(Brain) Graph Matching via Fast Approximate Quadratic Programming

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Abstract—Graph matching (GM)—the process of finding an optimal permutation of the vertices of one graph to minimize adjacency disagreements with the vertices of another—is rapidly becoming an increasingly important computational problem, arising in fields ranging from machine vision to chemical engineering to neuroscience. Because GM is NP-hard, exact algorithms are unsuitable for today's massive graphs; yet, scalable GM algorithms have received short shrift. GM can be formulated as a quadratic program with linear and binary constraints. We develop a fast approximate quadratic (FAQ) assignment algorithm to approximately solve a relaxed quadratic program with only linear constraints. FAQ scales cubicly with the number of vertices, and demonstrates marked improvements over previous state-of-the-art on nearly all benchmarks. Moreover, our non-convex formulation facilitates multiple restarts; 2-3 wisely chosen initial conditions yield the best objective function on *all* benchmarks. We find qualitatively similar results for our motivating application: brain-graph matching. Unfortunately, the computational complexity of FAQ scales too poorly to use it for mammalian braingraphs, with millions or billions of vertices. To inspire further development of approximate solutions to these problems, this work is available from on the first author's website, http://jovo.me.

Index Terms—graph theory, network theory, statistical inference, structural pattern recognition, connectome.

1 Introduction

RAPH matching—the process of finding an opti-**J** mal permutation of the vertices of one graph to minimize adjacency disagreements with the vertices of another—is a famously computationally daunting problem (see, for example, "Thirty Years of Graph Matching in Pattern Recognition" [1]). Specifically, graph matching is an \mathcal{NP} -hard problem, in particular, we do not know whether a polynomial time algorithm can solve it [2]. Perhaps the most prominent special case of graph matching is the traveling salesman problem [3]. As such, performance of graph matching algorithms are usually evaluated on graphs with ≈ 10 vertices, or at maximum ≈ 100 (see [4] for a description of the standard set of benchmarks). Yet, it is increasingly popular to represent large data sets by a graph, and thus increasingly desirable to consider matching large graphs.

The motivating application for this work is *brain-graph matching*. A brain-graph (aka, a connectome) is a graph for which vertices represent (collections of) neurons and edges represent connections between them [5], [6]. Via Magnetic resonance (MR) imaging, one can image the whole brain and estimate connectivity across

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voxels, yielding a voxelwise connectome with up to $\approx 10^6$ vertices and $\approx 10^9$ edges [7]. Comparing brains is an important step for many neurobiological inference tasks. For example, it is becoming increasingly popular to diagnose neurological diseases via comparing brain images [8]. To date, however, these comparisons have largely rested on anatomical (e.g., shape) comparisons, not graph comparisons. This is despite the widely held doctrine that many psychiatric disorders are fundamentally "connectopathies", that is, disorders of the connections of the brain [9]-[12]. Currently available tests for connectopic explanation of psychiatric disorders hedge upon first choosing some number of graph invariants to compare across populations. The graph invariant approach to classifying is both theoretically and practically inferior to comparing whole graphs via matching [13].

More generally, state-of-the-art inference procedures for essentially any decision-theoretic or inference task follow from constructing interpoint dissimilarity matrices [14]. Thus, we believe that graph matching of large graphs will become a fundamental subroutine of many statistical inference pipelines operating on graphs. Because the number of vertices of these graphs is so large, exact matching is intractable. Instead, we require inexact matching algorithms (also called "heuristics") that will scale polynomially or even linear [1]. We develop an approach to graph matching based on a relaxation of the quadratic programming problem (QAP). Our approach is only cubic in the number of vertices, and outperforms previously proposed approximate graph matching heuristics on a wide range of benchmark datasets as well as our motivating application.

2 GRAPH MATCHING

A labeled graph $G=(\mathcal{V},\mathcal{E})$ consists of a vertex set \mathcal{V} , where $|\mathcal{V}|=n$ is number of vertices, and an edge set \mathcal{E} . Note that we are not restricting our formulation to be directed or exclude self-loops. Given a pair of graphs, $G_A=(\mathcal{V}_A,\mathcal{E}_A)$ and $G_B=(\mathcal{V}_B,\mathcal{E}_B)$, where $|\mathcal{V}_A|=|\mathcal{V}_B|=n$, let Π be the set of permutation functions (bijections), $\Pi=\{\pi\colon \mathcal{V}_A\to \mathcal{V}_B\}$. Now consider the following two closely related problems:

- Graph Isomorphism (GI): Does there exist a $\pi \in \Pi$ such that $(u, v) \in \mathcal{E}_A$ if and only if $(\pi(u), \pi(v)) \in \mathcal{E}_B$.
- Graph Matching (GM): Which $\pi \in \Pi$ minimizes adjacency disagreements between \mathcal{E}_A and the permuted \mathcal{E}_B ?

Both GI and GM are computationally difficult. GM is at least as hard as GI, since solving GM also solves GI, but not vice versa. It is not known whether GI is in complexity class \mathcal{P} [15]. In fact, GI is one of the few problems for which, if $P \neq \mathcal{NP}$, then GI might reside in an intermediate complexity class called \mathcal{GI} -complete. GM, however, is known to be \mathcal{NP} -hard. Yet, for large classes of GI and GM problems, linear or polynomial time algorithms are available [16]. Moreover, at worst, it is clear that GI is only "moderately exponential," for example, $\mathcal{O}(\exp\{n^{1/2+o(1)}\})$ [17]. Unfortunately, even when linear or polynomial time GI or GM algorithms are available for special cases of graphs, the constants are typically unbearably large. For example, if all vertices have degree less than k, there is a linear time algorithm for GI. However, the hidden constant in this algorithm is $512k^3$! (yes, that is a factorial!) [18].

Because we are interested in solving GM for graphs with $\approx 10^6$ or more vertices, exact GM solutions will be computationally intractable. As such, we develop a fast approximate graph matching algorithm. Our approach is based on formulating GM as a quadratic assignment problem (QAP).

3 Graph Matching as a QAP

Graph matching can be formulated as a quadratic assignment problem (QAP). Let $A=(a_{uv})\in\{0,1\}^{n\times n}$ and $B=(b_{uv})\in\{0,1\}^{n\times n}$ correspond to the adjacency matrix representations of two graphs that we desire to match. That is, let $a_{uv}=1$ if and only if $(u,v)\in\mathcal{E}_A$, and similarly for b_{uv} . Moreover, let \mathcal{P} be the set of $n\times n$ permutation matrices $\mathcal{P}=\{P:P^\mathsf{T}\mathbf{1}=P\mathbf{1}=\mathbf{1},P\in\{0,1\}^{n\times n}\}$, where $\mathbf{1}$ is an n-dimensional column vector. We therefore have the following problem:

(QAP)
$$\underset{\pi \in \Pi}{\operatorname{argmin}} \sum_{i,j \in [n]} (a_{ij} - b_{\pi(i)\pi(j)})^2 =$$
 (1a)

$$\underset{P \in \mathcal{P}}{\operatorname{argmin}} \left\| A - PBP^{\mathsf{T}} \right\|_{F} = \tag{1b}$$

$$\underset{P \in \mathcal{P}}{\operatorname{argmin}} \operatorname{tr}(A - PBP^{\mathsf{T}})^{\mathsf{T}}(A - PBP^{\mathsf{T}}) = (1c)$$

$$\underset{P \in \mathcal{P}}{\operatorname{argmin}} - tr(BP^{\mathsf{T}}AP), \tag{1d}$$

where the last equality follows from dropping terms that cancel because P is a permutation matrix. Note that the above algebraic formulation of GM facilitates generalizing the original problem statement. In particular, one can now search for the permutation that minimizes a particular objective function, $f(P) = -tr(BP^\mathsf{T}AP)$. Moreover, it is natural to consider "weighted graph matching" problems, in which each edge is associated with a weight, $a_{uv} \in \mathbb{R}$.

Our approach follows from relaxing the above binary constraints to be non-negative constraints, yielding a quadratic program with *linear* constraints. Thus, the feasible region expands to the convex hull of the permutation matrices: the doubly stochastic matrices, $\mathcal{D} = \{P : P^\mathsf{T}\mathbf{1} = P\mathbf{1} = \mathbf{1}, P \succeq 0\}$, where \succeq indicates an element-wise inequality:

(rQAP) minimize
$$-tr(BP^{\mathsf{T}}AP)$$
 (2a)

subject to
$$P \in \mathcal{D}$$
. (2b)

rQAP—the above relaxed Quadratic Assignment Problem—is quadratic but not necessarily convex, because the Hessian of its objective function is not necessarily positive definite:

$$\nabla^2 f(P) = -B \otimes A - B^\mathsf{T} \otimes A^\mathsf{T},\tag{3}$$

where \otimes indicates the Kronecker product. This means that the solution space will potentially be multimodal, making initialization important. With this in mind, below, we describe an algorithm to find a local optimum of rQAP.

4 FAST APPROXIMATE QUADRATIC ASSIGN-MENT PROBLEM ALGORITHM

Our algorithm, called FAQ, has three components:

- A. Choose a suitable initial position.
- B. Find a local solution to rQAP.
- C. Project onto the set of permutation matrices.

Below, we provide details for each component.

A: Find a suitable initial position. While any doubly stochastic matrix would be a feasible initial point, we choose the "flat doubly stochastic matrix," $J = \mathbf{1} \cdot \mathbf{1}^T / n$, which is the barycenter of the feasible region.

B: Find a local solution to rQAP. As mentioned above, rQAP is a quadratic problem with linear constraints. A number of off-the-shelf algorithms are readily available for finding local optima in such problems. We utilize the Frank-Wolfe algorithm (FW), a successive linear programing problem originally devised to solve quadratic problems with linear constraints [19], [20].

Although FW is a relatively standard solver, especially as a subroutine for QAP algorithms [21], below we provide a detailed view of applying FW to rQAP. Given an initial position, $P^{(0)}$, iterate the following four steps.

Step 1: Compute the gradient $\nabla f(P^{(i)})$: The gradient f with respect to P is given by

$$\nabla f(P^{(i)}) = -AP^{(i)}B^{\mathsf{T}} - A^{\mathsf{T}}P^{(i)}B.$$
 (4)

Step 2: Compute a new putative point $\widetilde{P}^{(i+1)}$: The new putative point is given by the argument that minimizes a first-order Taylor series approximation to f(P) around the current estimate, $P^{(i)}$. The first-order Taylor series approximation to f(P) is given by

$$\widetilde{f}^{(i)}(P) \stackrel{\triangle}{=} f(P^{(i)}) + \nabla f(P^{(i)})^{\mathsf{T}} (P - P^{(i)}). \tag{5}$$

Thus, Step 2 of FW is

$$\widetilde{P}^{(i+1)} = \operatorname*{argmin}_{P \in \mathcal{D}} f(P^{(i)}) + \nabla f(P^{(i)})^{\mathsf{T}} (P - P^{(i)}) \quad \text{ (6a)}$$

$$= \underset{P \in \mathcal{D}}{\operatorname{argmin}} \nabla f(P^{(i)})^{\mathsf{T}} P. \tag{6b}$$

As it turns out, Eq. (6b) can be solved as a *Linear Assignment Problem* (LAP). The details of LAPs are well known [3], so we relegate them to the appendix. Suffice it to say here, LAPs can be solved via the "Hungarian Algorithm", named after three Hungarian mathematicians [22]–[24]. Modern variants of the Hungarian algorithm are cubic in n, that is, $\mathcal{O}(n^3)$, or even faster in the case of sparse or otherwise structured graphs [3], [25]. The $\mathcal{O}(n^3)$ computational complexity of FW was the primary motivating factor for utilizing FW; generic linear programs can require up to $\mathcal{O}(n^7)$.

Step 3: Compute the step size $\alpha^{(i)}$ Given $\widetilde{P}^{(i+1)}$, the new point is given maximizing the *original* optimization problem, rQAP, along the line segment from $P^{(i)}$ to $\widetilde{P}^{(i+1)}$ in \mathcal{D} .

$$\alpha^{(i)} = \operatorname*{argmin}_{\alpha \in [0,1]} f(P^{(i)} + \alpha^{(i)} \widetilde{P}^{(i)}). \tag{7}$$

This can be performed exactly, because f is a quadratic function.

Step 4: Update $P^{(i)}$ Finally, the new estimated doubly stochastic matrix is given by

$$P^{(i+1)} = P^{(i)} + \alpha^{(i)} \widetilde{P}^{(i+1)}. \tag{8}$$

Stopping criteria Steps 1–4 are iterated until some stopping criterion is met (computational budget limits, $P^{(i)}$ stops changing much, or $\nabla f(P^{(i)})$ is close to zero). These four steps collectively comprise the Frank-Wolfe algorithm for solving rQAP.

C: Project onto the set of permutation matrices. Let \widehat{D} be the doubly stochastic matrix resulting from the final iteration of FW. We project \widehat{D} onto the set of permutation matrices, yielding

$$\widehat{P} = \underset{P \in \mathcal{P}}{\operatorname{argmin}} - \langle \widehat{D}, P \rangle, \tag{9}$$

where $\langle \cdot, \cdot \rangle$ is the usual Euclidean inner product, i.e., $\langle X, Y \rangle \stackrel{\triangle}{=} tr(X^{\mathsf{T}}Y) = \sum_{ij} x_{ij} y_{ij}$. Note that Eq. (9) is a LAP (again, see appendix for details).

5 RESULTS

5.1 Algorithm Complexity and leading constants

As mentioned above, GM is computationally difficult; even those special cases for which polynomial time

algorithms are available, the leading constants are intractably large for all but the simplest cases. We therefore determined the average complexity of our algorithm and the leading constants. Figure 1 suggests that our algorithm is not just cubic in time, but also has very small leading constants ($\approx 10^{-9}$ seconds), making using this algorithm feasible for even reasonably large graphs.

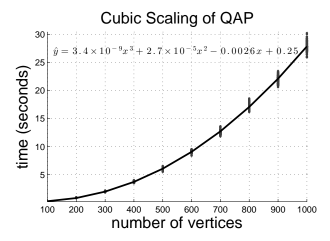


Fig. 1: Running time of FAQ as function of number of vertices. Data was sampled from an Erdös-Rényi model with p = log(n)/n. Each dot represents a single simulation, with 100 simulations per n. The solid line is the best fit cubic function. Note the leading constant is $\approx 10^{-9}$ seconds. FAQ the optimal objective function value in every simulation.

5.2 QAP Undirected Benchmarks

We next assess the computational properties of FAQ in comparison with other the previous state-of-the-art algorithms. We therefore compare FAQ to other approaches using a selection of the QAP benchmark library, QAPLIB [4]. Specifically, [26] created a path following algorithm (PATH) based on a convex and concave relaxation of QAP. In that manuscript, the authors considered 16 datasets from the QAPLIB benchmark, the same 16 datasets as were used in [27], which are known to be "particularly difficult". PATH was shown to outperform other state-of-the-art algorithms on 14 of 16 tests. Specifically, PATH was compared to the Quadratic Programming Bound approach (QGP) of [28], the graduated assignment algorithm (GRAD) of [29], and Umeyama's algorithm (U) [30]. Because either PATH or QBP outperformed GRAD and U on every dataset, Table 1 shows the performance of FAQ versus PATH and QBP. FAQ outperforms both of the previous state-of-the-art cubic algorithms on 13 out of 16 benchmarks. Figure 2 presents the same data graphically. The top panel compares both FAQ and PSOA—which is the minimum of the previous state-of-the-art (either PATH or QBP here) to the absolute minimum; FAQ get closer than PSOA to the minimum on 13 of 16. The bottom panel shows the ratio of FAQ to PSOA.

TABLE 1: Comparison of FAQ with the optimal objective function value and previous state-of-the-art on a set of 16 standard benchmarks from QAPLIB. The best (lowest) value is in **bold**. The number of vertices for each problem is the number in its name (second column).

#	Problem	Optimal	FAQ	PATH	QBP
1	chr12c	11156	13072	18048	20306
2	chr15a	9896	27584	19086	26132
3	chr15c	9504	11936	16206	29862
4	chr20b	2298	3068	5560	6674
5	chr22b	6194	8482	8500	9942
6	esc16b	292	320	300	296
7	rou12	235528	253684	256320	278834
8	rou15	354210	371458	391270	381016
9	rou20	725522	743884	778284	804676
10	tai10a	135028	157954	152534	165364
11	tai15a	388214	397376	419224	455778
12	tai17a	491812	529134	530978	550862
13	tai20a	703482	734276	753712	799790
14	tai30a	1818146	1894640	1903872	1996442
15	tai35a	2422002	2460940	2555110	2720986
16	tai40a	3139370	3227612	3281830	3529402

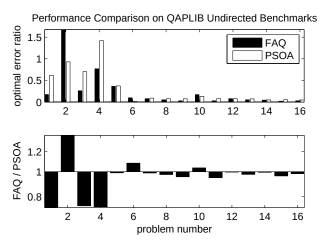


Fig. 2: Performance of FAQ relative to the previous state-of-the-art (PSOA) algorithms on the undirected QABLIB benchmarks. Top: The optimal error ratio is defined: $(\hat{f} - f^*)/f^*$, for \hat{f} being the minimum function value found by each algorithm, and f^* is the optimal (minimum) value. Bottom: Ratio of FAQ minimum to PSOA minimum. Note that both panels indicate that FAQ gets closer to the minimum on 13 of 16 benchmarks.

5.3 QAP Directed Benchmarks

Nothing in the development of our algorithm depends on the graphs being simple; indeed, FAQ applies equally well to directed graphs. To assess the performance of FAQ on directed graphs, we compare the performance of our algorithm to the previous state-of-the-art. Liu et al. recently developed an extended path following algorithm for directed graphs [31]. They compare the performance of their algorithm (EPATH) with several other algorithms on a set of 16 benchmarks from QAPLIB. In particular, they consider U and GRAD, as well as an

algorithm called QCV, which solves a convex relaxation similar to our approach. The EPATH algorithm achieves at least as low objective value as the other algorithms on 15 of 16 benchmarks. Our algorithm, FAQ, always gets the best of the five algorithms. Table 2 shows the numerical results comparing FAQ to EPATH and GRAD, which sometimes did better than EPATH. Note that some of the algorithms achieve the absolute minimum on some benchmarks. Figure 3 compares FAQ to whichever other algorithm did best, clearly indicating that FAQ is the best on these benchmarks.

TABLE 2: Comparison of FAQ with optimal objective function value and previous state-of-the-art for undirected graphs. The best (lowest) value is in **bold**. Asterisks indicate achievement of the global minimum. The number of vertices for each problem is the number in its name (second column).

ш	Problem	O ti1	D20	DD 3 BII	CDAD
#		Optimal	FAQ	EPATH	GRAD
1	lipa20a	3683	3791	3885	3909
2	lipa20b	27076	27076*	32081	27076*
3	lipa30a	13178	13571	13577	13668
4	lipa30b	151426	151426*	151426*	151426*
5	lipa40a	31538	32109	32247	32590
6	lipa40b	476581	476581*	476581*	476581*
7	lipa50a	62093	62962	63339	63730
8	lipa50b	1210244	1210244*	1210244*	1210244*
9	lipa60a	107218	108488	109168	109809
10	lipa60b	2520135	2520135*	2520135*	2520135*
11	lipa70a	169755	171820	172200	173172
12	lipa70b	4603200	4603200*	4603200*	4603200*
13	lipa80a	253195	256073	256601	258218
14	lipa80b	7763962	7763962*	7763962*	7763962*
15	lipa90a	360630	363937	365233	366743
16	lipa90b	12490441	12490441*	12490441*	12490441*

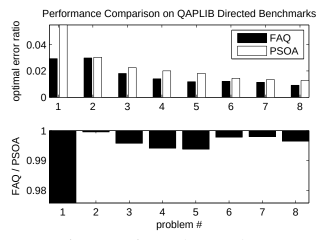


Fig. 3: Performance of FAQ relative to the previous state-of-the-art (PSOA) algorithms on the undirected QABLIB benchmarks. Top and Bottom panels as in Figure 2. Note that FAQ gets closer to the minimum on all 8 benchmarks for which the FAQ and PSOA answer differ.

5.4 rQAP solves QAP in certain special cases

The above numerical results can be strengthened by the below theoretical results. Note that rQAP relaxes the constraints of Eq. (1d), which suggests that in certain important special cases, the minimum of rQAP will be identical to the minimum of QAP. On the other hand, the equality between Eq. (1c) and Eq. (1d) follows from dropping cross-terms that fall out of the optimization because P is constrained to be a permutation matrix. If we had relaxed the constraints prior to canceling those terms, this equality would not follow. This leads us to wonder in which circumstances are the objective functions of QAP and rQAP equal. The following lemma clarifies:

Lemma 1. If A and B are the adjacency matrices of simple graphs (symmetric, hollow, and binary) that are isomorphic to one another, then the minimum of rQAP is equal to the minimum of QAP.

Proof: Because any feasible solution to QAP is also a feasible solution to rQAP, we must only show that the optimal objective function value to rQAP can be no better than the optimal objective function value of QAP. Let $A = PBP^{\mathsf{T}}$, so that $\langle A, PBP^{\mathsf{T}} \rangle = 2m$, where m is the number of edges in A. If rQAP could achieve a lower objective value, then it must be that there exists a $D \in \mathcal{D}$ such that $\langle A, DBD^{\mathsf{T}} \rangle > \langle A, PBP^{\mathsf{T}} \rangle = 2m$ (remember that we are minimizing the negative Euclidean inner product). For that to be the case, it must be that $(DBD^{\mathsf{T}})_{ij} \geq 1$ for some (u,v). That this is not so may be seen by the submultiplicativity of the norm induced by the ℓ_{∞} norm: $\|Dx\|_{\infty} \leq \|D\|_{\infty,\infty} \|x\|_{\infty}$. Applying this twice (once for each doubly stochastic matrix multiplication) yields our result.

5.5 Multiple Restarts

Although FAQ outperformed PSOA on 13 of 16 undirected benchmarks, and always did the best amongst 16 of 16 directed benchmarks, it was annoying to us that we did not do best on all 32 benchmarks. We utilize the non-convexity of rQAP is as a feature, although it can equally well be regarded as a bug (because rQAP is non-convex so the solution found by FAQ depends on the initial condition). It is a feature, however, if (i) we have some reason to believe that better solutions exist (many algorithms efficiently compute relatively tight lower bounds [32]), and (ii) we can efficiently search the space of initial conditions. Although we lack any supporting theory of optimality, we do know how to sample feasible starting points. Specifically, we desire that our starting points are "near" the flat matrix, and satisfy the conditions. Therefore, we sample $K \in \mathcal{D}$, a random doubly stochastic matrix using 10 iterations of Sinkhorn balancing [33], and let our initial guess be $P^{(0)} = (J + K)/2$, where J is the doubly flat matrix. We can therefore use any number of restarts with this approach.

Table 3 shows the performance of running FAQ 3 and 100 times, reporting only the best result (indicated by FAQ3 and FAQ100, respectively), and comparing it to the best performing result from Table 1. It only required three restarts to outperform all other cubic algorithms on all 16 of 16 benchmarks. Moreover, after 100 restarts, FAQ finds the absolute minimum on 3 of the 16 benchmarks. Figure 4 graphically demonstrates these results. Note that restarting FAQ a fixed number of multiple times is still cubic. Future work could investigate performance as a function of the number of restarts.

TABLE 3: Comparison of FAQ with optimal objective function value and the best result from Table 1 on undirected benchmarks. Note that FAQ restarted 100 times finds the optimal objective function value in 3 of 16 benchmarks, and that FAQ restarted 3 times finds a minimum better than the PSOA on all 16 benchmarks.

#	Problem	Optimal	FAQ ₁₀₀	FAQ3	min(FAQ ,PSOA)
1	chr12c	11156	12176	13072	13072
2	chr15a	9896	9896*	17272	19086
3	chr15c	9504	10960	14274	16206
4	chr20b	2298	2786	3068	3068
5	chr22b	6194	7218	7876	8482
6	esc16b	292	292*	294	296
7	rou12	235528	235528*	238134	253684
8	rou15	354210	356654	371458	371458
9	rou20	725522	730614	743884	743884
10	tai10a	135028	135828	148970	152534
11	tai15a	388214	391522	397376	397376
12	tai17a	491812	496598	511574	529134
13	tai20a	703482	711840	721540	734276
14	tai30a	1818146	1844636	1890738	1894640
15	tai35a	2422002	2454292	2460940	2460940
16	tai40a	3139370	3187738	3194826	3227612

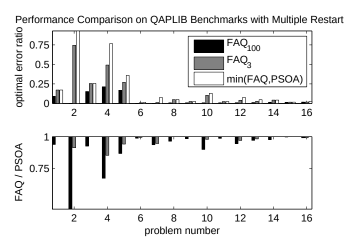


Fig. 4: Performance of FAQ with multiple restarts on the undirected benchmarks. FAQ3 yields a lower objective function value than the best result from Figure 2, and FAQ100 finds the absolute optimal permutation on 3 of the 16 benchmarks. Note that no other algorithm compared ever found the optimal for any of the benchmarks.

5.6 Brain-Graph Matching

A "connectome" is a brain-graph in which vertices correspond to (collections of) neurons, and edges correspond to connections between them. The Caenorhabditis elegans (C. elegans) is a small worm (nematode) with 302 labeled vertices. We consider the subgraph with 279 somatic neurons that form edges with other neurons [34], [35]. Two distinct kinds of edges exist between vertices: chemical and electrical "synapses" (edges). Any pair of vertices may have several edges of each type. Moreover, some of the synapses are hyper-edges amongst more than two vertices. Thus, the connectome of a C. elegans may be thought of as a weighted multi-hypergraph, where the weights are the number of edges of each type. FAQ natively operates on weighted or unweighted graphs. We therefore conducted the following synthetic experiments.

Let $A_{ij;z} \in \{0,1,2,\ldots\}$ be the number of synapses from neuron i to neuron j of type z (either chemical c or electrical e), and let $A_z = \{A_{ij;z}\}_{i,j \in [279]}$ for $z \in \{e,c\}$ correspond to the electrical or chemical connectome. To generate synthetic data, we let $B_z^{(k)} = Q_z^{(k)} A_z Q_z^{(k)}^{\mathsf{T}}$, for some $Q_z^{(k)}$ chosen uniformly at random from \mathcal{P} , effectively shuffling the vertex labels of the connectome. Then, we try to graph match A_z to $B_z^{(k)}$, for $z \in \{e,c\}$ and for $k=1,2,\ldots,10$, that is, we repeat the experiment 100 times. We define accuracy as the fraction of vertices correctly assigned. We always start with the doubly flat matrix.

Figure 5 displays the results of FAQ along with three previous state-of-the-art algorithms on the two connectomes: (i) Umeyama's algorithmU, (ii) a quadratic convex relaxation QCV (which follows from relaxing the permutation matrix constraint to the doubly stochastic constraint in Eq. (1c)), and (iii) PATH. The top left panel indicates that FAQ *always* found the optimal solution for the chemical connectome, whereas none of the other algorithms *ever* found the optimal solution. On the other hand, the top right panel shows that for the electrical connectome, none of the four algorithms ever found the optimal. One hundred restarts of FAQ failed to significantly improve the results.

The bottom panels compare the wall time of the various algorithms, running on an 2.2 GHz Apple MacBook. Note that we have only a Matlab implementation of FAQ, whereas the other algorithms are implemented in C. Nonetheless, FAQ runs nearly as quickly as both U and QCV, and significantly faster than PATH, for both connectomes.

The properties of these connectomes are analyzed in [35]; a cursory evaluation of the properties of these graphs does not suggest to us why the chemical connectome was so much easier to graph match than the electrical one.

To investigate the performance of FAQ on undirected graphs, we ran FAQ on binarized symmeterized versions of the graphs $(A_{ij;z}=1)$ if and only if $A_{ij;z}\geq 1$ or

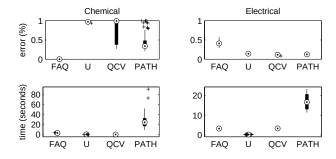


Fig. 5: Performance of U, QCV, PATH, and FAQ on synthetic C. elegans connectome data, that is, graph matching the true connectomes with permuted versions of themselves. Error is the fraction of vertices correctly matched. Circle indicates the median, thick black bars indicate the quartiles, thin black lines indicate extreme but non-outlier points, and plus signs are outliers. The top panels indicate error (fraction of misassigned vertices), and the bottom panels indicate wall time on a 2.2 GHz Apple MacBook. The left panels show the chemical connectome results, and the right panels show the electrical connectome results. For the chemical connectome, FAQ always obtained the optimal solution, whereas none of the other algorithms ever found the optimal. On the other hand, for the electrical connectome, none of the algorithms ever found the optimal. FAQ also ran very quickly, nearly as quickly as U and QCV, and much faster than PATH, even though our FAQ implementation is in Matlab, and the others are in C.

 $A_{ji;z} \geq 1$). The resulting errors are nearly identical to those presented in Figure 5, although speed increased by greater than a factor of two. Note that the number of vertices in this brain-graph matching problem—279—is several times larger than the largest of the 32 benchmarks used above.

6 Discussion

This work presents a fast approximate quadratic assignment problem algorithm called FAQ for approximately solving large graph matching problems, motivated by brain-graphs. Our key insight was to relax the binary constraint of QAP to its continuous and non-negative counterpart—the doubly stochastic matrix—which is the convex hull of the original feasible region. Numerically, we demonstrated that not only is FAQ cubic in time, but also its leading constants are quite small— 10^{-9} suggesting that it can be used for graphs with hundreds or thousands of vertices. Moreover, it achieves better performance than previous state-of-the-art cubic-time algorithms on 29 of the 32 standard QAP benchmarks, including both directed and undirected graph matching problems. Because rQAP is non-convex, we also consider multiple restarts, and achieve improved performance for the remaining three benchmarks using only two or

three restarts. We then demonstrate that the solution to our relaxed optimization problem, rQAP, is identical to that for QAP whenever the two graphs are simple and isomorphic to one another. Finally, we used it to match C. elegans connectomes to permuted versions of themselves. For the chemical connectome, of the four stateof-the-art algorithms considered, FAQ achieved perfect performance 100% of the time, whereas none of the other three algorithms ever achieved perfect performance. On the other hand, all the algorithms struggled with the electrical connectome. Moreover, FAQ ran about as fast as two of them, and significantly faster than PATH, even though FAQ is implemented in Matlab, and the others are implemented in C. Note that these connectomes have 302 vertices, several-fold more than even the largest benchmarks.

Fortunately, our work is not done. Even with very small leading constants for this algorithm, as n increases, the computational burden gets quite high. For example, extrapolating the curve of Figure 1, this algorithm would take about 20 years to finish (on a standard laptop from 2011) when n = 100,000. We hope to be able to approximately solve rQAP on graphs much larger than that, given that the number of neurons even in a fly brain, for example, is $\approx 250,000$. Therefore, more efficient implementations are of interest.

Additional future work might generalize FAQ in a number of ways. First, many (brain-) graphs of interest will be errorfully observed [36], that is, vertices might be missing and putative edges might exhibit both false positives and false negatives. Explicitly dealing with this error source is both theoretically and practically of interest [13]. Second, for many brain-graph matching problems, the number of vertices will not be the same across the brains. Recent work from [26], [37] and [38] suggest that extensions in this direction would be both relatively straightforward and effective. Third, the most "costly" subroutine is LAP. Fortunately, LAP is a quadratic optimization problem with linear constraints. A number of parallelized optimization strategies could therefore potentially be brought to bear on this problem [39]. Fourth, our matrices have certain special properties, namely sparsity, which makes more efficient algorithms (such as "active set" algorithms) readily available for further speed increases. Fourth, for brain-graphs, we have some prior information that could easily be incorporated in the form of vertex attributes. For example, position in the brain, cell type, etc., could be used to measure "dissimilarity" between vertices. Finally, although this approach natively operates on both unweighted and weighted graphs, multi-graphs are a possible extension.

In conclusion, this manuscript has presented an algorithm for approximately solving the quadratic assignment problem that is fast, effective, and easily generalizable. Yet, the $\mathcal{O}(n^3)$ complexity remains too slow to solve many problems of interest. To facilitate further development and applications, all the code and data used in this manuscript is available from the first author's website, http://jovo.me.

APPENDIX

The standard way of writing a Linear Assignment Problem (LAP) is

(LAP) minimize
$$\sum_{u,v \in [n]} a_{u\pi(v)} b_{ij}$$
 (10a)

subject to
$$\pi \in \Pi$$
, (10b)

which can be written equivalently in a number of ways using the notion of permutation matrix introduced in the main text:

$$\underset{P \in \mathcal{P}}{\operatorname{argmin}} \|PA - B\|_F = \tag{11a}$$

$$\underset{P \in \mathcal{P}}{\operatorname{argmin}} \ tr(PA - B)^{\mathsf{T}}(PA - B) = \tag{11b}$$

$$\underset{P \in \mathcal{P}}{\operatorname{argmin}} - tr(PAB^{\mathsf{T}}) = \underset{P \in \mathcal{P}}{\operatorname{argmin}} - \langle P^{\mathsf{T}}, AB^{\mathsf{T}} \rangle =$$
(11c)
$$\underset{P \in \mathcal{P}}{\operatorname{argmin}} - \langle P, AB^{\mathsf{T}} \rangle,$$
(11d)

$$\underset{P \in \mathcal{P}}{\operatorname{argmin}} - \langle P, AB^{\mathsf{T}} \rangle, \tag{11d}$$

where $\langle \cdot, \cdot \rangle$ is the usual Euclidean inner product, i.e., $\langle X, Y \rangle \stackrel{\triangle}{=} tr(X^{\mathsf{T}}Y) = \sum_{ij} x_{ij} y_{ij}$. While the objective function and the first two constraints of LAP are linear, the binary constraints make solving even this problem computationally tricky. Nonetheless, in the last several decades, there has been much progress in accelerating algorithms for solving LAPs, starting with exponential time, all the way down to $\mathcal{O}(n^3)$ for general LAPs, and even faster for certain special cases (e.g., sparse matrices) [3], [25].

That Eq. (6b) is a LAP is evident by considering Eq. (11d). If $\hat{A} = \nabla_P^{(i)}$ and B = I (the $n \times n$ identity matrix), then Eq. (6b) is identical to Eq. (11d).

To solve a LAP, consider a continuous relaxation of LAP, specifically, relaxing the permutation matrix constraint to a doubly stochastic matrix constraint:

$$\begin{array}{lll} \text{(rLAP)} & \underset{P}{\text{minimize}} & -\langle P, AB^\mathsf{T} \rangle & \text{(12a)} \\ & \text{subject to} & P \in \mathcal{D}. & \text{(12b)} \end{array}$$

subject to
$$P \in \mathcal{D}$$
. (12b)

As it turns out, solving rLAP is equivalent to solving LAP.

Proposition 1. LAP and rLAP are equivalent, meaning that they have the same optimal objective function value.

Proof: Although this proposition is typically proven by invoking total unimodularity, we present a simpler proof here. Let P' be a solution to LAP and let $\hat{P} =$ $\sum_{i \in [k]} \alpha_i P^{(i)}$ be a solution to rLAP for some positive integer k, permutation matrices $\{P^{(i)}\}_{i \in [k]}$, and positive real numbers $\{\alpha_i\}_{i \in [k]}$ such that $\sum_{i \in [k]} \alpha_i = 1$. Note that

$$\langle P, AB^{\mathsf{T}} \rangle = \langle \sum_{i \in [k]} \alpha_i P^{(i)}, AB^{\mathsf{T}} \rangle = \sum_{i \in [k]} \alpha_i \langle P^{(i)}, AB^{\mathsf{T}} \rangle$$

$$\leq \sum_{i \in [k]} \alpha_i \langle P', AB^{\mathsf{T}} \rangle = \langle P', AB^{\mathsf{T}} \rangle \leq \langle P, AB^{\mathsf{T}} \rangle,$$

because P' is feasible in rLAP.

This relaxation motivates our approach to approximating QAP.

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RESPONSE TO REVIEWERS

We would like to thank the two helpful reviewers and the editor for their insightful comments. We have extensively revised the manuscript in light of the comments we received. Below are a few general remarks about the current revision, followed by specific responses to reviewer comments (reviewer comments are in **bold**):

- We have significantly revised the text to clarify that we have devised a fast and *approximate* algorithm for solving quadratic assignment problems (QAP). That graph matching can be cast as a QAP is useful because it ties our algorithm into a greater literature on QAP. We were motivated to derive an approximate solution because our application of interest has between hundreds and billions of vertices, so exact algorithms are completely out of the question. We hope that the text of this version of the manuscript conveys that sentiment clearly. In fact, although our algorithm seems to be the best cubic-time approximate algorithm for solving graph matching problems, it is not nearly fast enough. The incoming massive brain-graph data will hopefully motivate others to further improve on our work, which we view merely as a starting point.
- Our algorithm now fundamentally has an initial starting position that is fixed at the barycenter of the feasible region. This is despite that the problem we seek to solve is non-convex, and we do not convexify it. However, by starting at a sensible initial condition (the barycenter of the feasible region), we found that our algorithm outperforms the previous state of the art on 29 of the 32 standard benchmarks, as well as our brain-graph matching problem of interest.
- Submission of this manuscript to TPAMI was motivated by our reading of the very nice paper of Zaslavskiy et al. [26], which was recently published in TPAMI. Zaslavskiy developed a novel PATH following algorithm, that was demonstrated to exceed the previous state of the art on 14 of 16 benchmark datasets selected from the QAP library [4]. Although the authors of that manuscript did not explain their choice of 16, it is the exact same 16 used in a previous comparison of various approximate graph matching algorithms [27]. Moreover, in [27], the authors explain that their choice of those 16 of the >100 datasets was based on those being "particularly difficult". Our algorithm outperformed the PATH algorithm on 13 of the 16 datasets (always starting at the barycenter). If we elect to use only 2 or 3 restarts, which we can only fruitfully do because our objective function is non-convex, we outperform all previous algorithms on all datasets we considered.
- Another paper was published in TPAMI while our manuscript was under review [31]. This paper extended the PATH paper to deal with directed graphs, developing an algorithm called EPATH. The authors of this paper consider 16 directed benchmarks from the QAP library, and demonstrated improve performance on 15 of 16 benchmarks. We applied our FAQ algorithm to those same benchmarks, again always starting at the barycenter, and did at least as well or better than all previous algorithms on all 16 benchmarks.
- Below, we have responded in detail to the reviewers helpful comments. Reviewer comments are in bold, responses are plain text, quotes from the main text are in red.

Reviewer 1

• 1) A principle flaw: The suggested approach samples an generally NP-problem at n=1,3 or 100 random points and search for a local minima in the usually highly non-convex function (the relaxed QAP). So, for increasing problem size the probability to obtain a good solution drops drastically.

In the original draft of this manuscript, we overemphasized the multiple restart aspect of this work. In fact, our algorithm outperforms the previous state of the art cubic-time QAP algorithms on 13 of the 16 benchmarks. We have substantially revised the text to reflect this. Specifically, in the first step of Section 4, we now write: **I: Find a suitable initial position.** While any doubly stochastic matrix would be a feasible initial point, we choose the "flat doubly stochastic matrix," $J = \mathbf{1} \cdot \mathbf{1}^T/n$, which is the barycenter of the feasible region. We have added a subsection in the results to merely remark that multiple restarts are a possibility (Section

- 2) Insufficient evaluation: A report on statistical properties of the approach would make an evaluation of the method possible, i.e, how often is the best solution found by the sampling depending on the the problem size. However such information is not provided in the report.
 - FAQ now always starts with the barycenter of the feasible region. Given this starting position, FAQ outperforms the previous state-of-the-art algorithm on 29 of the 32 benchmarks considered; the particular benchmarks considered were the exact same ones used by the two most recent publications in TPAMI on graph matching [26], [31]
- 3) Too few experiments: The authors have performed 16 experiments on the qaplib dataset which has in its standard form more than 100 problem instances.
 - We have now buttressed our results with another 16 benchmarks, the exact same benchmarks as those used in reference [31] (see Section 5.3). Moreover, the 16 that we used were exactly the 16 datasets used in recent TPAMI publication on the PATH algorithm for graph matching reference [26]. Although the authors of the PATH paper did not explain their choice of 16, they are the same 16 used in reference [27], in which several different graph matching algorithms were compared. In reference [27], it was explained that these 16 are "particularly difficult". Section 5.2 elaborates on this in the text:
 - Specifically, PATH was compared to the Quadratic Programming Bound approach (QGP) of [28], the graduated assignment algorithm (GRAD) of [29], and Umeyama's algorithm (U) [30]. Because either PATH or QBP outperformed GRAD and U on every dataset, Table 1 shows the performance of FAQ versus PATH and QBP.
- Developments reported for example in "N. Brixius and K. Anstreicher. Solving quadratic assignment problems using convex quadratic programing relaxations. Optimization Methods and Software." are completely ignored.
 - Our algorithm, like PATH, outperformed all the algorithms considered in the above mentioned citation. We have now clarified this in Section 5.3:
 - FAQ outperforms both of the previous state-of-the-art inexact cubic algorithms on 13 out of 16 benchmarks. Figure 2 presents the same data graphically. The top panel compares both FAQ and PSOA—which is the minimum of the previous state-of-the-art (either PATH or QBP here)—to the absolute minimum; FAQ get closer than PSOA to the minimum on 13 of 16. The bottom panel shows the ratio of FAQ to PSOA.
- On a different level it seems a bit unrealistic to assume that all the graphs they wish to compare are of equal size, however it is not discussed or even mentioned why the authors have reason to assume that the graphs are equally sized.
 - Indeed, although for some brain-graphs the number of vertices will be the same, for many, they will not be. We have modified Section 6 accordingly:
 - Second, for many brain-graph matching problems, the number of vertices will not be the same across the brains. Recent work from [26], [37] and [38] suggest that extensions in this direction would be both relatively straightforward and effective.
- The authors report a good performance on a quite large graph with 302 nodes, but do not indicate how the graph looks like. That makes it difficult to speculate about reasons that make this problem solvable by just sampling at a few positions followed by the Frank-Wolfe algorithm. It might be that the structure of the problem is more easy (Can the assignment be found by a spectral analysis of the adjacency matrix?) We have clarified Section 5.6 to point out extensive analyses of the brain-graphs by reference [35]:
 - The properties of these connectomes are analyzed in [35]; a cursory evaluation of the properties of these graphs does not suggest to us why the chemical connectome was so much easier to graph match than the electrical one
- minor comments: (The line numbers at the beginning refer to both columns...)
 Thank you, we have corrected all the minor comments.

Reviewer 2

• Though this is a proposal of a QAP inexact solver, it is in principle limited to graphs of the same size. We have now mentioned the possibility of extending this work to deal with different numbers of vertices, and have modified the Section 6 accordingly:

Second, for many brain-graph matching problems, the number of vertices will not be the same across the brains. Recent work from [26], [37] and [38] suggest that extensions in this direction would be both relatively straightforward and effective.

- Even in this case, there is no reference neither experimental nor theoretical to the GA (Graduated Assignment) Algorithm (Gold and Rangarajan, PAMI'1996).
 - We have now clarified in Section 5.2 that our algorithm outperforms the GRAD algorithm on all benchmarks tested:
 - Specifically, PATH was compared to the Quadratic Programming Bound approach (QGP) of [28], the graduated assignment algorithm (GRAD) of [29], and Umeyama's algorithm (U) [30]. Because either PATH or QBP outperformed GRAD and U on every dataset, Table 1 shows the performance of FAQ versus PATH and QBP.
- This is quite simplistic when comparing to the power of deterministic annealing implicit in GA to avoid local minima. A serious comparison between both schemes should be desirable.

 Because our algorithm outperformed a previously published algorithm in TPAMI which outperformed GA (the PATH algorithm), this seemed unnecessary.
- In [25] it is proposed an algorithm outperforming QBP. The method proposed here is quite similar to that in [25] (which is more focused on labeled graphs but based on FW) so I judge the proposal quite incremental. Indeed, our algorithm is similar to the PATH algorithm of reference [25] of the original, [26] in the revision. Yet, FAQ outperforms PATH on 13 of 16 benchmarks, which we believe to be significant. Moreover, our algorithm also addressed directed graphs, while PATH only deals with the undirected case. Reference [31] extends PATH for directed graphs, and our algorithm outperforms their EPATH algorithm (as well as GRAD, QCV, and U) on all 16 benchmarks.
- See similar algorithm in "Many-to-Many Graph Matching: a Continuous Relaxation Approach" in 2010 (arxiv).
 - Thank you for pointing out this reference, as we had not yet seen it. Indeed, this manuscript which is contemporaneous with ours proposes a similar algorithm to FAQ. However, they only consider many-to-one and many-to-many problems. Moreover, our brain-graph matching problem is unique, and motivates the development of algorithms with significantly better scaling rules. We have modified Section 6 accordingly: Second, for many brain-graph matching problems, the number of vertices will not be the same across the brains. Recent work from [26], [37] and [38] suggest that extensions in this direction would be both relatively straightforward and effective.
- Anyway in the experiments only 16 tests where done and the analysis was focuses on the number of restarts. The proposed algorithms require 1-100 starts where in 12 of 16 it only requires 1 start (the van der Waerden matrix).
 - As mentioned above, the 16 tests performed are identical to the 16 tests performed by the previous state of the art, reference [25] in the original submission (the PATH algorithm). And those 16 are the same 16 as used in a previous publication testing various approximate algorithms (reference [30] in the new submission). Moreover, the authors of [30] state that those 16 were chosen because they were "particularly difficult". We have significantly rewritten the text to emphasize that our algorithm fundamentally is initialized in the barycenter of the feasible region. A secondary point in the new submission remarks that random restarts can yield superior performance, time permitting. Section elaborates on this in the text:
 - Specifically, [26] created a path following algorithm (PATH) based on a convex and concave relaxation of QAP. In that manuscript, the authors considered 16 datasets from the QAPLIB benchmark, the same 16 datasets as were used in [27], which are known to be "particularly difficult".
 - Moreover, we have now added another 16 benchmarks, again, the exact same benchmarks used in a previous TPAMI publication, reference [31], to evaluate performance of FAQ on directed graphs. FAQ achieves a superior objective value function for all 16 benchmarks than all previous state-of-the-art algorithms, as described in Section 5.3.
- No comments on the recently embedding methods for graph matching: see for instance: "Graph matching through entropic manifold alignment. CVPR 2011". Indeed only same-size graphs seem to be explored. Thank you for pointing out this omission, we have rectified the text to refer to this work as mentioned above.