Quantum Multi-Model Fitting

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Abstract

Geometric model fitting is a challenging but fundamental computer vision problem. Recently, quantum optimization has been shown to enhance robust fitting for the case of a single model, while leaving the question of multi-model fitting open. In response to this challenge, this paper shows that the latter case can significantly benefit from quantum hardware and proposes the first quantum approach to multi-model fitting (MMF). We formulate MMF as a problem that can be efficiently sampled by modern adiabatic quantum computers without the relaxation of the objective function. We also propose an iterative and decomposed version of our method, which supports real-world-sized problems. The experimental evaluation demonstrates promising results on a variety of datasets. The source code is available at: \url{https://github.com/FarinaMatteo/qmmf}.

1. Introduction

Since the data volumes that AI technologies are required to process are continuously growing every year, the pressure to devise more powerful hardware solutions is also increasing. To face this challenge, a promising direction currently pursued both in academic research labs and in leading companies is to exploit the potential of quantum computing. Such paradigm leverages quantum mechanical effects for computations and optimization, accelerating several important problems such as prime number factorization, database search \cite{shor94} and combinatorial optimization \cite{grover96}. The reason for such acceleration is that quantum computers leverage quantum parallelism of qubits, i.e., the property that a quantum system can be in a superposition of multiple (exponentially many) states and perform calculations simultaneously on all of them.

Among the two quantum computing models, i.e., gate-based and Adiabatic Quantum Computers (AQC), the latter recently gained attention in the computer vision community thanks to advances in experimental hardware realizations \cite{farr05, far16, far17, mag19, men19, arr19}. At the present, AQC prof-

![](chart.png)

Figure 1. \textit{Left:} Differences between our method and HQC-RF \cite{far17}. While HQC-RF considers only a single model and is tested on quantum hardware with synthetic data, QuMF is also evaluated on real data on real quantum hardware. Although devised for multiple models, our method supports a single model likewise. \textit{Right:} Qualitative results of QuMF on motion segmentation on the AdelaideRMF dataset \cite{maz17}.

vide sufficient resources in terms of the number of qubits, qubit connectivity and admissible problem sizes which they can tackle \cite{mcc17}, to be applied to a wide range of problems in computer vision. The AQC model is based on the adiabatic theorem of quantum mechanics \cite{ah02} and designed for combinatorial problems (including NP-hard) that are notoriously difficult to solve on classical hardware. Modern AQCs operate by optimizing objectives in the \textit{quadratic unconstrained binary optimization} (QUBO) form (see Sec. 2). However, many relevant tasks in computer vision cannot be trivially expressed in this form. Hence, currently, two prominent research questions in the field are: 1) \textit{Which problems in computer vision could benefit from an AQC?}, and 2) \textit{How can these problems be mapped to a QUBO form in order to use an AQC?}

Several efforts have been recently undertaken to bring classical vision problems in this direction. Notable examples are works on graph matching \cite{zhu14, yu15}, multi-image matching \cite{far15}, point-set registration \cite{far16, mag19}, object detection \cite{chu16}, multi-object tracking \cite{men19}, motion segmentation \cite{farr05} and robust fitting \cite{far17}. Focusing on geometric model fitting, Doan \textit{et al.} \cite{far17} proposed an iterative consensus maximization approach to robustly fit a single geometric model to noisy data. This work was the first to demonstrate the advantages of quantum hardware in robust single-model fitting with error bounds, which is an important and chal-
lenging problem with many applications in computer vision (e.g., template recognition in a point set). The authors proposed to solve a series of linear programs on an AQC and demonstrated promising results on synthetic data. They also showed experiments for fundamental matrix estimation and point triangulation using simulated annealing (SA) [25]. SA is a classical global optimization approach, that, in contrast to AQCs, can optimize arbitrary objectives and is a frequent choice when evaluating quantum approaches (see Sec. 2). HQC-RF [17] takes advantage of the hypergraph formalism to robustly fit a single model. It is not straightforward to extend it to the scenario where multiple models are required to explain the data.

Multi-model fitting (MMF) is a relevant problem in many applications, such as 3D reconstruction, where it is employed to fit multiple rigid moving objects to initialize multi-body Structure from Motion [3, 37], or to produce intermediate interpretations of reconstructed 3D point clouds by fitting geometric primitives [32]. Other scenarios include face clustering, body-pose estimation, augmented reality and image stitching, to name a few.

This paper proposes QUMF, i.e., the first quantum MMF approach. We propose to leverage the advantages of AQCs in optimizing combinatorial QUBO objectives to explain the data with multiple and disjoint geometric models. Importantly, QUMF does not assume the number of disjoint models to be known in advance. Note that the potential benefit from AQCs for MMF is higher than in the single-model case: when considering multiple models the search space scales exponentially with their number, making the combinatorial nature of the problem even more relevant. Furthermore, we show that QUMF can be easily applied to single-model fitting even though not explicitly designed for this task. We perform an extensive experimental evaluation on quantum hardware with many large-scale real datasets and obtain competitive results with respect to both classical and quantum methods. Figure 1 depicts a visual comparison between HQC-RF [17] and QUMF.

Contributions. In summary, the primary technical contributions of this paper are the following:

- We bring multi-model fitting, a fundamental computer vision problem with combinatorial nature, into AQCs;
- We introduce QUMF, demonstrating that it can be successfully used both for single and multiple models;
- We propose DEQUMF, a decomposition policy allowing our method to scale to large-scale problems, overcoming the limitations of modern quantum hardware. The following section provides the background on AQCs and how to use them to solve QUBO problems. After introducing QUMF and DEQUMF in Sec. 3, we discuss related work in Sec. 4. Experiments are given in Sec. 5. Limitations and Conclusion are reported in Sec. 6 and 7, respectively.

2. Background

The **QUBO formulation** in a binary search space is an optimization problem of the following form:

$$\min_{y \in B^d} y^T Q y + s^T y,$$

(1)

where $B^d$ denotes the set of binary vectors of length $d$, $Q \in \mathbb{R}^{d \times d}$ is a real symmetric matrix and, in a QUBO with $d$ variables, their linear coefficients are packed into $s \in \mathbb{R}^d$. When the problem is subject to linear constraints of the form $Ay = b$, a common approach is to relax such constraints and reformulate the QUBO as:

$$\min_{y \in B^d} y^T Q y + s^T y + \lambda ||Ay - b||_2^2,$$

(2)

where $\lambda$ must be tuned to balance the contribution of the constraint term. Unravelling such a term leads to:

$$\min_{y \in B^d} \tilde{Q} y + \tilde{s}^T y$$

(3)

where the following simple substitutions are implied:

$$\tilde{Q} = Q + \lambda A^T A, \quad \tilde{s} = s - 2\lambda A^T b$$

(4)

For the proof, the reader can refer to, e.g., Birdal et al. [6].

**Adiabatic Quantum Computers (AQCs)** are capable of solving QUBO problems. An AQC is organized as a fixed and architecture-dependent undirected graph, whose nodes correspond to physical qubits and whose edges correspond to couplers (defining the connectivity pattern among qubits) [14]. Such graph structure, in principle, can be mapped to a QUBO problem (1) as follows: each physical qubit represents an element of the solution (i.e., of the binary vector $y$), while each coupler models an element of $Q$ (i.e., the coupler between the physical qubits $i$ and $j$ maps to entry $Q_{ij}$). An additional weight term is assigned to each physical qubit, modelling the linear coefficients in $s$.

Because AQC graphs are not fully connected, mapping an arbitrary QUBO to quantum hardware requires additional steps. This concept is easier to visualize if the notion of **logical graph** is introduced: the QUBO itself can also be represented with a graph, having $d$ nodes (called **logical qubits**1), corresponding to the $d$ entries of $y$ and edges corresponding to the non-zero entries of $Q$.

The logical graph undergoes minor embedding, where an algorithm such as [9] maps the logical graph to the physical one. During this process, a single logical qubit can be mapped to a set of physical qubits, i.e., a chain. All the physical qubits in a chain must be constrained to share the same state during annealing [38]. The magnitude of such

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1Hereafter we will sometimes omit the term “logical” or “physical” as it will be clear from the context which type of qubits are being involved.
constraint is called chain strength and the number of physical qubits in the chain is called chain length. An equivalent QUBO that can be directly embedded on the physical graph is obtained as the output of minor embedding.

Then, optimization of the combinatorial objective takes place: we say that the AQC undergoes an annealing process. In this phase, the physical system transitions from a high-energy initial state to a low-energy final state, representing the solution of the mapped QUBO problem, according to the laws of quantum mechanics [18, 33]. At the end of the annealing, a binary assignment is produced for each physical qubit. A final postprocessing step is needed to go from physical qubits back to logical ones, thus obtaining a candidate solution for the original QUBO formulation (prior to minor embedding). Due to noise during the annealing process, the obtained solution may not correspond to the global optimum of the QUBO problem. In other terms, the annealing is inherently probabilistic and has to be repeated multiple times. The final solution is obtained as the one that achieves the minimum value of the objective function (lowest energy). Performing multiple annealings of the same QUBO form is called sampling.

It is common practice [2, 17, 41, 53] to test the QUBO objective on classic computers using simulated annealing (SA) [25], a probabilistic optimization technique able to operate in the binary space. SA is probabilistic, so solutions are sampled multiple times, as it happens during quantum annealing. In contrast, SA operates on the original QUBO without embedding (i.e. directly on the logical graph) since no physical graph is involved. Importantly, SA can be used to evaluate performance on larger problem instances, in anticipation of larger AQC architectures.

3. The Proposed Method

In this section we derive our approach: first, we define an optimization function for the multi-model fitting task; then, we align it to the standard QUBO formulation, in order to enable quantum optimization; finally, we propose a decomposed version that can manage large-scale problems.

3.1. Problem formulation

Consider a set of data points \( X = \{x_1, \ldots, x_n\} \) and a set \( \Theta = \{\theta_1, \ldots, \theta_m\} \) of tentative and redundant (parametric) models, which, in practice, are obtained by a random sample procedure as done in RANSAC [19]. Specifically, the models are instantiated by randomly selecting the minimum number of points necessary to unambiguously constrain the parameters of a model (e.g., 2 points for a line, 8 for a fundamental matrix). The aim of multi-model fitting is to extract from \( \Theta \) the “best” models that describe all the data \( X \). Equivalently, MMF can be seen as a clustering problem, where points belonging to the same model must be grouped together. This is an ill-posed task since different interpretations of the same data are possible and recovering the best models requires the definition of a proper notion of optimality. To this end, the problem has been cast as the optimization of an objective function encoding different regularization strategies to disambiguate between interpretations and include additional prior about the data. The choice of the objective function is critical and determines the performance of the algorithm.

3.2. Choice of Optimization Objective

In this paper, we focus on the objective function of RANSACOV (Random Sample Coverage) [29] where MMF is treated as a set-cover problem. This choice is due to: i) the simplicity of the objective, which does not involve complex regularization terms as well as does not require to know in advance the number of models; ii) its combinatorial nature, that is well capitalized by a quantum formulation; iii) its generality, as this approach manages the broad situation of fitting multiple models, in contrast to previous work on quantum model fitting [17]. RANSACOV [29] casts MMF as an integer linear program. Such formulation is based on the notion of Preference-Consensus Matrix, which is commonly denoted by \( P \) and defined as:

\[
P[i, j] = \begin{cases} 
1 & \text{if } \text{error}(x_i, \theta_j) < \epsilon \\
0 & \text{otherwise}
\end{cases}
\]  (5)

where: \( x_i \) and \( \theta_j \) are the \( i \)-th point in the dataset and the \( j \)-th sampled model respectively; \( \text{error} \) expresses the residual of point \( x_i \) w.r.t. model \( \theta_j \) and \( \epsilon \) is the inlier threshold to assign the point to the model. \( P \) is a matrix of size is \( n \times m \), where \( n \) and \( m \) are the number of points and models, respectively. Intuitively, the columns of \( P \) represent the consensus sets of models, while the rows are points’ preference sets.

In this setup, selecting the subset of sampled models that best describe the data, is equivalent to selecting the minimum number of columns of \( P \) that explain all the rows. More formally:

\[
\min_{z \in \mathbb{B}^m} \mathbb{1}_n^Tz \quad \text{s.t. } Pz \geq \mathbb{1}_n
\]  (6)

where \( \mathbb{1} \) denotes a vector of ones (whose length is given as subscript) and \( z \) is a binary vector representing which models (i.e., columns of the preference matrix \( P \)) are selected. Solving (6) entails minimizing the number of selected models, while ensuring that each point is explained by at least one of them (as expressed by the inequality constraint). This problem is also known as set cover: we are looking for a cover of the data (i.e., a collection of sets whose union contains all the points) consisting of a minimum number of models, thus discouraging redundancy.

Our goal is to express MMF as a QUBO, in order to make it suitable for optimization on an AQC and capitalize all the advantages of quantum computing. In general, posing a
computer vision problem as a QUBO may not be straightforward, hence a careful choice of assumptions is essential. First, in this paper, we assume that the true number of models/clusters (denoted by $k$) is unknown, so our approach is general. Second, we assume that outliers are not present in the data, i.e., there exist exactly $k$ models that describe all the points. This is not a major limitation because, when outliers are present in the data, the models fitted to outliers could be easily recognized and discarded a-posteriori, using a-contrario reasoning [35], as usually done by MMF algorithms based on clustering [28, 42]. Third, we focus on the case where disjoint models are being sought, which is highly common in practical scenarios. In this situation, the inequality in (6) can be turned into an equality constraint.

These assumptions allow to rewrite the problem in terms of disjoint set cover, which is the basis of our work:

$$\min_{\mathbf{z} \in \mathbb{B}^m} \mathbf{1}_m^T \mathbf{z} \quad \text{s.t.} \ P \mathbf{z} = \mathbf{1}_n. \tag{7}$$

The above constraint, instead of ensuring that each point is explained by at least one selected model, forces each point to be explained by exactly one model, hence imposing the consensus sets of the selected models are disjoint. Problem (7) is the starting point of our QUBO formulation.

### 3.3. QUBO Formulation

Problem (7) is an integer linear program which is known to be NP-hard, hence it represents a suitable candidate to exploit the advantages of quantum optimization. To accomplish such a task, we need to turn (7) into a QUBO. Being a linear program, Problem (7) is, in particular, a special case of a quadratic program with linear equality constraints. Hence, if we pose constraints as soft ones (instead of hard ones), then we can rewrite (7) as (2) using the following correspondences:

$$Q = 0, \quad \mathbf{s} = \mathbf{1}_m, \quad \mathbf{A} = P, \quad \mathbf{b} = \mathbf{1}_n \tag{8}$$

where $m$ is the number of remaining columns in $P$ and $s$ is the number of remaining columns in $P$. Also, $A$, $b$, and $s$ are the choice of a subset of optimal models. Putting together the equalities in (4) and (8), it follows that:

$$\tilde{Q} = \lambda P^T P, \quad \tilde{\mathbf{s}} = \mathbf{1}_m - 2\lambda P^T \mathbf{1}_n \tag{9}$$

where no matrix of quadratic coefficients is present in $\tilde{Q}$ as the considered formulation only contains linear terms, i.e., $Q = 0$. Hence, Problem (7) can be turned into a QUBO of the form (3), namely:

$$\min_{\mathbf{z} \in \mathbb{B}^m} \lambda \mathbf{z}^T (P^T P) \mathbf{z} + (\mathbf{1}_m - 2\lambda P^T \mathbf{1}_n)^T \mathbf{z} \tag{10}$$

Once a QUBO formulation has been obtained for a specific problem as in (10), an AQC can be leveraged to find an optimal solution, following the procedure from Sec. 2. This gives rise to our quantum approach for multi-model fitting, hereafter named QuMF.

### 3.4. Iterative Decomposed Set-Cover

While the capabilities of quantum hardware are rapidly evolving [7, 24], current AQC architectures offer a limited number of physical qubits, thus the maximum dimension of the problems that can be solved is constrained in terms of $m$. In order to overcome this limitation, we present a decomposed approach that iteratively prunes out columns of the preference matrix $P$ and reduces the dimension of the problem to fit the number of available physical qubits.

Our iterative pruning procedure is named DEQuMF and is detailed in Algorithm 1. Instead of directly solving the problem encoded by the preference matrix $P$, at each iteration we partition the columns of $P$ into multiple submatrices $P_{\mathcal{J}}$, where $\mathcal{J}$ indicates the set of selected column indices and $|\mathcal{J}| = s$. We choose the subproblem size $s$ so that the partitioned problems can be solved with high confidence by QuMF. Each subproblem encoded by $P_{\mathcal{J}}$ is independently solved using QuMF (see Sec. 3.3), yielding an optimal binary vector $\mathbf{z}$, that represents the subset of selected models for the current subproblem. Note that models not selected in $\mathbf{z}$ are unlikely to be solutions to the original problem since, by construction, $\mathbf{z}$ maximizes the coverage on all points with only the restricted subset of models $\mathcal{J}$. Then, after all subproblems have been solved, we prune $P$ by retaining only the columns that are selected in the corresponding optimal $\mathbf{z}$. This has the effect of significantly decreasing the dimensionality of the search space (see Fig. 2).

We iteratively apply the above procedure until the number of remaining columns in $P$ falls below $s$, at which point the problem is directly and reliably solved using QuMF.

### 4. Related Work

The proposed approach is the first to address multi-model fitting using AQC. The most related work is HQC-RF [17] that leverages quantum computing as well for the scenario of single-model fitting.
Figure 2. Illustration of the iterative pruning technique applied to $P$ in DeQuMF over consecutive iterations. At time $t$, $P$ is partitioned in different sub-problems of the same size (in the figure, 4 colour-coded problems with 6 models each) which are independently solved with QuMF. At time $t + 1$, models belonging to the sub-solutions (highlighted in red in the figure) are retained; the others are discarded.

4.1. Multi-Model Fitting

Robust fitting is a central problem in computer vision, and the case of multiple models counts several works that can be organized across two main directions, namely clustering-based and optimization-based methods.

Clustering-based methods cast multi-model fitting as a clustering task where points belonging to the same model must be grouped together. Several procedural algorithms have been presented leveraging different schemes: hierarchical clustering [28, 31, 32, 44, 55], kernel fitting [10, 12], robust matrix factorization [30, 43], biclustering [16, 42], higher order clustering [1, 21, 23, 54] and hypergraph partitioning [27, 39, 47, 48, 50].

Our approach falls instead in the category of optimization-based methods. Compared with procedural approaches based on clustering, methods based on the optimization of a precise objective function provide a quantitative criterion for evaluating the quality of the retrieved solution. In particular, our QuMF belongs to consensus-based approaches that generalize RANSAC by maximizing the consensus of the sought models. Sequential RANSAC [46], Multi-RANSAC [56] and RANSA Cov [29] belong to this category as well. Other sophisticated techniques, that integrate consensus with additional priors, have also been proposed: Pearl [22], Multi-X [4] and Prog-X [5].

Our approach is related to RANSA Cov [29] as we consider the same data structure, i.e., the preference matrix, and a similar optimization objective. However, there are substantial differences. RANSA Cov addresses the task via integer linear programming, as Magri and Fusiello [29] use the MATLAB function intlinprog, which employs linear programming relaxation followed by rounding the found solution, in a combination with some heuristics. In case of failure it reduces to branch and bound. Our approach, instead, leverages quantum effects to optimize the objective directly in the space of qubits, where global optimality is expected with high probability after multiple anneals.

4.2. Quantum Computer Vision

Recently, many vision problems have been formulated for an AQC in QUBO forms, hence giving rise to the new field of Quantum Computer Vision (QCV). AQC are especially promising for problems involving combinatorial optimization (see Sec. 2) such as graph matching [40, 41], multi-image matching via permutation synchronization [6], single-model fitting [17], multi-object tracking [53], removing redundant boxes in object detection [26], and motion segmentation [2]. Moreover, solving non-combinatorial problems on quantum hardware is of high general interest as well, since new formulations have unique properties, can be compact and would allow seamless integration with other quantum approaches in future. Thus, several methods address point set alignment [20, 34] and approximate rotation matrices by the exponential map with power series.

Depending on the problem, QUBO formulation can be straightforward [26, 40, 53] or require multiple analytical steps bringing the initial objective to the quadratic form [2, 6, 17]. In most cases, problem-specific constraints must be included in a QUBO as soft linear regularizers, as done in (2). This strategy is followed by the majority of works in the literature, including [2, 6, 17, 40, 53] and our approach. Alternatively, it has been recently shown that it is possible to tackle linearly-constrained QUBO problems via the Frank-Wolfe algorithm [52], which alleviates the need for hyper-parameter tuning, at the price of iteratively solving multiple QUBO problems and sub-linear convergence.

All available approaches can be classified into single QUBO ("one sweep") and iterative methods. The first category prepares a single QUBO that is subsequently sampled on an AQC to obtain the final solution [2, 6, 20, 26, 40]. These methods can solve comparably small problems on current hardware: in the recent QuMoSeg [2], the maximum problem size solved with high accuracy has 200 qubits, which correspond to five images with two motions having ten points each. In QuantumSync [6], real experiments involve matching four keypoints in four images only. The second category of methods, instead, alternates between QUBO preparation on a CPU and QUBO sampling on an AQC, until convergence or a maximum number of iterations is reached [17, 34, 41]. This is done either to overcome the limit on the maximum problem size solvable on
current quantum hardware [34, 41] or because the QUBO represents only a small portion of the whole problem, hence requiring additional (classical) steps as in Q-Match [41] and HQC-RF [17]. Closely related to iterative methods are decomposition approaches (e.g., [53]) that partition the original objective into QUBO subproblems. This paper proposes both a single QUBO method (see Sec. 3.3) and a decomposition-iterative pruning approach (see Sec. 3.4).

**Differences to HQC-RF.** Most related to ours is a hybrid quantum-classical approach for robust fitting, HQC-RF [17], that can be regarded as a pioneer in bringing robust fitting into a QUBO-admissible form, thus paving the way to the adoption of AQCs for such a class of hard combinatorial problems. HQC-RF and our approach have substantial differences. First, although being both based on linear programs, the two formulations do not share common steps: HQC-RF considers the hypergraphs formalism, relying on multiple QUBOs in an iterative framework; our formulation, instead, is more compact as it involves a single QUBO in its one-sweep version. Second, HQC-RF considers a single model, whereas we consider the more general case of multiple models, which is relevant in practical applications and more challenging in terms of the search space and the number of unknowns involved. Extending Doan et al. [17] to MMF is not straightforward (it is not clear so far whether it is possible). Finally, HQC-RF is evaluated with an AQC on synthetic data only, whereas we test our method on several synthetic and real datasets on real quantum hardware, as shown in Sec. 5. Fig. 1 summarizes the core differences between HQC-RF [17] and our approach.

**Remark.** Finally, it is also worth mentioning the technique for robust fitting (single model) by Chin et al. [11] evaluated on a simulator and designed for a different type of quantum hardware than AQC, i.e., a gate-based quantum computer (GQC). GQCs operate differently from AQCs, i.e., they execute unitary transformations on quantumly-encoded data, and currently possess a much smaller number of qubits compared to AQCs [24]. We focus on methods that can run on real quantum hardware. Unfortunately, modern AQCs cannot run algorithms designed for GQCs.

**5. Experiments**

In this section, we illustrate several experiments on synthetic and real datasets to evaluate the proposed approaches. The chosen metric is the misclassification error, that is the percentage of wrongly classified points, as it is common practice in this field. Our comparisons mainly concentrate on the classical RANSACOV [29] and the hybrid quantum-classical HQC-RF [17], which are the most related to our approaches, as explained in Sec. 4. Quantum experiments run on the Dwave Advantage System 4.1, a modern AQC by Dwave built with the Pegasus topology [7], comprising approximately 5000 physical qubits. Communication with the

![Figure 3. Misclassification Error for several methods on the Star5 dataset [44]. The number of points is fixed to n = 30, the number of ground-truth structures is fixed to k = 5, and the number of sampled models m is arranged on the x-axis.](image)

![Figure 4. Relationship between physical qubits and logical qubits in embeddings produced with small-scale preference matrices from the Star5 dataset [44]. AQC is managed via the Ocean software stack [13] and every experiment is executed with an annealing time of 20μs.](image)

**5.1. Synthetic Data**

We consider the synthetic Star5 dataset [44], that is used to fit five lines (depicting a star) given a set of 2D points. Outliers are removed to satisfy our assumptions.

**Small-scale Data.** In this experiment, we select a subset of n = 30 points. The number of ground-truth models is fixed to k = 5, and additional m − k excess models are sampled in order to study performances with increasing problem size. Note that this corresponds to including false models (i.e., columns) in the preference matrix, thereby representing noise. For each configuration, 20 preference ma-

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3We refer to optimal solution probability as the ratio between the anneals ended in a globally optimal state and the total number of anneals.
Figure 5. Misclassification Error for several methods on the Star5 dataset [44]. The number of points \( n \) is fixed to \( n = 250 \), the number of ground-truth structures is fixed to \( k = 5 \), and the number of sampled models \( m \) (corresponding to the dimension of the search space) is arranged on the x-axis.

Matrices are generated and averaged results are reported. We consider our single-QUBO quantum approach QuMF and its CPU counterpart QuMF (SA). We also compare our approaches with RANSACov [29], whose code is available online\(^4\), that can be viewed as a baseline implementing a sophisticated integer programming solver. All the methods receive as input the same preference matrix and optimize the same objective, as already observed in Sec. 4.1.

Results are given in Fig. 3, showing that QuMF can exactly solve problems up to 40 qubits, but it can not solve larger problems with high accuracy: this is an expected behaviour (in line with previous quantum papers, see Sec. 4.2), as current quantum hardware is limited and far from maturity. It is worth noting that our QuMF (SA) achieves excellent results (on par with RANSACov) for all the analyzed problem sizes: considering that QuMF (SA) uses the same QUBO as QuMF, we can thus view the former as an indication on the performances the latter can achieve in the future thanks to the continuous improvements in the quantum hardware.

**Physical vs Logical Qubits.** We analyze here how the allocation of QPU resources (physical qubits) grows with increasing problem size (logical qubits). Similarly to our previous experiment, we use synthetic preference matrices generated from the Star5 dataset with increasing number of columns. However, we consider here all of the 250 points in the dataset in order to study denser cases. Our analysis is displayed in Fig. 4, showing that physical qubits grow super-linearly with respect to logical ones. Such relationship makes it infeasible to allocate real world problems, as they typically require thousands of logical qubits. This finding, given the current state of the hardware, strengthens the need for a decomposition approach. Hence, we analyze our decomposition policy in the next paragraph.

**Large-scale Data.** In this experiment, we consider up to 1000 models in the input preference matrix (also including the \( k = 5 \) ground-truth models) as well as all of the \( n = 250 \) points in the Star5 dataset. Since QuMF is not able to manage such large-scale data, it is not considered here. Instead, it is used as a sub-component of our decomposed approach DeQuMF: each sub-problem has a dimension that can be solved with high accuracy by QuMF, namely 20 qubits. As a reference, we also include QuMF (SA) and its decomposed version DeQuMF (SA) as well as RANSACov [29].

Results are given in Fig. 5, showing that our DeQuMF (SA) achieves nearly zero misclassification error in all the cases, on par with RANSACov. Average DeQuMF errors lie in the range \([0, 10]\)% for all problem sizes, showing the great effectiveness of the decomposition principle. As for QuMF (SA), this specific dataset represents a failure case: after inspecting the solution, we hypothesize that a possible reason is the sparsity pattern of the preference matrix, causing the algorithm to select more models than needed.

### 5.2. Real Data

In this section, we test our framework on several real world datasets, each associated to a different task. Since these are large-scale problems, we do not consider QuMF but its CPU variant QuMF (SA).

**AdelaideRMF dataset** [49]. From this benchmark, we consider the 15 sequences involving multiple fundamental matrices (which, in turn, can be used to perform motion segmentation in two images) and we remove outliers beforehand. All the analyzed methods are given the same preference matrices as input. Results are given in Tab. 2. Both QuMF (SA) and DeQuMF (SA) significantly outperform RANSACov [29] in this scenario. In particular, DeQuMF (SA) performs better than its quantum version DeQuMF, in agreement with our previous experiments.

**Hopkins benchmark** [45]. From this dataset, we consider the Traffic2 and Traffic3 sequences, which comprise 31 and 7 scenes, respectively. Here, the task is to fit multiple subspaces to point trajectories in the context of video motion segmentation. This benchmark provides outlier-free data, hence no pre-processing is needed. Results are given in Tab. 1: in addition to RANSACov [29], other multi-model fitting approaches have been considered, whose results are taken from the respective papers. All the variants of our approach\(^5\) perform reasonably well, with DeQuMF (SA) being on par with the state of the art.

**York Urban Line Segment Database** [15]. This dataset contains 102 images used for vanishing point detection from multiple line segments without outliers. Results are presented in Table 1, following the same rationale as per the Hopkins benchmark. Such results are in line with our previous experiments: in particular, DeQuMF (SA) exhibits competitive performances with the state of the art.

\(^4\)https://fusiello.github.io/demo/cov/index.html

\(^5\)We are not able to run QuMF on Traffic2 due to the limited amount of QPU time available in our monthly subscription combined with the large dimension of the dataset, hence results are not reported.
Table 1. Misclassification Error [%] on the Traffic3 and Traffic2 subsets of the Hopkins benchmark [45] and on the York DB [15].

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<th>Method</th>
<th>Outlier ratio</th>
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<tbody>
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<td>Multi-X [4]</td>
<td>10%</td>
</tr>
<tr>
<td>J-Linkage [44]</td>
<td>20%</td>
</tr>
<tr>
<td>T-Linkage [28]</td>
<td>Full sequences</td>
</tr>
<tr>
<td>RPA [30]</td>
<td></td>
</tr>
<tr>
<td>RANSAC [29]</td>
<td></td>
</tr>
<tr>
<td>QU-MF (SA)</td>
<td></td>
</tr>
<tr>
<td>DeQU-MF</td>
<td></td>
</tr>
<tr>
<td>DeQU-MF (SA)</td>
<td></td>
</tr>
</tbody>
</table>

Table 2. Misclassification Error [%] for several methods on the 15 multi-model sequences of the AdelaideRMF dataset [49].

Table 3. Misclassification Error [%] for quantum methods on the single-model sequences of the AdelaideRMF dataset [49].

Figure 6. A sample of the worst results of the considered quantum methods on the book image pair from the AdelaideRMF [49] with 20% outliers. Outliers and inliers are shown in red and green, respectively. On the right, HQC-RF [17] classifies all the points as outliers, whereas our method achieves 8.49% error.

6https://github.com/dadung/HQC-robust-fitting

5.3. Comparison with Quantum State-of-the-art

Our method is the first quantum solution to multi-model fitting, hence there are no quantum competitors in the literature. The most similar work to ours is HQC-RF [17], whose code is available online6, which addresses single-model fitting in the presence of outliers (see Sec. 4.2). HQC-RF has strong theoretical justifications, and the proposed algorithm terminates with either the globally optimal solution or a feasible solution paired with its error bound. Neither QU-MF or DeQU-MF have this property. On the other hand, our framework is general and can be used for a single model as well, allowing for an experimental comparison with HQC-RF [17]. Specifically, we exploit the information that only a single model is present in the data and we select, from the solution returned by set-cover, only the model with the maximum consensus set. All the other models in the solution are treated as outliers.

We focus on fundamental matrix estimation and use the 4 sequences of the AdelaideRMF dataset [49] characterized by the presence of a single model. Since this is a large-scale dataset, we omit QU-MF and consider QU-MF (SA), DeQU-MF and DeQU-MF (SA), as previously done. We analyze how performances vary with different outlier ratios: we consider 10% and 20% of outliers, in addition to the evaluation on the original sequences, where no fixed outlier ratio is imposed. Results are summarized in Tab. 3.

HQC-RF shows remarkable behaviour in controlled scenarios with low outlier ratios; however, it is worth noting that our method is more robust to higher outlier ratios, although not explicitly designed for single-model fitting. See also Fig. 6 for qualitative results of worst-case scenarios.

6https://github.com/dadung/HQC-robust-fitting

6https://github.com/dadung/HQC-robust-fitting

6https://github.com/dadung/HQC-robust-fitting

7. Conclusion

We brought a new, very important problem into AQC-admissible form. Our method shows promising results on a variety of datasets for MMF and even surpasses the quantum SotA on single-model fitting, although not explicitly designed for such task. We believe it offers many avenues for future research.

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