

# Quantum Annealing-Based Software Components: An Experimental Case Study with SAT Solving

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## ABSTRACT

Quantum computers have the potential of solving problems more efficiently than classical computers. While first commercial prototypes have become available, the performance of such machines in practical application is still subject to exploration. Quantum computers will not entirely replace classical machines, but serve as accelerators for specific problems. This necessitates integrating quantum computational primitives into existing applications.

In this paper, we perform a case study on how to augment existing software with quantum computational primitives for the Boolean satisfiability problem (SAT) implemented using a quantum annealer (QA). We discuss relevant quality measures for quantum components, and show that mathematically equivalent, but structurally different ways of transforming SAT to a QA can lead to substantial differences regarding these qualities. We argue that engineers need to be aware that (and which) details, although they may be less relevant in traditional software engineering, require considerable attention in quantum computing.

## KEYWORDS

Quantum Computing, Quantum Annealing, Boolean Satisfiability, Experimental Performance Analysis

### ACM Reference format:

Tom Krüger and Wolfgang Mauerer. 2020. Quantum Annealing-Based Software Components:

An Experimental Case Study with SAT Solving. In *Proceedings of IEEE/ACM 42nd International Conference on Software Engineering Workshops*, Seoul, Republic of Korea, May 23–29, 2020 (ICSEW’20), 7 pages.

DOI: 10.1145/3387940.3391472

## 1 INTRODUCTION

The upcoming end of Moore’s law and the trend towards energy efficient systems, but the likewise ever-growing need for more computational power pose substantial challenges to systems engineering and software architecture. New computational approaches that substantially diverge from technologies established during the last decades start to graduate from research laboratories into first working prototypes. Especially quantum computing has gained

substantial attraction during the last years [24]. Programming quantum computers (QC) differs drastically from previously established techniques and approaches. Integrating QC into existing appliances must not only be addressed at the level of algorithmic implementation, but also concerns many of the broader issues investigated in software engineering [4]. In this paper, we argue that the problem at the current stage of development must be considered at a much lower level of abstraction than is customary in software engineering, and illustrate this by a case study of how to transition a core computational primitive—solving binary satisfiability problems—from classical to quantum in existing software architectures. Our study illustrates that defining and testing specific quality properties of QC components is one of the crucial challenges. These properties do not play a central role in traditional engineering, but must be considered in software architectures with quantum components. We illustrate this by analysing different approaches—one of which has been specifically designed for this paper, and improves considerably on the state-of-the-art—to solving the binary satisfiability (SAT) problem. We hope this helps readers to form a realistic intuition of near- and mid-term capabilities, potentials and challenges of augmenting software with quantum components.

## 2 QUANTUM ANNEALING

By utilising quantum mechanical properties, QCs are expected to solve some problems more efficiently than their classical counterparts. Simulations of quantum systems [23] and chemical reactions [25], breaking of cryptographic codes [28], but also optimising portfolios [37] are among the list of candidate problems. Recent advances—although not undisputed—claim *quantum supremacy* [3], even if for extremely artificial problems. Real-world adoption of quantum computing, as it matters to software engineering, is likely to happen in an evolutionary way than by disruptive revolution.

We base our considerations on quantum *annealers*: Many early potential industrial use-cases [22, 31, 32] rest on this class of machines, in part because they were among the first offerings available for commercial use (discussions about the full quantum mechanical nature of such machines [27] are not relevant for our purposes).

Especially NP-complete problems, which are known to be classically intractable for inputs of growing size when non-approximate solutions are desired, are candidates for which (polynomial) quantum speed-ups would be desirable. Many NP-complete problems of practical interest are known. Especially the Boolean satisfiability problem (SAT) has received substantial attention because many use cases, from system verification to constrained planning problems [13], have SAT at their core. Quantum annealers are particularly well suited to process problems of this type [16]. They

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ICSEW’20, Seoul, Republic of Korea

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DOI: 10.1145/3387940.3391472

differ considerably from gate-based approaches in their physical realisation, and in the ways programs are engineered.

## 2.1 Using Quantum Annealing Primitives in Existing Software Architectures

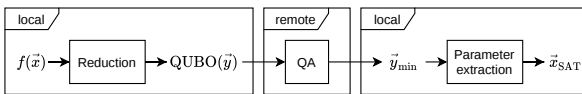
Software engineering is (ignoring many aspects that we cannot address for the lack of space) concerned with development, integration, and testing (verification, validation, performance, quality, ...) of software [4]. This impacts quantum software development:

**2.1.1 Development.** A considerable body of previous research devoted to developing languages for programming quantum systems focuses on gate-based approaches (e.g., [11, 30, 34]). A growing number of quantum programming languages has been devised for this hardware class (e.g., [10, 17, 33]). Roughly speaking, gate based quantum computers relate to quantum annealers like imperative programming languages do to declarative approaches. A deep understanding of quantum mechanics is not required to use current QA hardware, which is beneficial from a software point of view. Engineers can resort to techniques known from constraint programming, optimisation, and problem reduction.

**2.1.2 Integration.** As Knill [14] discussed as early as 1996, quantum computers will not entirely replace traditional machines, but will be part of hybrid quantum-classical architectures, not unlike GPUs [1, 12] or other accelerators (TPUs, FPGAs, ...). Quantum annealers can be seen as hardware accelerators for approximating quadratic unconstrained binary optimisation (QUBO) problems.

The ability to easily replace functional components of a software architecture is a crucial element of component-based software engineering [35, 36], and many existing applications are designed along these ideas. In the following, we consider that a SAT solving component is supposed to be replaced by a QA device in an existing software architecture.

Replacing a library function call to a traditional solver by a network-based job submission to a QA device is an easy programming tasks that we do not consider any further. However, two data conversions are necessary, as Figure 1 illustrates: The propositional calculus formula for which a solution is sought must be mapped to a QUBO. Once the result of the optimisation process is available, it must be translated back to the original SAT model. Both steps can be trivially abstracted by an interface.



**Figure 1: Interface wrapping: Classical SAT solvers can be replaced with a QA based implementation with limited effort.**

**2.1.3 Testing.** Miranskyy and Zhang [20] discuss testing aspects related to verification and validation of quantum programs. Fundamental properties of NP-complete problems guarantee that solutions can be verified in polynomial time [29], and consequently, validation and verification of QA programs is not a core challenge.

However, quantum annealers usually only deliver *approximate* solutions to problems, and the quality of approximation is closely related to how “programs” (in the form of mathematical reductions) are created. We focus on the issues arising from this scenario in the rest of the paper.

## 2.2 Workflow

The workflow for solving problems on quantum annealers is more involved than for classical constraint optimisation. The necessary process comprises five stages, and choices in some of the stages can greatly influence performance and accuracy of computations. Consequently, some knowledge of the inner working of the quantum annealing process are useful. An AQO computation proceeds along the following stages [12, 18]:

**PROBLEM REDUCTION:** Like classical constraint optimization solvers, QA machines can optimise a specific class of models. Annealers can find solutions to *quadratic unconstrained binary optimization* (QUBO) problems [15], which are given by

$$\min[\vec{x}] \left( \sum_i c_{ii} x_i + \sum_{i,j} c_{ij} x_i x_j \right) \quad (1)$$

with  $x \in \{0, 1\}$  and  $c \in \mathbb{R}$ . A QUBO can be represented by a weighted graph with nodes  $x_i$  and associated weights  $c_{ii}$ . Weighted edges are given by  $c_{ij}$ .

Reducing a given problem  $p$  to a QUBO,  $p \leq \text{QUBO}$ , requires no knowledge of quantum mechanics, and is similar to well-known reductions to Boolean satisfiability problems. As we will discuss later, structurally different (but logically equivalent) reductions can lead to drastically different performance on contemporary hardware.

**HARDWARE EMBEDDING:** Software solvers can react dynamically to input, and easily build arbitrary data structures. For QA, the “data structure” used to represent a given input is fixed in hardware. This step “translates” an input onto the hardware structure [1, 7] (see Figure 2). Mathematically equivalent reductions can lead to pronounced differences in solution quality, as we show in Section 3.

**HARDWARE PROGRAMMING:** The problem embedding needs to be transferred to the machine. The physical details of this operation are irrelevant to programmers, except that some parameters—most importantly, the duration of the annealing process—can be influenced. Finding an optimal value is currently a matter of experimentation.

**EXECUTION:** The machine finds a solution to the optimisation problem by “executing” a physical process.

**POST PROCESSING:** Results obtained in the previous step are usually only *close* to the desired optimum. Classical post-processing can improve solution quality [9]. We will ignore this step in this paper since we are interested in the capabilities of QA as such, and not of classical data processing.

## 2.3 Experimentation Platform

All experiments that we discuss in the following were performed on a D-Wave 2000Q quantum annealer, model DW\_2000Q\_2.1. The machine can be remotely accessed via a Python-based API. Performing computations requires to specify a problem QUBO, and

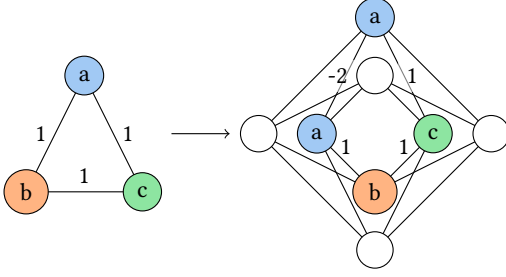


Figure 2: Example for embedding a *logical* graph that describes couplings between qubits (left) into a *physical* qubit structure (right) with limited connectivity. Node “a” is mapped to a chain of two nodes representing “a”, which illustrates that the amount of physical qubits required to represent a problem is larger than the amount needed for a structural description.

(essentially) anneal time and desired sample size  $n$ . Once the QA has evaluated the problem, a result set with  $n$  samples is returned. Each sample contains an assignment for all qubits.

While it is possible to arbitrarily weigh the interaction between qubits as specified by term  $c_{ij}$  from Eq. (1), there are substantial restrictions on which qubit  $i$  can *physically* and *directly* interact with which qubits  $j$  (see Ref. [5] for details on the available hardware graph structure). This limited connectivity poses a major practical challenge when mapping logical to physical problems, since a pair of nodes that requires a logical connection (a non-zero entry  $c_{ij}$  in Eq. (1)) must be represented by a chain of multiple nodes on the hardware graph. This considerably limits the number of *effectively usable* qubits as compared to the number of *physically available* qubits—Figure 6 exemplifies the problem visually. In general, longer chains lead to more undesirable physical perturbation, and decrease result quality [26]. As a rule of thumb, the number of usable logical qubits is only about 5-10% the number of physical qubits.

### 3 QUALITY ASSESSMENT OF QUANTUM 3-SAT

Let us now turn our attention to discussing how implementation details of quantum computational primitives can influence qualities of software. We focus on the problem of finding and comparing reductions of the problem to a machine specific structure. Such low-level issues are usually not of much relevance in software engineering, and are justifiably perceived as implementation details—however, we show that this level of abstraction is far from reached on quantum machines yet.

The  $k$ -SAT problem, the cornerstone of NP-completeness [13], serves as an example. We first discuss different reductions of  $k$ -SAT  $\leq_P$  QUBO, and show how differences arise from seemingly small details. We then offer guidance on comparing reductions.

#### 3.1 Problem Definition

The problem of Boolean satisfiability is well known: Let  $X := \{x_1, x_2, \dots, x_n\}$  be a set of Boolean variables, and let *literals* be defined as  $L := \{l | l \in \{x, \bar{x}\}, x \in X\}$ . The set of all *clauses* is given by

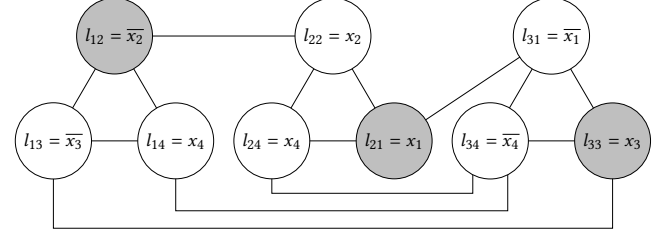


Figure 3: Graphical illustration of a QUBO formula that represents  $f(\vec{x}) = (\bar{x}_2 \vee \bar{x}_3 \vee x_4) \wedge (x_1 \vee x_2 \vee x_4) \wedge (\bar{x}_1 \vee x_3 \vee \bar{x}_4)$  using Choi’s reduction. Grey nodes represent a satisfying assignment  $[x_1 \mapsto 1, x_2 \mapsto 0, x_3 \mapsto 1]$ .

$C := \{C_i | i \in [1; n], C_i \subset L, |C_i| = k\}$ . For each  $x_i \in X$ , there exists at least one  $C_j$  such that  $x_i \in C_j$ . A function  $f(\vec{x}) = \bigwedge_{C_i \in C} \bigvee_{l \in C_i} l$  that satisfies these conditions is called a  $k$ -CNF function. Given a  $k$ -CNF function  $f(\vec{x})$ , the  $k$ -SAT problem is to find an assignment  $\vec{x}_t$  such that  $f(\vec{x}_t) = \text{true}$ . It is textbook knowledge [29] that every CNF formula can be cast in 3-CNF form.

The  $k$ -SAT problem is the cornerstone of NP-completeness, but not all specific instances are difficult to solve. The hardness of an instance depends on the ratio of clauses per variable  $\alpha = \frac{|C|}{|X|}$  [6]. For instances with few clauses per variable (small  $\alpha$ ), it is easy to find satisfying assignments. For instances with many clauses per variable, it is easy to find contradictions. Instances with large or small values of  $\alpha$  tend to be easy to solve. In an  $\alpha$  region surrounding  $\alpha_c \approx 4.25$ , the probability that a random  $k$ -SAT formula can be satisfied drops abruptly from 1 to 0 [6, 21], and the hardest instances are contained in this parameter regime. Improvements in SAT solving are therefore most desirable around this *phase transition*.

#### 3.2 Choi’s Standard Reduction

Choi [8] gives a standard reduction from  $k$ -SAT to a QUBO. Let  $l_{ij}$  be the literal of variable  $x_j$  in clause  $i$ . Two literals  $l_{ij}$  and  $l_{i'j}$  are *in conflict* if  $l_{ij} = \bar{l}_{i'j}$ . Satisfying a Boolean function in CNF implies at least one satisfied, but conflict-free literal per clause.

The reduction assigns a negative weight  $-\omega$  to  $l_{ij}$ :  $-\sum_{l_{ij}} \omega l_{ij}$ . All literals of a clause are fully connected with positive weighted edges:  $\sum_{l_{ij}, l_{i'j'} \in C_i} \delta l_{ij} l_{i'j'}$ . All conflicting literals of the same variable are pairwise connected with *positive* edge weights:  $\sum_{l_{ij} = \bar{l}_{i'j}} \delta l_{ij} l_{i'j}$ . The last two sums are pure penalty terms and evaluate to 0 for correct assignments. This leads to a definition illustrated in Figure 3:

**Definition 3.1** ( $k$ -SAT  $\leq_P$  QUBO (MIS)). Let  $f(\vec{x})$  be a boolean  $k$ -CNF function. The literal of a variable  $x_j$  in  $\vec{x}$  in a clause  $C_i \in C$  is given by  $l_{ij} \in \{0, 1\}$ . Under the constraint  $\forall \delta, \omega : \delta > \omega > 0$ ,

$$\min[\vec{x}] \left( - \sum_{l_{ij}} \omega l_{ij} + \sum_{l_{ij}, l_{i'j'} \in C_i} \delta l_{ij} l_{i'j'} + \sum_{l_{ij} = \bar{l}_{i'j}} \delta l_{ij} l_{i'j} \right) \quad (2)$$

finds a satisfying assignment for  $f(\vec{x})$  if one exists.[8].

### 3.3 Backbone Reduction

To demonstrate the effect of different reductions on various aspects of QA performance, consider a different reduction that we have devised for this paper, and that improves (as we will analyse later) on the reduction given in Eq. (2):

*Definition 3.2* ( $k$ -SAT  $\leq_P$  QUBO (Backbone)). Let  $f(\vec{x})$  be a Boolean function in  $k$ -CNF, and let  $l_{ij}$  be a literal of  $x_j \in \vec{x}$  in  $C_i \in C$ , with  $l_{ij}, x_j \in \{0, 1\}$ . Then

$$q(\vec{x}) = \omega \left( \sum_{l_{ij}, l_{ij'}} l_{ij} l_{ij'} + \sum_{l_{ij}=x_j} -l_{ij} x_j + \sum_{l_{ij} \neq x_j} -l_{ij} + l_{ij} x_j \right) \quad (3)$$

with  $\omega > 0$  describes a QUBO  $q(\vec{x})$  for which  $\min[\vec{x}]$  represents a satisfying assignment of  $f(\vec{x})$  if one exists.

Mathematical details of the derivation are given in Appendix A.

### 3.4 Quality Criteria for Reductions

Quality criteria for software are plentiful, and many of them also apply to the relative merits of reductions. Since the development of quantum computers is mainly driven by the desire for more computational power, we focus on two indicators: Performance and scalability. There is (despite recent [standardisation efforts](#)) no universally applicable (and accepted) definition of how to measure performance of quantum computers; this is particularly hard for QA, where the run-time is not determined by the input, but chosen as a parameter—the annealing time. Consequently, we consider solution *quality*—how likely is it to obtain a correct answer that does not violate constraints, and how accurate is the answer (*i.e.*, how close is it to the optimal achievable value)—as proxy for performance. Scalability considers the question of how large problems can be solved on a hardware of given size (*i.e.*, number of physical qubits).

The achievable accuracy of a reduction depends on its structure (how well do logical connections between qubits match the available physical structure?) and on hardware parameters. While the adiabatic theorem ensures that longer annealing times (runtimes) results in better accuracy, flaws and approximate implementations of the scheme in real hardware lead to less direct relations. Like with traditional approximation algorithms, increasing the amount of computes samples also leads to more accurate solutions.

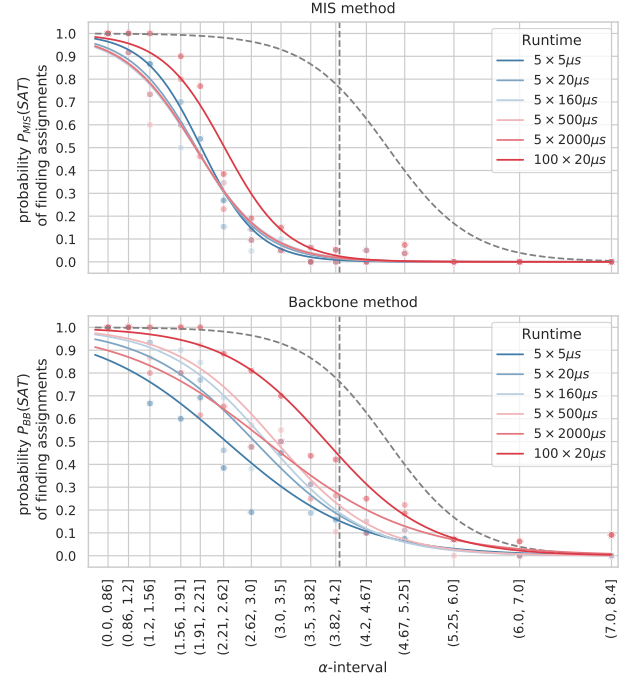
### 3.5 Generating Instance Datasets

Owing to the lack of a published, physically accurate model of the quantum annealer that includes imperfections and noise,<sup>1</sup> determining scalability and accuracy is currently only possible with experimental means [19].

When SAT is used to model constrained optimisation problems in practical applications, the resulting SAT instances often exhibit specific structural properties, which can guide the generation of useful test instances for determining quality properties of reductions. This is, of course, not unlike the well-studied problem of generating tailored input data for general software testing problems [2].

We are interested in a general comparison of reductions, and therefore base our input data generation on general properties of 3-SAT. We have discussed that the problem exhibits different regimes

<sup>1</sup>It is unlikely that such a physically accurate model will be available in the near- or mid-term future.



**Figure 4: Influence of the embedding method on the probability of finding correct satisfying assignments for randomly generated 3-SAT instances with varying ratios  $\alpha$  of clauses to variables. The horizontal dashed lines marks the critical value  $\alpha_c$  (accompanied by a peak increase in required computing time when using traditional numeric solvers). The dashed curve represents the probability distribution of finding a satisfying assignment with optimal solvers. To ease comparing quantum and classical result, a logistic regression curve is given for each parameter variation. Both (mathematically equivalent) methods arrive at correct conclusions less often than classical solvers, which is caused by imperfections and limitations of the available hardware.**

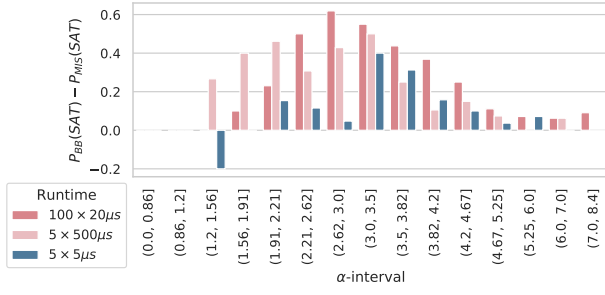
in Section 3.1, and systematically generate random 3-SAT instances that cover these by sweeping across different values of  $\alpha$ . For the number of needed qubits,  $k|C|$  is the dominant term for both reduction approaches. Keeping  $|C|$  fixed and varying  $|V|$  produces stable QUBO sizes across the  $\alpha$ -spectrum, which guarantees a consistent hardware graph utilization.

### 3.6 Experimental Results

We generate a data-set containing 250 random 3-SAT instances with 42 clauses each. In total, six runs with varying annealing times (5 to 2000  $\mu$ s) and samples sizes (5 and 100) were performed on the quantum annealer described in Section 2.3. Figure 4 shows results for the two different reduction methods.

**3.6.1 Accuracy.** For Choi’s MIS-based reduction, the annealing time does not substantially effect the accuracy. Using probability amplification by performing a larger number of runs  $100 \times 20 \mu$ s, does





**Figure 5: Accuracy difference (in percentage points) between MIS and backbone method.**

improve the accuracy slightly. Results obtained with the backbone method, in contrast, improve with increasing annealing time, and increasing the number of runs is also accompanied by a larger improvement as compared to the MIS method. It is also important to note that using an excessively long annealing time of 2000 $\mu$ s results in a *decrease* of result quality<sup>2</sup>.

Recall from the above discussion that solving SAT instances in the critical parameter region around  $\alpha_c$  is most involved for classical solvers, and improvements by quantum computers are most desirable in this region. Unfortunately, the MIS method delivers satisfying solutions in this range with almost zero probability.

Figure 4 directly compares accuracy results. The difference in accuracy reaches up to 60%, and the backbone method is consistently more accurate for all scenarios. The decreasing difference in accuracy at  $\alpha > \alpha_c$  is a consequence of the low number of satisfiable instances in this region. Around the critical region, we observe marked differences of around 35%.

**3.6.2 Scalability.** Figure 6 compares scalability of the two reductions by analysing the amount of required physical qubits, and the mean length of chains necessary to connect qubits without direct physical connections (we use the *minorminer* tool provided as part of the D-Wave API to embed QUBOs into the hardware graph).

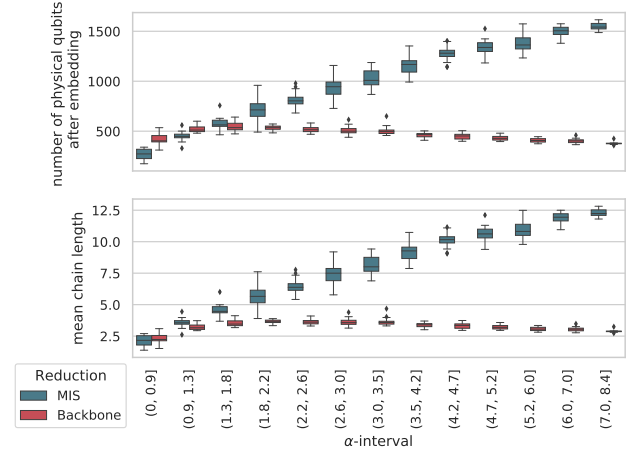
For the MIS-based method, the amount of physical qubits and mean chain length grow essentially linear with an increasing  $\alpha$ , which follows from the pairwise links between conflicting literals.

The backbone method improves upon both aspects because the QUBO is less densely populated, which makes it easier to find embeddings. Especially around the critical value  $\alpha_c$ , the amount of required physical qubits is only half of what is required for the MIS method, which in turn implies that substantially larger problem sets can be solved on a hardware of given size.

## 4 CONCLUSION

Development and evaluation of quantum software components must address well-established engineering concerns of traditional SWE. Based on the scenario of replacing SAT solving, a key element of many applications, with a quantum component, we have

<sup>2</sup>This observation does contradict the adiabatic theorem; the effect is likely caused by a large amount of noise leading incorrect initial configurations or random energy level jumps during the annealing process. Both effects occur with growing probability for increasing annealing times.



**Figure 6: Number of required physical qubits to after embedding a QUBO for a given ratio of variables and clauses (top) and median chain lengths necessary to connect qubits without direct interconnections (bottom).**

shown that careful attention is required in defining and evaluating relevant qualities. We have argued that scalability and accuracy are of particular relevance for early existing quantum annealers. While replacing classical with quantum components is not particularly involved from a programming perspective, our experiments indicate that engineers must be aware of crucial details that might be perceived as irrelevant in traditional SWE to make informed decisions on potentials and pitfalls of quantum computing.

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## A ALTERNATIVE REDUCTIONS

**Loosened Clause Penalties** Choi's reduction (theorem 3.1), is, in essence, a reduction from  $k$ -SAT to the problem of finding the maximal independent set (MIS) of a given graph. Assume a  $k$ -SAT instance is reduced to QUBO as described in theorem 3.1, and let  $G_f$  be the graph representation. Consider a MIS of  $G_f$ , which is given by the largest set of vertices such that there are no connected vertices. In theorem 3.1 this property is enforced by the constraint  $\delta > \omega$ . Solving a QUBO defined by theorem 3.1 also solves the MIS problem for  $G_f$ . The problem of finding the MIS  $G_f$  corresponds to the problem of finding the maximal number of satisfiable clauses (MAX- $k$ -SAT) in  $f$ . The relation between MAX- $k$ -SAT and  $k$ -SAT is trivial.

**THEOREM A.1.** *Setting  $\delta = \omega$  in theorem 3.1 does not change the correctness of the assignment derived from the QUBO solution.*

**PROOF.** Let  $q(x)$  be a sub-QUBO representing one clause like described in theorem 3.1. Under  $\delta = \omega$  the following holds:  $\min(E(n)) = -\omega$  if  $E(n)$  is the energy of a clause sub-QUBO with  $n$  satisfied literals. It is straight forward to see that  $E(n) = -n\omega + \binom{n}{2}\omega$ . Therefore,  $E(0) = 0$  and  $E(1) = E(2) = -\omega$ . The inequality  $E(n) < E(n+1)$  evaluates to  $-n < -1$  which is true for all  $n > 1$ . This leads to the conclusion that  $\min(E(n)) = -\omega$ .

Consider a clause  $C_i$  and its corresponding sub-QUBO  $q_i(x)$ . Now,  $\min(q_i(x)) = -\omega$  for one or two satisfied literals in  $q_i(x)$ . Therefore, the minimization of  $q_i(x)$  leads to a satisfied clause  $C_i$ . For two conflict-free clauses  $C_i$  and  $C_j$  the combined minimum energy is given by  $\min(q_i(x) + q_j(x)) = -2\omega$ . Now we introduce a conflict between  $C_i$  and  $C_j$ . That activates an additional penalty term  $p_{ij} = \omega$  which leads to  $\min(q_i(x) + q_j(x) + p_{ij}) = -\omega > -2\omega$ . This shows that conflicts between clauses always lead to a higher energy level and thus should be avoided when minimizing the complete QUBO  $q(x)$ . For all satisfiable  $k$ -SAT instances  $f(x)$  with  $n$  clauses the minimal energy of their corresponding QUBOs  $q(x)$  will be  $\min(q(x)) = -n\omega$ . Every function  $f(x)$  with minimal QUBO value  $\min(q(x)) > -n\omega$  cannot be satisfied.  $\square$

**Backbone** Choi's reduction represents variables solely by their literals. To avoid conflicts, we need to ensure that  $l_{ij} \neq l'_{ij}$  for all pairs  $(l_{ij}, l'_{ij}) \Leftrightarrow (x_j, \bar{x}_j)$ . An edge in the QUBO connects the literals as penalty term, which leads to highly connected graphs for instances with large values of  $\alpha$ . The degree of connectivity can be improved by introducing a *backbone* for variables, which allows us to transitively express equivalence between literals by linking them to their corresponding variable. The reduction is given in Definition 3.2 on page 4.

**CORRECTNESS OF THEOREM 3.2.** Let  $E(n)$  be the energy of a clause sub-QUBO with  $n$  satisfied literals. The difference between sub-QUBOs in theorem 3.1 and theorem 3.2 is that in the latter node weights of literals  $l_{ij} \neq x_j$  are moved to the edges  $(l_{ij}, x_j)$ . For every literal, there exists exactly one edge to its corresponding variable. Therefore, edge weights can be viewed as node weights, and it follows that  $E(n) = -n\omega + \binom{n}{2}\omega$ . Consequently,  $\min(E(n)) = -\omega$

also holds for theorem 3.2. If two literals  $l_{ij}$  and  $l'_{i'j}$  conflict, one of  $\omega l_{ij}x_j$  or  $\omega l'_{i'j}x_j$  evaluates to  $\omega$ , while the other evaluates to 0. The rest of the argument follows Theorem A.1.  $\square$