient one, while GNCCP and our ABPF-G are more computationally expensive than other algorithms.

8 Conclusion

In this paper, we proposed a novel branching path following strategy for graph matching aiming to improve the matching performance. To avoid the pitfalls at singular points in the original path following strategy, our new strategy first discovers singular points and subsequently branches new paths from them seeking for potentially better solutions. Furthermore, an adaptive path estimation method is devised to accelerate the convergence of searching along each path. We applied the strategy to a state-of-the-art graph matching algorithm that utilizes the original path following strategy. Experimental results reveal that, our new graph matching algorithm, named ABPF-G, gains remarkable improvement on matching performance compared with the original algorithm, and also outperforms other state-of-the-art algorithms.

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