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#### Full length article

# Evaluating the potential of quantum machine learning in cybersecurity: A case-study on PCA-based intrusion detection systems

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#### ARTICLE INFO

# Keywords: Quantum computing Quantum machine learning QML Evaluation Framework Impact PCA Principal component analysis Network intrusion detection

#### ABSTRACT

Quantum computing promises to revolutionize our understanding of the limits of computation, and its implications in cryptography have long been evident. Today, cryptographers are actively devising post-quantum solutions to counter the threats posed by quantum-enabled adversaries. Meanwhile, quantum scientists are innovating quantum protocols to empower defenders. However, the broader impact of quantum computing and quantum machine learning (QML) on other cybersecurity domains still needs to be explored. In this work, we investigate the potential impact of QML on cybersecurity applications of traditional ML. First, we explore the potential advantages of quantum computing in machine learning problems specifically related to cybersecurity. Then, we describe a methodology to quantify the future impact of fault-tolerant QML algorithms on real-world problems. As a case study, we apply our approach to standard methods and datasets in network intrusion detection, one of the most studied applications of machine learning in cybersecurity. Our results provide insight into the conditions for obtaining a quantum advantage and the need for future quantum hardware and software advancements.

#### 1. Introduction

Network security

Quantum computing combines concepts from computer science, mathematics, and physics to provide a novel computational model. A *quantum computer* is a programmable physical system that obeys the laws of quantum physics. Writing code for a quantum computer means specifying the evolution of a quantum mechanical system such that the system's final description encodes the output of the computation. Expected to expand the way we process information radically, quantum computing provides new primitives unavailable in classical information processing.

The impact of quantum computing extends across a wide range of domains. Quantum simulation, for example, allows researchers to model complex quantum systems that are intractable for classical computers, with significant implications for material science, chemistry, and fundamental physics. Additionally, quantum computing is driving the development of the quantum internet, which aims to utilize quantum entanglement for ultra-secure communication. The emerging field of quantum software engineering focuses on creating tools, languages,

and frameworks to efficiently design and optimize quantum algorithms, ensuring practical and scalable implementations.

In cryptography, the influence of quantum computing has been particularly profound. In 1994, Peter Shor published an algorithm for factoring integers and solving the discrete logarithm problem in polynomial time (Shor, 1994), mining the security of current asymmetric cryptography. Additional algorithmic primitives potentially threaten the security parameters of symmetric cryptography and hashing algorithms. These developments profoundly impacted cryptography, leading to the dawn of post-quantum cryptography (Bernstein and Lange, 2017), a research line striving to devise cryptographic primitives and protocols resilient to quantum-enabled attackers. Concurrently, quantum scientists are developing protocols to aid quantum-enabled defenders (Pirandola et al., 2020). Cryptography, however, is not the sole domain expected to transform with the advent of practical quantum computing.

Another promising field is machine learning (ML), where quantum computers are expected to enable processing more extensive and complex datasets, potentially leading to more accurate and efficient algorithms. The interest in quantum machine learning (QML) can be traced back to 2009 when Harrow et al. (2009) proposed a quantum algorithm

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to solve linear systems in time that depends only poly-logarithmically on the system's size and polynomially on other parameters. This result paved the way for new quantum linear algebra that proved very useful in ML (Biamonte et al., 2017).

Simultaneously, classical<sup>1</sup> machine learning has been successfully applied to a wide range of cybersecurity tasks, from malware and binary analysis (Ucci et al., 2019; Rieck et al., 2011) to detecting network intrusions (Buczak and Guven, 2015) and financial frauds (Carminati et al., 2018). As ML algorithms become integral to both defenders' and attackers' arsenals for securing (Arp et al., 2022) and attacking (Godefroid et al., 2017; Yamaguchi et al., 2011) computer systems, the potential impact of QML on cybersecurity becomes a natural question.

Recent literature has begun exploring the performances of nearterm heuristic QML algorithms for computer security problems, such as denial of service or botnet detection. However, the absence of theoretical guarantees for these near-term heuristics and the limitations of current hardware and simulators hinder researchers from predicting the scaling and performances of these algorithms on realistic datasets. The research question of how QML can impact cybersecurity in the long run when fault-tolerant quantum computers are available has been hardly explored in any previous literature to our knowledge.

In this work, we address this question by presenting a theoretical analysis of the potential advantages of quantum computing in ML. We build a simple framework to identify the conditions under which a QML algorithm for a fault-tolerant quantum computer outperforms a classical one. The goal of the framework is twofold:

- Study how the errors of QML algorithms which are a particular kind of randomized approximation algorithms affect the performance of quantum models with respect to their classical counterparts;
- Compare the running times of quantum and classical ML models to discover the settings where quantum algorithms provide an advantage.

This framework enables cybersecurity experts and practitioners to assess whether quantum machine learning algorithms can effectively address specific practical problems. It provides a benchmarking tool for fair comparisons of future QML solutions in the cybersecurity domain, which will surely be proposed in the next few years.

As a demonstrative case study, we apply the methodology to PCA-based quantum machine learning models for network intrusion detection, a typical ML application in cybersecurity. In particular, we study how the algorithmic errors introduced during the QML training affect the performance of intrusion detection models on real datasets. To do so, we implement and test a numerical simulation of essential quantum subroutines — amplitude estimation, phase estimation, and tomography of pure states. These subroutines are critical building blocks that enable simulations of many existing QML algorithms. In our case study, we evaluate the performances of quantum clustering (q-means Kerenidis et al., 2019) and quantum dimensionality reduction (QPCA Bellante et al., 2022) for intrusion detection.

In summary, this paper contributes to elucidate the potential impact of quantum machine learning on cybersecurity by providing a theoretical framework for assessing the advantages of quantum algorithms over classical ones, as well as conducting a case study on PCA-based quantum algorithms for network intrusion detection, shedding light on their performance and feasibility in real-world scenarios.

The manuscript is structured as follows. In Section 2, we discuss the importance of machine learning in cybersecurity, the recent literature on near-term heuristic-based QML, and describe the opportunities for QML algorithms in the fault-tolerant regime. Section 3 provides readers with the necessary quantum background to understand the proposed

framework. Section 4 presents the evaluation framework. Finally, in Section 5, we present the case study on PCA-based quantum algorithms for network intrusion detection.

#### 2. Quantum machine learning algorithms for computer security

In the rapidly evolving landscape of computer security, the intersection of quantum computing and machine learning has emerged as a frontier with transformative potential. As we witness the advancements in quantum technologies, researchers have embarked on developing quantum machine learning (QML) algorithms that could find cybersecurity applications. This section explores the connections between computer security and quantum machine learning, discussing the current landscape and the promising horizons within the fault-tolerant regime.

#### 2.1. Machine learning for cybersecurity

Machine learning algorithms have found widespread success in cybersecurity applications (Arp et al., 2022; Ucci et al., 2019; Sabau, 2012).

In the unsupervised learning domain, Principal Component Analysis (PCA) stands as an indispensable tool for preprocessing and analyzing large, complex datasets. Its applications span anomaly and intrusion detection, feature selection, and privacy protection. In anomaly detection (Brauckhoff et al., 2009; Ringberg et al., 2007), PCA helps identify unusual patterns in network traffic, system logs, or user behavior, thereby making it easier to detect deviations that may signal cyberattacks or malicious activities. For Intrusion Detection Systems (Sultana et al., 2019) (IDS) and Malware Detection, PCA aids in analyzing network traffic data and malware samples, respectively, and reduces the feature space to preserve the most pertinent information. Regarding feature selection in machine learning-based cybersecurity applications (Vasan and Surendiran, 2016), PCA selects relevant features, discarding less significant ones and enhancing the model's efficiency and accuracy. Additionally, the application of clustering algorithms (e.g., Hierarchical, k-means, and spectral clustering) has earned significant attention as an effective tool for identifying patterns and anomalies within large datasets of security-relevant phenomena like network intrusions (Horng et al., 2011; Jianliang et al., 2009; Ma et al., 2016) and malware (Pai et al., 2017).

At the same time, supervised learning techniques such as Support Vector Machines (SVMs), linear regression, Neural Networks (NNs), and Convolutional Neural Networks (CNNs) are used for multiple tasks. SVMs find use in malware classification (Zhao et al., 2011) and intrusion detection (Mukkamala et al., 2002), as they can handle high-dimensional data and can segregate malicious activity from normal through hyperplane separation in feature space. Linear regression, though traditionally employed in predictive modeling, can also forecast the behavior of security metrics over time, thereby aiding proactive threat management. Neural Networks are used, for instance, in phishing detection (Mohammad et al., 2014) and vulnerability identification (Wu et al., 2017), thanks to their ability to learn complex patterns in data. Convolutional Neural Networks have proven to be particularly effective in malware analysis (Kalash et al., 2018; Vasan et al., 2020).

As machine learning establishes itself as a pivotal tool in cybersecurity, the potential applications of quantum machine learning (QML) become increasingly evident. Accelerating computational processes through quantum computing has the potential to significantly enhance security measures, particularly in solving complex problems such as the analysis of massive-scale datasets, symbolic execution-based malware analysis, and automated vulnerability discovery through fuzzing.

 $<sup>^{\</sup>rm 1}\,$  In this manuscript, the word "classical" means non-quantum.

#### 2.2. Current research trends

In recent years, researchers have started exploring applying quantum machine learning to cybersecurity problems. Most ongoing applied research focuses on designing and testing QML algorithms for NISQ (Noisy Intermediate-scale quantum) devices — small, non-error-corrected quantum computers. This research line expects to ease the hardware requirements imposed by error-corrected architectures and start benefiting from the first quantum computers. The main idea is to program the quantum computer with parameterized gates, akin to classical neural networks, and optimize the circuit parameters using a classical computer. For a comprehensive review of the subject, we refer to Bharti et al. (2022).

Along this research line, Kalinin and Krundyshev (Kalinin and Krundyshev, 2023) evaluated the application of Quantum SVM and CNN to intrusion detection. Suryotrisongko and Musashi (Suryotrisongko and Musashi, 2022) investigated variational quantum algorithms for botnet detection based on domain generation algorithms (DGA). Payares and Martínez-Santos (Payares and Martínez-Santos, 2021) discuss applications of QML using variational algorithms for SVM, Neural Networks, and an ensemble to detect denial of service (DOS) attacks. Masum et al. (2022) use variational SVM and Neural Networks to investigate supply chain attacks. Beaudoin et al. (2022) investigate using variational algorithms for quantum LSTM and NN to detect Trojan-infected circuits.

Despite the simplicity offered by these variational algorithms, obtaining provable guarantees about their performances remains challenging. Some recent results fuel the conjecture that it will be hard for parameterized quantum circuits to outperform significantly and consistently classical algorithms in many relevant areas (Aharonov et al., 2023). Moreover, the lack of theoretical guarantees makes quantum simulators and hardware crucial for evaluating their impact on practical applications, restricting the dimensions of the datasets on which it is possible to run experiments. We believe that the proper test bench of these quantum algorithms will be when NISQ-scale quantum computers are available. The same difficulties stand for quantum annealing and machine learning approached through QUBO formulations (Arthur et al., 2021). For these reasons, we focus on QML algorithms with a provable speedup over their classical counterparts.

#### 2.3. Opportunities in the fault-tolerant regime

Besides NISQ and QUBO algorithms, researchers have been investigating what fault-tolerant quantum computers equipped with a classically writable memory can do for machine learning. Leveraging techniques from quantum linear algebra, researchers developed "quantized" versions of emblematic machine learning algorithms (Biamonte et al., 2017). They aim to reduce the running time requirements of the classical counterparts to process data in shorter times and analyze more data with the same computational power. These algorithms come with theorems and analyses that bound their running times. Examples of fault-tolerant machine learning algorithms with provable running times are the following.

Unsupervised learning. There are several algorithms for unsupervised learning. Among the many, we cite quantum algorithms for hierarchical clustering (Aïmeur et al., 2013), spectral clustering (Kerenidis and Landman, 2021), nearest neighbor (Wiebe et al., 2015; Lloyd et al., 2013; Bellante et al., 2023), k-means (Kerenidis et al., 2019), Gaussian mixture models (Kerenidis et al., 2020b), quantum algorithms for PCA ed eigenvalue-based techniques (Bellante et al., 2022), and for learning sparse representations (Bellante and Zanero, 2022).

**Supervised learning.** The literature on supervised algorithms is equally proceeding. For instance, we have quantum algorithms for Support Vector Machines (Rebentrost et al., 2014), linear regression (Chakraborty et al., 2023), slow feature

analysis (Kerenidis and Luongo, 2020), and many others. There are also some quantum algorithms for deep learning, like quantum neural networks, convolutional neural networks, and perceptron algorithms (Roget et al., 2022; Liao et al., 2021; Kerenidis et al., 2020a).

Most of these algorithms have the following important characteristic: once provided quantum access to the data, the complexity of the quantum algorithms depends only polylogarithmically on the number of data points. This starkly contrasts with classical algorithms, where the dependence is at least linear. The theoretical guarantees that come with analyzing these algorithms make them suitable for evaluating the impact of quantum computing in the future, allowing researchers to investigate their use on realistic datasets without the need for quantum simulators or hardware (e.g., see experiments in Bellante et al. (2022), Kerenidis and Luongo (2020), Dalzell et al. (2023)). For this reason, we will focus on evaluating the impact of quantum machine learning algorithms in the fault-tolerant regime. In the reminder, we use QML to refer to quantum machine learning algorithms with provable guarantees in fault-tolerant settings.

#### 2.4. Motivation and relevance to cybersecurity

The cybersecurity field faces growing challenges from increasingly sophisticated threats, requiring continual advancements in detection and mitigation techniques. While traditional machine learning algorithms have demonstrated considerable success in areas such as intrusion detection, anomaly detection, and malware analysis, these methods are increasingly strained by the scale and complexity of modern security problems. Quantum machine learning offers a potential pathway to overcoming these limitations by enabling computational speedups and enhanced algorithms that could significantly improve cybersecurity defenses.

The primary motivation for this work is to explore the future implications of fault-tolerant quantum machine learning algorithms in cybersecurity applications. To date, much of the research in quantum computing for cybersecurity has centered around NISQ devices, which, while promising, are constrained by hardware limitations and lack theoretical guarantees. These limitations hinder their scalability and applicability to large, real-world datasets, such as those commonly used in cybersecurity. Our approach is different: we focus on QML algorithms designed for fault-tolerant quantum computers, which hold the potential to overcome these constraints and deliver provable performance guarantees. In this paper, we define a quantum algorithm as advantageous if it runs faster than the classical version while still delivering comparable results. However, the exact conditions under which quantum algorithms outperform their classical counterparts remain uncertain, particularly in practical cybersecurity applications. This uncertainty presents both a challenge and an opportunity for research.

Our work aims to address this gap by providing an evaluation framework that allows cybersecurity experts to systematically assess the potential advantages of QML algorithms in a fault-tolerant setting. We also offer a case study on PCA-based anomaly detection for network intrusion that illustrates how such a framework can be applied to standard machine learning tasks, offering insights into when and how quantum algorithms may achieve a meaningful advantage. While the case study serves to showcase the framework, the broader contribution lies in the methodology, which can be adapted to various cybersecurity domains and tasks. By focusing on fault-tolerant quantum computing and algorithms with provable guarantees, this paper establishes a foundation for future research that will be relevant as quantum hardware advances. The work not only highlights the potential impact of QML on cybersecurity but also equips practitioners with practical tools to evaluate this impact, ensuring that the cybersecurity community is prepared to harness the capabilities of quantum computing when the technology matures. In summary, the motivation behind this research

is twofold: first, to investigate the theoretical underpinnings and performance benefits of fault-tolerant QML algorithms in cybersecurity; and second, to provide a robust framework for practitioners to assess the practical feasibility and scalability of these algorithms as quantum computing technology continues to evolve.

#### 3. Quantum computing background

This section provides an overview of the key quantum computing concepts essential for evaluating fault-tolerant quantum machine learning (QML) algorithms. Readers seeking a brief introduction to quantum computing are directed to Appendix A, while a comprehensive one is available in Nielsen and Chuang (2010).

**Qubits and registers.** A qubit, or quantum bit, is the fundamental unit of quantum information. Differently from classical bits, qubits can exist in a superposition of  $|0\rangle$  and  $|1\rangle$  states and collapse to one of these states when measured. The state of a single qubit is described by a complex vector with two entries, from which it is possible to compute the probability of collapsing in one state or another. The state of multiple qubits can be described by a complex vector of size exponential in the number of qubits, and qubits can be correlated.

**Quantum algorithm.** A quantum algorithm is a computational procedure that consists of two key processes: evolving and measuring a specific initial quantum state multiple times. The evolution process involves applying a series of quantum operations or gates to the initial quantum state. These operations change the state of qubits in a controlled manner. Instead, measurement collapses the quantum state from its superposition of outcomes to a definite state, yielding the outcome of the computation. The process of reconstructing a quantum state requires statistics over several measurements and is called *tomography*. To perform tomography, one must re-create the quantum state by repeating the algorithms from scratch multiple times (e.g., Theorem B.4).

**Notation.** We adopt Dirac's notation, denoting a complex vector  $\mathbf{x} \in \mathbb{C}^n$  as  $|x\rangle$ , for some  $n \in \mathbb{N}$ , and its transposed complex conjugate  $x^\dagger$  as  $\langle x|$ . Tensor products  $|x\rangle \otimes |y\rangle$  are abbreviated as  $|x\rangle |y\rangle$ . For a matrix  $\mathbf{X} \in \mathbb{R}^{n\times d}$ ,  $\|\mathbf{X}\|$  represents the spectral norm (the greatest singular value),  $\kappa(\mathbf{X})$  denotes the condition number (the ratio between the greatest and the smallest singular values), and  $\mu(\mathbf{X})$  is a parameter bounded by the Frobenius norm (i.e.,  $\mu(\mathbf{X}) \leq \|\mathbf{X}\|_F = \sqrt{\sum_{ij} X_{ij}^2}$ , see Definition B.3 in Appendix B). Big-O notation is used for algorithmic complexity, with  $\widetilde{O}$  omitting polylogarithmic terms.

#### 3.1. Mapping quantum software on hardware

At the time of writing, numerous research institutions and companies invest considerable resources in developing quantum computers, with some prototypes already available in the cloud. Current quantum computers have qubits in the order of hundreds and can execute gates in the order of thousands. These prototypes have neither enough qubits nor enough quantum coherence to run any useful computation with a provable advantage over classical computers for problems of real interest. Nevertheless, in the past years, some hardware architectures have successfully solved computational problems of purely theoretical interest (i.e., with no practical application) faster than any classical computer.

When a quantum hardware platform can execute a task that is computationally infeasible for classical computers, it is said to have achieved quantum supremacy or quantum advantage (Madsen et al., 2022; Preskill, 2012). It is reasonable to expect that in the next decades, quantum computers will be mature enough to execute software for more relevant problems. Interested readers are referred to Saffman et al. (2010), Bruzewicz et al. (2019), Kjaergaard et al. (2020) for an overview of scalable quantum computer architectures. While quantum

architectures are expected to become faster and more reliable over time, they are not projected to surpass classical architectures in clock time (Babbush et al., 2021). Consequently, considerable speedups must come from algorithmic improvements in the number of operations executed.

In the reminder, we describe the steps to consider when compiling quantum software into quantum hardware. All these steps add some overhead that needs to be considered when estimating the wall-clock time of a quantum algorithm.

**Programming.** From a programmatic standpoint, conceptualizing quantum computers is akin to thinking of an FPGA, where the circuit is described for mapping and execution on the device. To facilitate the development of quantum software, quantum computers can be programmed in high-level languages. Although these languages are less advanced than the ones we can use for classical hardware, they facilitate specifying abstract gates and circuits (e.g., arbitrary qubit rotations, adders, multipliers) and common subroutines such as quantum Fourier transform and amplitude amplification and estimation. Often, the code generated by these high-level languages is not optimized for the task or the target hardware.

Compilation. Quantum hardware platforms typically execute only a basic set of universal quantum gates, i.e., gates that can be combined to build any other circuit. These gate sets often vary from architecture to architecture. During compilation, high-level instructions are decomposed into sequences of these gates. The Solovay–Kitaev theorem enables efficient transpilation of quantum software between architectures, though this process can introduce some overhead in the total number of gates (Maronese et al., 2022; Bouland and Giurgica-Tiron, 2021). Optimization techniques based on heuristics are available (Duncan et al., 2020) and can help reduce the circuit size and depth. Besides dealing with circuit decomposition and optimization, compilation will also need to take into account error correction and connectivity constraints.

Error correction. The primary challenge in achieving large-scale quantum computers is constructing noise-resilient qubits. Quantum Error Correction Codes (QECC) are a set of methods that help protect quantum information from noise. They usually embed one logical qubit in a bigger Hilbert space spanned by many physical qubits and work by detecting errors and applying corrective gates at runtime. QECC introduces overhead in terms of qubit quantity, number of operations, and classical interaction needed to decode the errors and control the quantum processing unit accordingly. With current technology, this overhead is such that asymptotic quadratic speedups are believed insufficient for practical scenarios (Babbush et al., 2021).

Connectivity. Another factor influencing compilation and effective quantum algorithm running time is hardware connectivity. In fact, some architectures limit connectivity to interactions between physically proximate qubits. To overcome this limitation, qubits must be swapped along the circuit using intermediate qubits to build long-range interactions. This introduces an overhead in the number of operations. During compilation, logical qubits are assigned to physical qubits to minimize long-range interactions, as connectivity constraints may limit circuit parallelization.

#### 3.2. Complexity of quantum algorithms

There are different complexity measures for quantum algorithms. As in the study of classical algorithm complexity, at a theoretical level, we are interested in the *asymptotic* scaling in the problem parameters.

The *query complexity* of an algorithm is the number of calls made to an *oracle* providing access to the problem's input. This measure – which is also standard in classical computer science – conceals the oracle's implementation cost and the processing cost between any two oracle calls. One of the reasons for its adoption in quantum (and classical) computer science is the use of techniques like the polynomial and

#### Example 1 - Query advantage $\Rightarrow$ gate advantage.

Consider a quantum application targeting a preimage attack scenario, where an oracle provides access to a hash function (e.g., SHA256), comparing the output with a target hash image. The brute-force query complexity, evaluated in terms of hash function calls, is on the order of  $O(2^m)$  for an m-bit hash function. Using Grover's search algorithm (Grover, 1997) with a quantum computer, a quantum computer reduces this complexity to  $O(2^{m/2})$ . Notably, this result only suggests a reduction in the number of hash function evaluations. However, in this case, an advantage in query complexity translates into an advantage in gate complexity, as the costs of the quantum and classical oracles are comparable and vastly dominate the cost of intermediate operations between oracle calls. To express the quantum cost in terms of time complexity, one must delve into the implementation details of the oracle (i.e., a quantum circuit for SHA256) and consider the overhead for error correction. An analysis of the physical resources (the number of qubits and the actual number of gates) and wallclock time estimation, with surface error correction codes on planar connectivity, can be found in Amy et al. (2016), Aggarwal et al. (2018).

adversarial methods to prove lower bounds in this model (Ambainis, 2000; Beals et al., 2001).

The *gate complexity* denotes the total number of one and two-qubit gates executed by the quantum circuit implementing the algorithm. This choice stems from the fact that any multi-qubit gate can be decomposed into a sequence of one- and two-qubit that form a universal gate set and that the asymptotic behavior of the gate complexity remains unaffected by the choice of the specific gate set. This complexity measure is the quantum equivalent of a classical Boolean circuit's *gate complexity* or the *circuit-size complexity*. Given the query complexity and a gate decomposition of both the oracle and the intermediate operations between the oracle calls, one can determine the gate complexity.

The *depth* of a quantum algorithm mirrors the classical *circuit-depth complexity* of a Boolean circuit. It is the length of the longest sequence of gates from the input to the output of a quantum circuit. Typically measured before circuit compilation on specific hardware architecture, it does not consider Quantum Error Correction (QECC) or connectivity issues. Depth complexity offers insight into the degree of parallelization achievable within a quantum algorithm.

The time complexity gauges the wall-clock time required to execute the algorithm on specific hardware. Unlike the previously mentioned asymptotic complexity measures, which abstract away implementation details, this metric reflects the actual execution time of a quantum algorithm on the chosen hardware platform, which may impose constraints on parallelism, available error-correcting codes, and compilation techniques. An estimate of the time complexity can be derived from the query complexity, along with the details on the implementation of the oracles and the details of the hardware architecture (such as compilation, error correction, connectivity, and parallelization). The process of estimating the time complexity of a quantum algorithm by expanding the oracle, optimizing the circuit, and considering all the hardware and technological constraints is called *resource estimation*.

We clarify the previous definitions with two simple examples. In Example 1, we observe that query complexity serves as a reliable measure to assess the efficiency of a quantum algorithm, as the advantages readily extend to gate and time complexity. Contrastingly, Example 2 emphasizes the need for caution when relying solely on query complexity. The quantum algorithm for the hidden subgroup problem exhibits a stark difference between query and gate complexity, challenging the assumption that reducing query complexity guarantees efficiency gains in other aspects.

Researchers commonly express the complexity of fault-tolerant quantum machine learning algorithms in terms of queries to input oracles. Analogous to Example 1, in QML, the negligible cost of intermediate operations between oracle calls often positions memory accesses as the bottleneck. This aligns with the understanding that, in quantum machine learning, query complexity remains a pertinent metric, offering valuable insights into the algorithm's efficiency.

#### 3.3. Classical data and quantum computers

To process classical data in a quantum computer, we must consider input and output operations, which we survey in this section.

**Input.** Along with a quantum computer, it is common to assume the availability of a classically-writable quantumly-readable quantum random access memory (QRAM). Here, a QRAM is a memory that can be written with a classical device and responds to queries in superposition from a quantum device. A memory  $[m_0, \ldots m_i, \ldots m_{M-1}]$  is an indexed array of size M, with entries encoded in p bits. A quantum memory is a unitary of the kind

$$U_{ORAM}: |i\rangle |0\rangle \mapsto |i\rangle |m_i\rangle,$$
 (1)

which maps the *i*th address to the memory entry  $m_i$ .

Some algorithms may require the binary encoding of  $m_i$  in a register of qubits, as produced by the unitary in Eq. (1). Others may need to encode the entries in the amplitudes of a quantum state  $|m\rangle = \frac{1}{\|m\|} \sum_{i=0}^{M-1} m_i |i\rangle$ . Others again require access to a unitary U (a circuit) such that  $\|A - \alpha(\langle 0|^{\otimes q} \otimes I)U(|0\rangle^{\otimes q} \otimes I)\| \leq \epsilon$ , where A is a matrix representation of a portion of the memory (i.e.,  $(\alpha, q, \epsilon)$ -block-encoding access to a matrix Low and Chuang, 2019). Efficient access to all these data representations can be achieved with a QRAM oracle, as described in Eq. (1).

While a basic multiplexer circuit can implement the mapping in Eq. (1), it would have a linear depth in memory size O(Mp). There exist more efficient circuit implementations for creating access to sparse data structures, such as sparse vectors or matrices (Camps et al., 2024; Matteo et al., 2020). Fortunately, better general architectures for implementing Eq. (1) exist. For instance, the bucket-brigade architecture maintains a total circuit size of O(Mp) gates but has a depth only logarithmic in the memory size (Giovannetti et al., 2008) and is quite resilient to generic errors (Hann et al., 2021). Although the first log-depth QRAM architecture was presented in 2008, quantum random access memories only recently started to become an active topic in the research community. For instance, different research proposals exist for implementing this architecture on a different kind of quantum hardware (Hann et al., 2019) and in data centers (Liu et al., 2022).

We have *efficient access to a quantum memory* if the mapping in Eq. (1) can be performed in time  $O(\text{poly}(\log{(M)}, p))$ .

**Output.** To retrieve data out of a quantum computer, the possibilities are more narrow. In QML, the output is often encoded in the amplitudes of a quantum state that the algorithm produces. Subroutines like amplitude estimation (Theorem B.6) can be used for a single-number output. In cases where the output is a vector, one can use quantum tomography of pure states (Theorem B.4). Quantum pure-state tomography enables the reconstruction of a quantum state by sampling it. Given a quantum state  $|x\rangle$ , the tomography algorithm outputs an estimate  $\overline{x}$  such that  $||x-\overline{x}||_2 \le \delta$  by performing  $N = \widetilde{O}\left(\frac{d}{\delta^2}\right)^2$  measurements (Kerenidis and Prakash, 2020b).

 $<sup>^2</sup>$  Note that tomography subroutines have been recently improved in general settings, offering a further quadratic speedup in the precision  $\delta$  (Apeldoorn et al., 2023).

#### Example 2 - Query advantage ⇒ gate advantage.

An illustrative example highlighting a significant disparity between query complexity and gate complexity is the quantum algorithm for the hidden subgroup problem (Ettinger et al., 2004). In this case, the algorithm requires only a polynomial number of queries to the function oracle, seemingly indicating efficiency. However, the number of gates between successive calls results in an exponential gate complexity. For this algorithm, query complexity is not a good proxy for time complexity.

#### 4. Evaluating the quantum advantage

This section introduces a framework designed to assess the quantum advantage in machine learning applications. We focus on cybersecurity problems. However, the methodology is general and can be adapted for evaluating the impact of QML algorithms in various fields. First, we analyze fault-tolerant quantum machine learning algorithms, emphasizing the challenges posed by the data loading problem and the complexities in estimating their running times on real-world datasets. Subsequently, we detail the methodology framework, discussing its merits and limitations.

#### 4.1. Fault-tolerant quantum machine learning

We discuss the caveats arising from the cost of loading classical data into a quantum device and the challenges associated with evaluating the complexity and impact of a quantum machine learning algorithm.

#### 4.1.1. The cost of data loading

Quantum machine learning algorithms can process both native quantum data, originating from controllable quantum algorithms and processes, and classical data, derived from conventional sensors and machinery. When assessing the cost of processing classical data, one must also consider the expenses associated with compiling the unitaries that provide input to the data.

In applying QML to contemporary cybersecurity challenges, like network intrusion detection or malware analysis, the criticality of the classical data loading cost onto quantum devices becomes evident. In some cases, refraining from considering this cost by assuming that the data is pre-stored in some quantum-accessible memory may lead to inaccurate assessments of the quantum algorithms' efficiency, as emphasized by Aaronson (Aaronson, 2015).

QML algorithms capitalize on the ability to store a size d vector using only  $\lceil \log d \rceil$  qubits and an  $N \times d$  matrix using  $\lceil \log N \rceil + \lceil \log d \rceil$  qubits. Amplitude encodings or block-encodings facilitate this storage. With efficient quantum memory access (Eq. (1)) and some dataset preprocessing, these encodings can be generated in polylogarithmic time in the matrix dimensions. Detailed methods for leveraging QRAM for this purpose are outlined in Appendix B.

It is important to highlight that these encodings necessitate preprocessing the input data on a classical device. This step consists of preparing a specific data structure to store in the quantum memory rather than the raw matrices or vectors. The preprocessing time for a  $N\times d$  matrix is  $\widetilde{O}(Nd)$  and is executed only once during loading. This process is highly parallelizable, efficient to update  $(\widetilde{O}(1))$  per matrix entry), and is an essential factor in evaluating the quantum algorithm's speed compared to classical methods.

Considering this preprocessing input step, the expected exponential speedup of the quantum procedure is often lost, as one needs to pay a linear time data loading (Aaronson, 2015). As an example, this data loading cost implies that quantum algorithms may not significantly expedite the inference phase of classical machine learning models whose prediction time is linear in the number of features. In such cases, the time required to input the test data point into a quantum memory would be comparable to the prediction time itself. Nevertheless, in situations where a model's training or inference phase incurs a polynomial time cost, quantum algorithms could still offer a comparative polynomial advantage over classical alternatives.

#### 4.1.2. Complexity evaluation

The complexity of fault-tolerant quantum machine learning algorithms is often expressed in terms of queries to a quantum memory or in the total number of gates to execute considering efficient access to a quantum memory (i.e., polylogarithmic, similar to the memory access cost in the classical RAM model). Similar to Example 1 - unlike Example 2 - in QML algorithms, these two costs are equivalent up to polylogarithmic factors. This equivalence makes the QRAM query complexity a robust metric for quantifying the number of simple operations required. We leverage this fact in our evaluation methodology, enabling decisions based on query complexity before executing detailed resource estimations. The QML algorithms considered here are randomized approximation algorithms. Their running times depend on some probability of failure  $\gamma$  and approximation error  $\epsilon$ . It is possible to trade off these parameters to expedite the algorithm's execution at the expense of performance and reliability. A cybersecurity expert might want to tailor the amount of tolerable error and failure probability to the problem they are trying to solve, striking the best trade-off between time efficiency and accuracy/reliability.

While theorems provide asymptotic query complexity for QML algorithms, comparing this complexity to classical algorithms is not straightforward. Indeed, quantum and classical running times often depend on different dataset parameters. For instance, a classical algorithm for machine learning might depend solely on the dataset's size. In contrast, the query complexity of its quantum counterpart might depend on the effective condition number  $\kappa(X)$ , some approximation error  $\epsilon$ , a failure probability  $\delta$ , one choice of  $\mu(X)$  (Definition B.3), and other dataset-dependent parameters. These dataset-dependent parameters are critical in evaluating regimes in which the quantum algorithm can exhibit advantages over its classical counterparts (see Examples 3 and 4).

**Failure probability.** QML algorithms fail with a probability smaller than  $\gamma > 0$ , similar to classical randomized algorithms. This probability can often be minimized by incurring in a multiplicative cost in the running time of  $O(\log{(1/\gamma)})$ , which is negligible in practice. Once the cybersecurity expert fixes the tolerable failure probability based on the application, this factor accounts for a constant in the total running time.

**Approximation error.** The output of QML algorithms approximates the output of corresponding classical subroutines. For a vector output  $\overline{s}$ , we consider an  $\ell_2$  or  $\ell_\infty$  approximation error over the classical output s, i.e., a vector  $\overline{s}$  such that  $\|s-\overline{s}\|_2 \le \epsilon$  or  $\|s-\overline{s}\|_\infty \le \epsilon$ . In the case of a scalar a, we consider the absolute or relative error between the classical and the quantum output, i.e.,  $\|a-\overline{a}\| \le \epsilon$  or  $\|a-\overline{a}\| \le \epsilon a$ .

Other dataset-dependent parameters. Several parameters impact QML algorithms' performance, such as the maximum norm of vectors in the dataset (see Theorem B.7, or  $\eta$  in Theorem B.10, or  $s_q$  in Definition B.3), the sparsity of the matrix, or a threshold  $\theta \in (0, \|X\|]$  for picking the chosen number of principal components (Theorem B.9). Another critical quantity is the condition number of the dataset matrix. In many real-world applications, matrices are not full-rank, meaning some small singular values are zero or nearly zero, potentially resulting in an unbounded condition number. Discarding singular values below a threshold reduces the effective condition number, potentially enhancing numerical stability and algorithm performance. The threshold depends on the dataset, the algorithm, and the application, and its optimization can benefit the numerical stability and the performances of the (Q)ML algorithm, as it can help regularize the model.

#### Example 3 - Approximation error vs Running time.

Consider an anomaly detection system based on Euclidean distances, where a test vector  $\vec{x}_t$  is flagged as an anomaly if it falls within a radius r of vector  $\vec{a}$ , i.e.,  $\|\vec{x}_t - \vec{a}\|_2 \le r$ . For vectors of d features, an exact and deterministic classical classifier would require O(d) operations. On the other hand, a quantum classifier that uses Theorem B.7 can evaluate the distance D with an error  $\epsilon$  (i.e., estimates  $\overline{D} \in [D - \epsilon, D + \epsilon]$  with high probability) with query complexity and extra number of gates scaling proportionately with the inverse error  $(\widetilde{O}(\frac{1}{\epsilon}))$ . Determining a suitable value for  $\epsilon$  involves striking a balance between speed and classification accuracy. If the error is too big, we risk misclassifying anomalous vectors, e.g.,  $D \le r$  but  $D + \epsilon > r$ . On the other hand, if the required error is too small, we end up with a quantum algorithm slower than its classical counterpart. In conclusion, the quantum algorithm seems advantageous whenever the problem tolerates an error that is larger than the inverse number of features ( $\epsilon \in \Omega(1/d)$ ). Conversely, the classical algorithm seems advantageous whenever the problem requires an error smaller than the inverse number of features ( $\epsilon \in \widetilde{O}(1/d)$ ). Overall, the advantage is not self-evident nor easy to evaluate, as it depends on the problem and its data.

#### Example 4 - Dataset parameters.

Consider a quantum and a classical algorithm for fitting a PCA machine learning model. The model consists of the first k right singular vectors, corresponding to the largest k singular values of a matrix  $X \in \mathbb{R}^{n \times d}$ . The running times of the classical and the quantum algorithms are:

$$\widetilde{O}_{\mathrm{c}}\left(\frac{kdn}{\sqrt{\epsilon_L}}\log(\frac{1}{\gamma})\right)\,\widetilde{O}_{\mathrm{q}}\left(\frac{kd}{\delta^2}\frac{\|\boldsymbol{X}\|}{\theta}\frac{1}{\sqrt{p}}\frac{\mu(\boldsymbol{X})}{\epsilon_Q}\log(\frac{1}{\gamma})\right)$$

Here,  $\epsilon_L$  represents the error related to the relative spectral gap between eigenvalues,  $\theta, \epsilon_Q$  and  $\delta$  are the quantum algorithm approximation errors, p is the amount of retained variance,  $\mu(X)$  as per Definition B.3 and  $\gamma$  is the failure probability. Note that the dependence on n is encapsulated in the  $\widetilde{O}$  notation, as the quantum algorithm depends only polylogarithmically on it. One needs to estimate the parameters governing the query complexity of the QML algorithm on real datasets and use these estimates to compare the performances of the two algorithms.

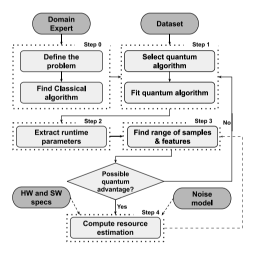


Fig. 1. Framework for evaluating speedups with quantum algorithms.

#### 4.2. The evaluation framework

Building on the considerations of the previous sections, we describe a framework for evaluating the impact of quantum machine learning algorithms on cybersecurity problems. The framework is summarized in Fig. 1.

**Step 0:** Formally define the problem and identify the best classical algorithm. Start by defining a machine learning problem. Collaborate with experts to formalize the problem, select the best classical algorithm, and choose a representative dataset for assessing speedup. The best classical algorithm might be the one with the best asymptotic complexity or the best performance in practice.

**Step 1:** Select a quantum machine learning model. Train the model and maximize the amount of approximation errors. Stop before the performance of the QML algorithm gets worse than tolerable. (a) Select a candidate

quantum machine learning algorithm to solve the problem. The quantum algorithm may not be present in literature or may need to be tailored to the problem, requiring the help of a quantum algorithm researcher. (b) Model the quantum algorithm with a classical algorithm that simulates the quantum procedure by artificially adding errors in the relevant steps, following the error types expected from the quantum algorithm's theoretical structure. Design the error to realistically/pessimistically model the approximation error expected in the actual quantum computation. Fit the quantum model and find the best set of parameters (e.g., approximation errors, effective condition number) that enables a satisfying performance, minimizing the running time. The parameters can be found manually or using algorithms for hyper-parameter tuning. Usually, a satisfying performance matches (or improves) the classical counterparts.

Step 2: Measure the dataset parameters that influence the running time of the quantum algorithm. Measure the other dataset parameters that influence the quantum algorithm's running time and cannot be traded for time efficiency. Examples can be the Frobenius norm of the dataset matrix, the maximum norm of the data points, the condition number, or other parameters specified in the theorem of the quantum algorithm and described in Section 4.1.2.

**Step 3:** Find the combinations of the number of data points and features that enable quantum advantage. Use the parameters found in the previous steps to quantify the query complexity of the quantum algorithm and the classical complexity as the number of data points and features increase. Estimate the dataset size (number of samples and features) at which the QML algorithm offers a significant advantage in query complexity compared to the classical approach.

**Decision:** If a practical dataset size is identified where the quantum algorithm exhibits an advantage in query complexity, proceed to step 4. Otherwise, return to step 1 and consider using a classical algorithm if no suitable quantum algorithms are found.

**Step 4:** *In-depth resource estimation.* Select a hardware architecture with a given connectivity and gate set, a noise model of the qubits and gates,

and an error correction code. Estimate the number of qubits and the resources needed for a better wall-clock time than the classical algorithm. In this context, the resource overhead associated with achieving fault tolerance — namely, the increased number of physical qubits, execution time, and overall cost — must be critically evaluated. If the required execution time per quantum gate is deemed unrealistic due to these overheads, reconsider the dataset dimensions identified in step 3.

This methodology can be used to identify promising QML applications and rule out the tasks for which a quantum advantage cannot be reasonably expected. It can be used to evaluate speedups both in the training and inference phases of QML algorithms. Under the assumption that the quantum clock-time will not become faster than the classical one (Babbush et al., 2021) and that the theoretical analyses of the algorithms are tight, the Decision step after step 3 suffices to understand which tasks might benefit from quantum advantage in the future. However, this decision process must consider the potential resource overhead introduced by fault-tolerant requirements. Once it is established that a quantum advantage might be plausible for a task, an in-depth resource estimation will provide further insights into the algorithm feasibility within the current state of the technology and into the necessary hardware improvements that could enable the advantage in the future. While we expect that improving quantum hardware will enable the practicality of many quantum advantages, the reader should also be mindful that classical architectures and computers are expected to improve, albeit at a slower pace.

#### 4.2.1. Advantages and limitations

The Decision step. This conceptual framework's most significant advantage and limitation lie in the *Decision* step.

On the positive side, ruling out the advantage of a QML algorithm based on its QRAM query complexity might spare the researchers the burden of performing an in-depth resource estimation, enabling cybersecurity experts and practitioners to expedite their evaluation of OML algorithms. Indeed, the parameter estimation steps (1–2) can be performed without the need for quantum simulators, enabling theoretical studies on large and realistic datasets. Rather than simulating the algorithm on a quantum simulator, practitioners will need to understand how to introduce artificial errors in the classical versions of the quantum algorithms, following the theoretical analysis of the quantum algorithm. Performing in-depth resource estimations requires deep knowledge of quantum technologies, and the scientific effort needed to make a proper one makes for a scientific contribution on its own. A preliminary analysis of the advantage in query complexity can be enough to understand the potential of a QML algorithm and spare the need for detailed resource estimation.

On the negative side, to successfully rule out an algorithm at the decision step, we need the following assumptions:

- The quantum computer's clock will not significantly outpace a classical one.
- The asymptotic query complexity is tight (and not loose because of proof artifacts).
- The models for the simulated errors align realistically with actual errors

Any deviation from these assumptions risks discarding a valuable candidate for quantum advantage. Consequently, the Decision step makes the evaluation framework a cautious approach, potentially missing advantageous QML candidates while avoiding misclassifying a suboptimal quantum algorithm as advantageous.

**Metrics for advantage.** The criteria for selecting a quantum algorithm in  $Step\ 1$  could benefit from further elaboration.

In this work, a quantum algorithm is considered advantageous if it demonstrates superior speed compared to its classical counterpart while retaining comparable performance. However, the selection criteria might be based on more complex security properties of QML algorithms, such

as their robustness and resilience to adversarial attacks. As of today, the robustness properties of many QML algorithms still need to be explored and require further research. This broader perspective could enhance the evaluation's depth, accounting for security beyond raw computational efficiency.

#### 5. Case study: PCA-based network IDSs

In this section, we demonstrate the evaluation framework on PCA-based QML methods and realistic datasets in one of the most studied applications of ML in cybersecurity: network intrusion detection. Specifically, we investigate potential quantum speedups in the training phase of three PCA-based anomaly detection algorithms: the Principal Component Classifier (PCC) by Shyu et al. (2003), the Reconstruction Loss method used in Verkerken et al. (2020), and our own extension of PCC, called Ensemble PCC, developed to enhance detection robustness through an ensemble of classifiers while preserving the simplicity of the original method. In all the three algorithms, PCA-extracted features are combined with the input sample in a binary classifier to distinguish between normal and anomalous network traffic. We use these algorithms on three standard network intrusion detection datasets: KDDCUP99, CICIDS2017, and DARKNET, with the goal of classifying normal versus anomalous traffic.

The machine learning training pipeline for these anomaly detection methods typically consists of three main stages: data preprocessing, PCA model extraction, and fine-tuning. Among these, the PCA model extraction step is often the most computationally intensive and resource-demanding, particularly in the classical setting. Our analysis specifically focuses on comparing the quantum and classical computational costs for this step, as it represents the primary computational bottleneck in the classical pipeline. However, classical preprocessing and fine-tuning can sometimes incur higher costs than the quantum model extraction step, highlighting the need for equivalent quantum algorithms to address these stages. While a thorough evaluation of quantum preprocessing and fine-tuning would be a valuable extension of this work, it lies beyond the scope of the present study.

We selected these PCA-based methods due to their simplicity and wide applicability. First, simple QML algorithms are essential building blocks for developing more complex quantum models. Second, simple ML algorithms are particularly valuable in cybersecurity domains — such as defense and cyber–physical systems — where interpretability and explainability are crucial (Geer Jr., 2018). These fields require transparency in automated decision-making processes, making simple, explainable models ideal for our evaluation.

Importantly, this paper does not aim to improve upon the latest state-of-the-art results in network intrusion detection. While more complex machine learning models may offer better detection performance, our goal is to demonstrate the application of our evaluation framework in investigating quantum speedups. The simplicity of the algorithms chosen is deliberate, as it allows us to focus on the core aim of assessing quantum versus classical computational efficiency. This section showcases how to apply our methodology's core steps (1, 2, 3, and Decision), with additional discussion on the resource estimation provided in Section 5.8. Although future work may explore quantum speedups for more sophisticated models, such as neural networks or more advanced ensemble methods, these comparisons are beyond the scope of this study.

#### 5.1. Anomaly detection algorithms

In the following, we denote the standardized input data matrix as  $X \in \mathbb{R}^{n \times d}$ , having n samples, d features, and rank r. The principal components of X are the eigenvectors of the covariance matrix  $X^TX \in \mathbb{R}^{d \times d}$ , denoted by  $\{e_i\}_i^r$ . Their index always corresponds to the one of the eigenvalues  $\{\lambda_i\}_i^r$ , ordered decreasingly  $\lambda_1 \geq \cdots \geq \lambda_r$ . Given a set  $S \in [r]$ , the variance explained by the components in S is  $p = \frac{\sum_{i \in S} \lambda_i}{\sum_{j=1}^r \lambda_j} \in [0, 1]$ .

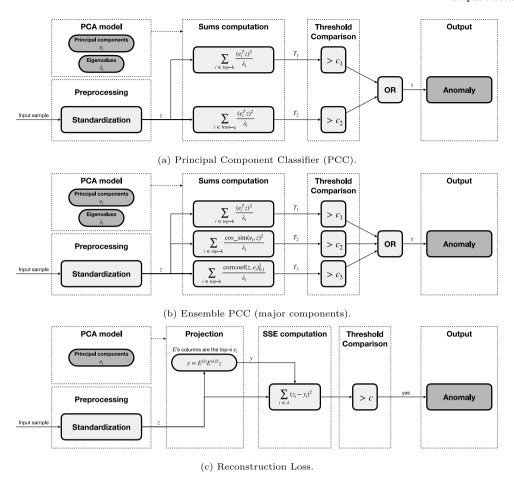


Fig. 2. Detection procedures of the three considered PCA-based anomaly detection classifiers.

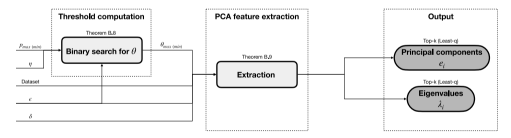


Fig. 3. Quantum PCA model extraction.

Fitting a PCA model means retrieving the principal components and corresponding eigenvalues that explain a total variance p. Usually, we are interested in the top-k components (major), though sometimes the least-q (minor) are of interest too. In this case, we denote the largest threshold such that the principal components with eigenvalues  $\sqrt{\lambda_i} > \theta$  explain variance  $p_{maj}$  as  $\theta \in (0, \sqrt{\lambda_1})$  and the smallest threshold such that the principal components with eigenvalues  $\sqrt{\lambda_i} < \theta_{\min}$  explain variance  $p_{min}$  as  $\theta_{\min} \in (0, \sqrt{\lambda_1})$ . These parameters are summarized in Table 1.

We proceed to describe the three anomaly detection algorithms and report the quantum routines that allow model fitting. The case study focuses on the advantages that the quantum routines can provide for the extracting the PCA model during the training phase, as this constitutes the algorithms' computational bottleneck. In all of these models, the PCA features are computed on a training set of normal (non-anomalous) data.

Principal Component Classifier (PCC). The first anomaly detection algorithm is by Shyu et al. (2003). Combining the input sample and

Table 1 PCA model parameters.

Parameters	Description
$e_i$	i <sup>th</sup> Principal component
$\lambda_i$	i <sup>th</sup> Eigenvalue
	of the covariance matrix
$p_{maj \text{ (min)}}$	Amount of variance explained
	by the top (least) eigenvalues
$\theta_{maj~(min)}$	Cut-off threshold for
	the top (least) eigenvalues
k	Number of top eigenvalues
q	Number of least eigenvalues

the extracted PCA features, the algorithm computes two sums and compares them against two anomaly detection threshold. The two sums are  $T_1 = \sum_{i=1}^k \frac{y_i^2}{\lambda_i}$  and  $T_2 = \sum_{i=r-q+1}^r \frac{y_i^2}{\lambda_i}$ , where k and q are the number of major and minor principal components (explaining variance  $p_{maj}$  and  $p_{min}$ ) and  $y_i = e_i^T z$ , with z vector of standardized observations and

 $e_i$  *i*th eigenvector corresponding to the  $\lambda_i$  eigenvalue. The algorithm classifies an observation z as an attack if  $(T_1 > c_1 \text{ or } T_2 > c_2)$ , where  $c_1$  and  $c_2$  are the outlier thresholds. The outlier threshold are computed on a validation set using a function of  $T_1$  and  $T_2$ , respectively, plus a parameter called false alarm rate  $\alpha \in (0,1)$  (Shyu et al., 2003). An increase of  $\alpha$  corresponds to a decrease in the outlier thresholds, leading the algorithm to detect more anomalies at the expenses of false positives. Sometimes we only use the sum on the top principal components, without computing  $T_2$  and  $T_2$ , we call this PCC with major components. We summarize the PCC detection procedure in Fig. 2(a).

**Ensemble PCC.** We extend the PCC algorithm by proposing two novel ways of computing  $y_i$ , using cosine similarity and correlation measures, in addition to the original dot product between  $e_i$  and z. The cosine similarity between  $e_i$  and z is computed as  $y_i = \frac{e_i^T z}{\|e_i\|\|z\|}$ , while the correlation is computed using the correcoef function from Numpy (Harris et al., 2020). Ensemble PCC computes three sums for the top-k principal components and three for the least-q. Like in PCC, each sum is compared again a threshold. If any sum exceeds this threshold, then the input sample is labeled as an anomaly. We call this method Ensemble PCC because the algorithm is equivalent to running three variations of PCC and labeling the sample as anomalous if any of the three models outputs so. We observe that this ensemble improves the performance of PCC. Fig. 2(b) shows the Ensable PCC detection pipeline with major components.

Reconstruction loss. The reconstruction loss anomaly detection algorithm has been widely employed for a large variety of tasks. One example of its use in network intrusion detection is given by Verkerken et al. (2020). The key idea in this algorithm is that the top PCA components extracted from a training set of normal data can be used to explain normal data, but introduce errors when used to express anomalous data. To classify an input sample, the algorithm projects it onto the space of principal components and then back into the original feature space. This process of projecting the sample into a lower dimensional space and back leads to a reconstruction error: the method computes the loss as the sum of square error (SSE) and use it as an anomaly score. The SSE sum is compared against a threshold and if the error is too large the sample gets labeled as an anomaly. The detection pipeline is depicted in Fig. 2(c).

**PCA model extraction.** The aim of the case study is to compare the classical and quantum running times needed to extract the PCA features from the training set of normal data points. These features can then be used in all the three detection algorithms explained above.

The quantum routines that can be used to fit the PCA models are explained in detail by Bellante et al. (2022). The important routines are reported in Theorems B.8 and B.9 and summarized in the following list.

- Quantum binary search for  $\theta$  (Theorem B.8). Given an amount p of target explained variance and a tolerance parameter  $\eta$ , the quantum routine finds a threshold  $\theta$  for the eigenvalues such that the top-k (least-q, with a minor fix) components selected by  $\theta$  explain at least  $\overline{p}$  variance, with  $\|p-\overline{p}\| \leq \eta$ . Because of the quantum phase estimation error, the routine also needs an error parameter  $\epsilon$  to estimate the eigenvalues on which the threshold  $\theta$  is learned. The routine runs in  $\widetilde{O}(\frac{\mu(A)\log(\mu(A)/\epsilon)}{\epsilon})$ .
- Quantum PCA extraction (Theorem B.9, note  $e_i = v_i$ ). Given a threshold  $\theta$ , the second routine enables extracting the eigenvalues  $\lambda_i$  and the top principal components  $e_i$ . The first task can be done in  $\widetilde{O}(\frac{\|X\|\mu(X)k\log(k)}{\theta\sqrt{\rho_e}\delta})$  to error  $\|\lambda_i \overline{\lambda_i}\| \leq 2\epsilon\sqrt{\lambda_i}$ , while the latter takes  $\widetilde{O}(dk\frac{\|X\|\mu(X)\log(k)\log(d)}{\theta\sqrt{\rho_e}\delta^2})$  to estimate the top-k principal components such that  $\|e_i \overline{e}_i\|_2 \leq \delta$ . This theorem can be modified (flip the condition of Alg. 4, step 3 Bellante et al., 2022) to extract the least-q principal components and eigenvalues. Provided a threshold  $\theta_{\min}$ , it can find the corresponding minor components in time  $\widetilde{O}(\frac{\theta_{\min}}{\sigma_{\min}}\frac{\mu(X)}{\epsilon}\frac{qd}{\sqrt{\rho_{\min}}})$  with guarantee  $\|e_i \overline{e}_i\|_2 \leq \delta$ .

 Table 2

 PCA model computation - running time parameters.

Parameters	Description
n	Number of training datapoints
d	Number of features
	or principal components' size
k (q)	Number of top (least) principal
	components/eigenvalues to extract
X	Dataset's spectral norm
$\mu(X)$	Dataset's normalization parameter
	(see Def Definition B.3)
$p_{maj \text{ (min)}}$	Amount of variance explained
, ()	by the top (least) eigenvalues
$\theta_{maj~(min)}$	Cut-off threshold for
, ()	the top (least) eigenvalues
$\sigma_{\min}$	Smallest value of all $\sqrt{\lambda_i}$
η	Max error allowed on $p_{mai \text{ (min)}}$
$\epsilon$	Max error allowed on each $\sqrt{\lambda_i}$
δ	Max error allowed on each $e_i$ ,
	in $\ell_2$ norm

The quantum PCA model extraction pipeline and the input parameters are represented in Fig. 3.

Depending on whether we require both the major and minor components or only the major ones, we compare the quantum running times with one of two classical alternatives: either the full singular value decomposition (SVD) with a complexity of  $O(\min(nd^2, n^2d))$ , or, if only the major components are needed, a randomized PCA variant with a lower complexity of  $O(ndk \log(k))$  (Halko et al., 2011). The running time parameters of these algorithms are summarized in Table 2.

#### 5.2. Experimental settings

To simulate the PCA model extraction and perform the training, we selected classical PCA-based methods implemented in scikit-learn (Pedregosa et al., 2011), which uses the numerical methods of LAPACK (Anderson et al., 1999), and modified them to model the error of the quantum subroutines.<sup>3</sup>

For the classical simulation of the quantum errors, we implemented Theorem B.9 for top-k singular vector extractor (and Theorem B.10 for clustering), which are based on quantum pure state tomography (Theorem B.4), amplitude estimation (Theorem B.6), and phase estimation (Theorem B.5). Theoretically, these QML routines show a running time advantage over their classical counterparts with high-dimensional data.

We fit our model on three publicly available datasets: KDDCUP99<sup>4</sup> (Tavallaee et al., 2009), CIC-IDS2017 (Sharafaldin et al., 2018), and DARKNET (Lashkari et al., 2020). In Appendix C, we report additional experiments on the tomography subroutine on the CIC-MALMEM-2022 (Carrier et al., 2022) dataset. For each dataset, we measure the parameters that influence the algorithms' running time and find the range of features and samples that would enable quantum advantage. The goal of this use case analysis is to compare the performance of training models using quantum algorithms against their classical counterparts by: ① studying the influence of the intrinsic error that affects quantum-machine-learning algorithms on the detection task and ② evaluating the expected quantum running time as the dataset grows

Considering the characteristics of the dataset under analysis (i.e., number of features, number of points, effective condition number,

<sup>&</sup>lt;sup>3</sup> https://github.com/tommasofioravanti/sq-learn

<sup>&</sup>lt;sup>4</sup> We are aware of selected datasets' limitations (especially for KDD-CUP99 (McHugh, 2000; Tavallaee et al., 2009)). We use such datasets only to fairly compare the performance of classical and quantum algorithms, with the goal of understanding the long-term impact of quantum machine learning.

approximation errors), we study the trade-off between the quantum algorithms' detection performance and running time, aiming at finding the "working point" that matches classical performances while minimizing the quantum algorithm's execution time. In particular, we fix the error parameters and evaluate the theoretical running time varying the number of samples and features. We then compare the running times of the classical and the quantum anomaly detection models to quantify the dataset dimensions needed to observe any quantum advantage, discussing which cybersecurity tasks may match such requirements.

In our analysis, we do not consider the time required to acquire the data in a classical (quantum readable) memory, as these procedures are to be performed only once when the data is received and need to be done in either case. In Appendix E, we conduct further experiments on clustering with PCA-based dimensionality reduction and the quantum version of k-means. For this problem, while using a cybersecurity-related dataset, we do not perform anomaly detection, but we compare the quantum and the classical algorithm on a clustering metric and show that the two algorithms offer similar performance.

**Tomography heuristics.** Appendix C discusses an analysis of our realistic simulation of the tomography subroutine. Tomography sample complexity scales as  $O\left(\frac{d\log d}{\delta^2}\right)$ , where d is the vector size. Experiments suggest that decreasing the number of samples by a significant constant factor still produces a near δ-accurate reconstruction of the state vector. We witness cases where we can save a constant factor up to  $\approx 10^3$  in sample complexity from for  $\delta = 0.03, d = 55$  (expected  $\approx 10^7$ ). This heuristic is used in the results of Section 5.5.

#### 5.3. PCC over KDDCUP99

We execute the principal component classifier over the KDDCUP99 dataset. While we perform a set of experiments varying the percentage of variance retained by the models, here we report the results of PCA70 and QPCA70, which retain 70% of the variance in the major components.

Regarding the dataset, we consider numerical features and split the dataset into a training set of 5,000 *normal* samples and a test set of 92,278 *normal* samples and 39,674 *attacks*. The training set is extracted using trimming to remove outliers and systematic random sampling (see ). Features are normalized and scaled for both test and training sets, with constant features removed.

**Performance analysis with Major Components only.** We classify a sample as an attack only if  $T_1 > c_1$  and normal otherwise (with  $c_1$  varying according to  $\alpha$ ). Results for PCA70 and QPCA70 are reported in Table 3, varying the false alarm rate  $\alpha \in (0,1)$ . For the quantum experiment, we consider the following error parameters: we use the quantum binary search of Theorem B.8 to estimate  $\theta$ , with parameters p=0.70,  $\epsilon_{\theta}=1$  as  $\epsilon$ , such that  $|\sigma_i-\overline{\sigma}_i|\leq \epsilon_{\theta}$ , and  $\eta=0.1$  such that  $|p-\overline{p}_{\theta}|\leq \eta$ ; we use Theorem B.9 to extract  $e_i$  and  $\lambda_i$  with parameters  $\epsilon=1$  to estimate singular values such that  $|\lambda_i-\overline{\lambda}_i|\leq \epsilon$ , and  $\delta=0.1$  is the error we tolerate in estimating singular vectors such that  $||e_i-\overline{e}_i||_2\leq \delta$ . With these error parameters, we match classical performances. As  $\alpha$  increases, recall increases, and precision decreases. This is expected because increasing  $\alpha$  increases the false alarm rate, resulting in a lower outlier threshold and more observations being classified as attacks. In Appendix D, we extend this experiment using the minor components,

Running time analysis with Major Components only. We compare the algorithms by plotting the classical and the quantum complexity, varying the number of samples n and features d. For the quantum running time, we consider the time complexity of the quantum binary search and the quantum top-k singular vector extraction. The first has a cost of  $\widetilde{O}\left(\frac{\mu(X)}{\epsilon\eta}\log(\frac{\mu(X)}{\epsilon})\right)$ , while the latter requires

Table 3
Comparison for classical © (PCA70) and quantum @ (QPCA70) principal components classifier with major components only over KDDCUP99.

α (%)	Recall (%)		Precision (%)		F1-score (%)		Accuracy (%)	
	c	q	c	q	c	q	c	q
1	93.14	92.84	98.63	98.68	95.81	95.67	97.55	97.47
2	93.19	92.88	98.18	98.23	95.62	95.48	97.43	97.35
4	96.04	95.75	96.51	96.57	96.28	96.15	97.76	97.69
6	98.51	98.12	94.20	92.21	96.30	96.12	97.72	97.62
8	98.67	98.36	92.01	92.07	95.22	95.11	97.02	96.69
10	99.44	99.12	90.05	90.10	94.51	94.40	96.53	96.46

 $\widetilde{O}\left(dk\frac{\|\mathbf{X}\|\mu(\mathbf{X})}{\theta\sqrt{p}\epsilon\delta^2}\log(k)\log(d)\right)$  queries to estimate the top-k right singular vectors and values. We compare the quantum running time with a randomized classical version of PCA, which has a complexity of  $O(ndk\log(k))$  (Halko et al., 2011) (since we are focusing on the major components only, there are better options than performing the full SVD, whose complexity is  $O(\min\{nd^2,n^2d\})$ ). Fig. 4(a) compares the quantum and classical running times in blue and green, respectively. We observe that the use of QML is not advantageous for small datasets. However, as the dataset's dimensionality increases, the query complexity advantage becomes evident (e.g., after  $\approx 4*10^6$  samples and  $\approx 50$  features).

#### 5.4. PCC and ensemble over CICIDS2017

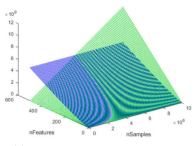
We consider the principal component classifier with major and minor components and the CICIDS2017 dataset with DDoS attacks and normal samples. The training set comprises 5,000 *normal* samples, while the test set of 87,300 *normal* and 70,000 *DDoS* samples. We follow the same preprocessing performed on the KDDCUP dataset.

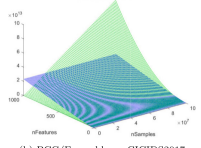
**Performance analysis.** In Table 4, we report the performance of QPCA70 (column q1) for PCC with major and minor components, using error parameters  $\delta=0.1, \epsilon=\epsilon_\theta=1, \eta=0.1, \gamma=\frac{1}{d}, \theta_{\min}=\sqrt{0.20},$  and varying the false alarm rate  $\alpha$ . Using these parameters, we observe that the QPCA-70 model matches the performances of the classical equivalent (similar to the analysis of Table 3). The performances of both the classical and quantum PCC algorithms on CICIDS2017 ( Table 4,  $q_1$ ) are significantly worse than the ones on KDDCUP99 with major components only ( Table 3).

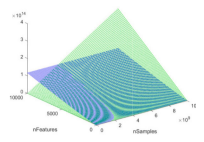
Keeping the same parameters, we assess the detection performance of the ensemble method (an improvement of PCC, first proposed here). The ensemble model improves the performance of PCC, particularly recall and accuracy, by utilizing six criteria for classifying an attack instead of two. This results in an improved recall, with a decrease in false negatives at the expense of false positives. Despite the drop in precision, the substantial recall improvements account for an overall F1-score increase. These results significantly improve over PCC, as seen in Table 4.

Running time analysis. We compare the running times by plotting the classical and the quantum complexity, varying the number of samples n and features d. We consider the classical full-SVD time complexity  $O\left(nd^2\right)$  to measure the classical running time, as we need all the components. For the quantum computation, instead, we consider the running time of quantum binary search and the quantum top-k singular vectors extractor to estimate the top-k components, plus the running time of the quantum least-q singular vectors extractor to estimate the minor components, which is  $\widetilde{O}\left(\frac{\theta_{\min}}{\sigma_{\min}}\frac{\mu(X)}{\sqrt{\rho_{\min}}}\frac{qd}{\sqrt{\rho_{\min}}}\right)$ , where  $\theta_{\min}$  is the custom  $\theta$  passed to the function to extract the least singular values and  $p_{\min}$  is the variance retained from the least components. In Fig. 4(b), the quantum and the classical running times are presented in blue and green, respectively.

As expected, the depicted running times are notably higher than those in the KDDCUP99 experiment, due to the comparison with classical full SVD complexity rather than the randomized one. Additionally,







(a) PCC major only on KDDCUP99.

(b) PCC/Ensamble on CICIDS2017  $\,$ 

(c) Rec. loss on CICIDS2017.

Fig. 4. Running time comparison of classical (green) and quantum (blue) algorithms. In Plot 4(a) we show the comparison of PCA and QPCA of Section 5.3 (KDDCUP99). In Plot 4(b) we show the running times of the algorithms discussed in Section 5.4 (CICIDS2017). In Plot 4(c) we show the comparison between PCA and QPCA over the CICIDS2017 dataset, as per Section 5.5.

Table 4 Comparison for QPCA70 with both major and minor components  $(q_1)$  and QPCA70 ensemble  $(q_2)$  over CICIDS.

or (0/s)	(%) Recall (%)		Precision (%)		F1-score (%)		Accuracy (%)	
α (%)								
	$q_1$	$q_2$	$q_1$	$q_2$	$q_1$	$q_2$	$q_1$	$q_2$
1	36.05	39.80	96.94	95.30	52.55	56.15	70.36	71.69
2	58.97	73.84	96.54	94.07	73.22	82.74	80.35	85.97
4	63.30	89.61	94.36	90.79	75.77	90.19	81.56	91.13
6	63.37	96.96	91.61	87.25	74.92	91.85	80.68	92.16
8	64.43	97.56	88.99	83.45	74.75	89.95	80.17	90.08
10	65.90	97.78	86.92	80.44	74.97	88.27	79.96	88.16

the quantum case involves not only binary search and top-k right singular vectors extraction but also the least-q extraction, contributing to increased quantum running time. A quantum advantage in query complexity emerges with a large dataset of  $\approx 2*10^9$  samples and  $\approx 100$  features.

The absence of a quantum advantage over classical machine learning, as demonstrated by the full classical SVD with a dataset of  $\approx 3*10^7$  samples and 500 features, suggests that quantum is not particularly beneficial for problems requiring the extraction of minor components. Moreover, leveraging a more efficient classical algorithm for extracting minor components, such as Minor Component Analysis (MCA) (Luo et al., 1997), would likely diminish the quantum advantage further, posing challenges for practical application in intrusion detection.

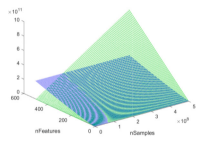
#### 5.5. Reconstruction loss over CICIDS2017

We fit the PCA-based model with reconstruction loss over the CI-CIDS2017 dataset, including all the kinds of attacks in the anomalous class, following the same preprocessing of Section 5.4. Unlike previous experiments, we perform hyper-parameter tuning on a validation set and preprocess the data with a quantile transform (see ). For the training set, we use 50,000 normal samples; for the validation set, 60,000 normal samples and 166,966 attacks; finally, for the test set, we use 140,000 normal samples and 389,590 attacks, following (Verkerken et al., 2020). Through hyper-parameters tuning, we found that the best PCA model has 12 principal components, which retain 94.88% of the variance. The outlier threshold is t = 0.425: each sample whose anomaly score (defined in Section 5.1) is higher than t is classified as anomalous.

**Performance analysis.** We found that the error parameters needed to match the performances of the classical algorithm are  $\epsilon_{\theta}=\epsilon=0.3$ ,  $\eta=0.00075$ , and we set  $\gamma$  to  $\frac{1}{d}$ . With these error parameters, we extract the same number of principal components of the classical model and explore perturbations on  $\delta$  (the error on the singular vectors). Setting  $\delta=0.01$ , we match the classical performances. By increasing  $\delta$ , the model tends to classify data points as attacks, as shown by the precision and recall trends in Table 5.

**Table 5** Comparison of the classical (c) PCA and quantum (q) QPCA principal components classifier with reconstruction loss over CICIDS, varying  $\delta$  error.

δ	Recall (%)		Precisio	Precision (%)		F1-score (%)		Accuracy (%)	
	q	c	q	c	q	c	q	c	
0.01	99.12		91.28	91.28	95.04	95.04	98.08	98.08	
0.1	99.17	00.10	91.23		95.03		98.08		
0.9	99.79	99.12	71.31		83.18		92.52		
2	100		27.30		42.89		50.66		



**Fig. 5.** Running time comparison of classical (green) and quantum (blue) algorithms for Rec. Loss over DARKNET. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

Running time analysis. For this experiment, we used the heuristic insights obtained on the tomography subroutine (see Section 5 and Appendix C): we target an approximation error  $\delta=0.01$  but divide the sample complexity of pure state tomography by a factor of 100. While this heuristic sacrifices provable guarantees, the model performs comparably to the classical counterpart in the testing phase.

Fig. 4(c) illustrates how the quantum and classical running times scale. This model on CICIDS2017 has more principal components than PCC with major components over KDDCUP99, resulting in higher running times. Specifically, the current model requires 32 components (retaining p=99.75% of the variance) compared to the PCC with major components only, which has 6 components (retaining p=70% of the variance). For this model, the QML algorithms achieve advantage after  $\approx 2*10^9$  samples.

#### 5.6. Reconstruction loss over DARKNET

We present the results obtained on PCA with Reconstruction loss over DARKNET. The test set comprises 21,000 normal samples and 14,000 anomalies, the training set uses 50,000 normal samples, and the validation set has 20,000 normal samples plus 10,000 anomalies. The d=85 features have been normalized and scaled to 0 mean and unit variance. The hyperparameter search optimizing the F1-score over

Table 6

Comparison of the classical (c) PCA and quantum (q) QPCA principal components classifier with reconstruction loss over DARKNET, varying  $\delta$  error. In the first row, we use the heuristic described in Appendix C. To get an error  $\delta=0.01$  we should use  $s=(36\frac{d\log d}{0.01})$  samples, but we see that h=s/100 suffice.

δ	Recall		Precision		F1-score		Accuracy	
	q	c	q	c	q	c	q	с
h	84.40		73.18		78.39		81.07	
0.9	87.08	84.17	68.12	73.71	76.44	78.59	78.17	81.35
2	91.47		59.97		72.44		71.69	

the classical PCA finds that the shorter running time is obtained with 35 principal components (which retain 99.65% of the variance) and outlier threshold t = 0.443.

**Performance analysis.** In the quantum model, the error parameters are set to  $\epsilon_\theta = \epsilon = 0.35, \eta = 0.0011$ , and p = 99.65%. With these parameters, 34 principal components are extracted, aligning with the classical case. Table 6 compares classical (c) and quantum (q) performances, showing how the model tends to classify data points as attacks as the error  $\delta$  increases, similar to the previous CICIDS2017 experiment. We achieve almost the same performances of the classical algorithm by using the heuristic and setting the theoretical error on singular vectors to  $\delta = 0.1$  (i.e., setting  $s = \frac{36d \log d}{\delta^2}$  with  $\delta = 0.01$  but taking s/100 samples).

**Running time analysis.** We plot the running time comparison in Fig. 5 using the parameters obtained from the hyperparameter search. We observe a quantum advantage in query complexity with  $10^8$  samples for dataset features up to hundreds.

#### 5.7. Decision

Upon initial examination, the dataset parameters required for the theoretical quantum advantage do not appear unrealistic, considering the volume of network packets companies receive daily. For instance, Microsoft's reported DDoS attack in January 2022 involved 3.47 Terabit of data per second (340 millions of packets per second) (Toh, 2021). Using a training dataset from such an attack and considering  $\approx 50$  features, the quantum model requires  $\approx 1.3 * 10^8$  operations against  $\approx 3.9 * 10^{10}$  of the randomized classical model (see Fig. 4(a)). Github also suffered a significant DDoS attack of about 1.35 Terabit per second with about 130 millions of packets per second (Kottler, 2018). In this case, a quantum model would require  $\approx 1.4 * 10^8$  steps against  $\approx 1.3 * 10^{10}$  of the classical one.

While this suggests a theoretical gap in operations of about two orders of magnitude, the advantage appears only for massive datasets. Therefore, the practical applicability of these QML algorithms for PCA-based network intrusion detection appears to be limited to large companies or organizations with the resources to handle vast amounts of data, computational power, time, and energy. In addition, for this particular PCA-based analysis, we witnessed high generalization performances by extracting the principal components on a small subset of data ( $\approx 5000$  or  $\approx 50000$  samples), rendering the need for such extensive datasets seemingly unrealistic for this application.

Despite these considerations, we proceed with a rough resource estimation to gauge the current state of quantum technology for these algorithms.

#### 5.8. Resource estimation

In this subsection, we analyze some key quantities for the oracle implementation of the QRAM, based on the code<sup>5</sup> and analysis of Matteo et al. (2020). While we refrain from performing an exhaustive resource estimation, this preliminary analysis of the execution time of a QRAM query indicates the current state of quantum hardware. We focus on the QRAM *Bucket Brigade parallel* circuit layout of Matteo et al. (2020), which provides reduced depth at the expenses of an increased number of auxiliary qubits. The analysis is performed on superconducting hardware with a defect-based error correcting surface code.

We consider a dataset with  $n=10^7$  rows and d=44 features, stored in a KP-tree (see Appendix B, near Theorem B.2) to allow quantum access to the dataset. This data structure consists of trees with a total of  $O(nd \log{(nd)})$  nodes. The content of these nodes can be stored using an address space of  $\lceil \log_2(nm \log_2(nm)) \rceil = 34$  bits, assuming a system word size of 1 bit. Although a 1-bit word size is optimistic, it simplifies the circuit, allowing us to directly apply the architectural estimates from Di Matteo et al. In practice, a 32-bit word size might be more realistic, but even with the optimistic 1-bit assumption, the results demonstrate the substantial resources required by the QRAM circuit.

This configuration would require approximately  $1.37 \times 10^{11}$  logical qubits and a circuit depth of 539 layers. This circuit also incurs a T-gate count of  $3.61 \times 10^{11}$ , a T-depth of 67, and a Clifford gate count of  $9.28 \times 10^{11}$ . For error correction, we base our analysis on a superconducting architecture with a defect-based surface code. Considering the same parameters of Di Matteo et al. we assume hardware with a gate error probability of  $10^{-5}$ , a failure probability for the magic states in the first concatenation layer of the QECC of  $10^{-4}$ , and a surface code cycle of 200 ns

While these assumptions are intentionally optimistic considering current hardware, they highlight the significant challenges that remain. Under these settings, a single QRAM query would take approximately 1.07 ms and require  $2.08\times10^{14}$  physical qubits. Looking more at the near future, less some realistic parameters would be hardware with a gate error probability of  $10^{-3}$ , a failure probability for the magic states in the first concatenation layer of the QECC of  $10^{-2}$ , and a surface code cycle of  $1\mu s$ . With these parameters, a single QRAM query would take approximately  $28.1~\rm ms$  and require  $7.31\times10^{16}$  physical qubits.

These numbers underscore the notable disparity in access time between a QRAM and a classical RAM, whose access time is in the scale of nanoseconds. This distinction in access time has cascading implications for the requirements on dataset size. Even a theoretical advantage of two orders of magnitude would not sufficiently compensate for the difference in memory access speed.

Finally, the sheer volume of physical qubits required is currently beyond the reach of today's technological trends. This underscores the practical limitations and emphasizes the considerable technological advancements needed to bridge the gap between theoretical potential and current quantum hardware capabilities for practical impacts.

#### 6. Conclusions

In this work, we analyzed the potential impact of QML on cyber-security tasks. We introduced a methodology for assessing quantum machine learning algorithms' impact on specific security problems. Besides laying out a clear methodology for studying quantum advantage in ML for cybersecurity, we demonstrated its application to fundamental algorithms used in network intrusion detection as a case study.

The results presented in this work show that QML techniques are unlikely to outperform classical methods for intrusion detection on small datasets. In our case study, a quantum advantage in query complexity becomes apparent as the problem size surpasses  $4*10^6$  samples and 50 features. There are settings where we observed significant theoretical gaps between the quantum and the classical models' running times. These requirements in the number of features and samples are potentially interesting for the intrusion detection domain since they align with publicly available datasets and real-world scenarios (Toh, 2021;

<sup>&</sup>lt;sup>5</sup> https://github.com/glassnotes/FT\_qRAM\_Circuits/tree/master

BBC, 2020; Kottler, 2018). However, the hardware slowdowns highlighted in Section 5.8 shift the advantage to bigger datasets, requiring even more physical qubits than estimated and making the advantage unlikely any soon.

Our case study suggests that fault-tolerant quantum computing could bring an asymptotic running time advantage, though more likely only for massive-scale datasets. However, the first generation of quantum computers is not expected to work on datasets of this size. Unless there is a significant advancement in hardware technology in the coming years, the sheer size of massive datasets will significantly exceed the capabilities of hardware implementations.

This evaluation framework can help the cybersecurity community find useful applications of quantum machine learning algorithms in practically relevant tasks and better distinguish the hype often associated with new technologies from their practical impact. As more experts and practitioners from the field start practicing with the technologies, it will be possible to acquire a better understanding of the potential of quantum machine learning and of the way it will impact future users, markets, and attack-defense dynamics.

#### CRediT authorship contribution statement

Armando Bellante: Writing – review & editing, Writing – original draft, Validation, Supervision, Methodology, Investigation, Conceptualization. Tommaso Fioravanti: Visualization, Software, Methodology, Investigation, Data curation. Michele Carminati: Writing – review & editing, Writing – original draft, Validation, Conceptualization. Stefano Zanero: Writing – review & editing, Supervision, Funding acquisition, Conceptualization. Alessandro Luongo: Writing – review & editing, Writing – original draft, Validation, Supervision, Methodology, Investigation, Conceptualization.

#### Acknowledgments

We thank Adithya Sireesh for useful discussions on error correction. A.B. would like to thank professors Donatella Sciuto and Ferruccio Resta for their support and acknowledges the financial support from ICSC - "National Research Centre in High Performance Computing, Big Data and Quantum Computing" Spoke 10, funded by European Union – NextGenerationEU - under the grant PNRR-CN00000013-HPC. Research at CQT is funded by the National Research Foundation, Singapore, the Prime Minister's Office, and the Ministry of Education, Singapore under the Research Centres of Excellence programme's research grant R-710-000-012-135. We also acknowledge funding from the Quantum Engineering Programme (QEP 2.0) under grant NRF2021-QEP2-02-P05.

#### Declaration of competing interest

The authors declare the following financial interests/personal relationships which may be considered as potential competing interests: Armando Bellante reports financial support was provided by Italian National Research Centre in High Performance Computing, Big Data and Quantum Computing. Alessandro Luongo reports financial support was provided by Inveriant Pte. Ltd. Armando Bellante reports a relationship with Italian National Research Centre in High Performance Computing, Big Data and Quantum Computing that includes: funding grants. Tommaso Fioravanti reports a relationship with IBM that includes: employment. Alessandro Luongo has patent #11741386 issued to Bull SA. Stefano Zanero serves as an editor for Computers & Security. If there are other authors, they declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

#### Appendix A. Brief introduction to quantum information

The fundamental unit of quantum information is the qubit. The state of a qubit can be expressed as a linear combination of vectors from an orthonormal basis of  $\mathbb{C}^2$ , such that the sum of squares of the absolute values of the coefficients sums up to one. Often, we express states using the computational basis  $|0\rangle = \begin{bmatrix} 1 & 0 \end{bmatrix}^T$ ,  $|1\rangle = \begin{bmatrix} 0 & 1 \end{bmatrix}^T$ , meaning that a generic qubit state can be written as  $|\varphi\rangle = \alpha \, |0\rangle + \beta \, |1\rangle$ , with  $\alpha, \beta \in \mathbb{C}$  and  $|\alpha|^2 + |\beta|^2 = 1$ . The coefficients are also called *amplitudes* and if more than one is non-zero, we say that the state is in a *superposition* of the basis states.

A quantum register is an ensemble of qubits. We can express the state of a quantum register as  $|v\rangle=\frac{1}{\|v\|}\sum_{i=0}^{2^n-1}v_i\,|i\rangle\in\mathbb{C}^{2^n}$ , where  $n\in\mathbb{N}$  is the number of qubits in the register, each  $v_i\in\mathbb{C}$ , and  $\sum_{i=0}^{2^n-1}|v_i|^2=1$ . Here  $|i\rangle$  is the ith vector of the computational basis of  $\mathbb{C}^{2^n}$ ; i.e.,  $|i\rangle=|i_0\rangle\otimes|i_1\rangle\otimes\cdots\otimes|i_{n-1}\rangle$ , where  $i_j$  denotes the jth bit of the binary encoding of i. This representation is analogous to the decomposition of a vector in the computational basis of a vector space. It is crucial to note that n qubits suffice to span a space of dimension  $2^n$ .

A quantum algorithm consists of *evolving* and *measuring* a certain initial state, multiple times. (a) The *evolution* of a *n*-qubits quantum state is described by a unitary matrix  $U \in \mathbb{C}^{2^n \times 2^n}$ ; i.e., a matrix such that  $U^\dagger U = U U^\dagger = I$ . These matrices preserve norms, mapping quantum states to other valid quantum states. Any quantum algorithm, excluding measurements, corresponds to a unitary matrix. These matrices can be decomposed in one and two-qubits quantum gates, the basic elements of quantum circuits.

Evolutions can be combined — multiplications and tensor products of unitary matrices are still unitary. In practice, the reader might find it convenient to think of multiplications of unitary matrices as circuits applied to the same qubits (in series) and tensor products as circuits applied to different qubits (in parallel). The evolution of quantum states is a unitary operation that requires quantum gates to be reversible. Classical non-reversible gates (such as the AND) have reversible counterparts (CCNOT), implemented at the cost of introducing some auxiliary qubits.

(b) Reading data from quantum states is not as immediate as accessing a memory location on a classical computer. While quantum states can be in a superposition of the computational basis, quantum mechanics does not allow us to retrieve all the information at once from these superpositions. *Measurements* are modeled through a set of measurement operators  $\{M_m\}_{m=0}^{2^n-1}$  such that  $\sum_{m=0}^{2^n-1} M_m^{\dagger} M_m = 1$ , with each  $M_m \in \mathbb{C}^{2^n}$ . The probability that an outcome m occurs for a state  $|\varphi\rangle$  is given by  $p(m) = \langle \varphi | M_m^{\dagger} M_m | \varphi \rangle$ , and after the  $m^{\text{th}}$  outcomes is measured, the state *collapses* on a new state  $|\varphi'\rangle = \frac{M_m |\varphi\rangle}{\sqrt{\langle \varphi | M_m^{\dagger} M_m | \varphi\rangle}}$ . In this work, we can restrict the measurement operators to  $\{|i\rangle\langle i|\}_{i=0}^{2^n-1}$ . In this case, the resulting state of the quantum register is a vector of the computational basis. In practice, if we measure a register (or a portion of it), we can only see one bit-string corresponding to one computational basis state that the register decomposes onto (a sequence of 0s and 1s, like in a classical register).

After the measurement, the register (or the qubits read) will lose the superposition and collapse on the measured state without providing further information on the amplitudes. The process of reconstructing the amplitudes of a quantum state, with respect to a given basis, requires statistics over several measurements and is called *tomography* of pure states. The reversibility of quantum mechanics prevents a generic copying algorithm from existing (no-cloning theorem). To perform tomography, one must re-create the quantum state by repeating the algorithms from scratch multiple times and sample (Theorem B.4).

#### Appendix B. Algorithms and subroutines implemented

In this section, we detail the algorithms that we simulated to carry on the case study. Starting from the input procedures, we precisely define the meaning of quantum access to a matrix.

**Definition B.1** (*Quantum Access to a Matrix (Kerenidis and Prakash, 2020a*)). We have quantum access to a matrix  $A \in \mathbb{R}^{n \times m}$ , if there exists a data structure that allows performing the mappings  $|i\rangle |0\rangle \mapsto |i\rangle |a_{i,\cdot}\rangle = |i\rangle \frac{1}{\|a_{i,\cdot}\|} \sum_{j=1}^{m} a_{ij} |j\rangle$ , for all i, and  $|0\rangle \mapsto \frac{1}{\|A\|_F} \sum_{i=1}^{m} \|a_{i,\cdot}\| |i\rangle$  in time  $\widetilde{O}(1)$ .

This definition directly extends to vectors, which are special kind of matrices. We say to have quantum access to a vector of size m if we can implement the mapping

$$U_{x}|0\rangle = \frac{1}{\|x\|} \sum_{i \in m} x_{i} |i\rangle \tag{B.1}$$

its controlled version, and inverse, in time  $\widetilde{O}(1)$ .

Having access to a quantum random access memory, it is possible to efficiently create quantum access to matrices and vectors. This requires preprocessing the input data and storing a tree data structure, also known as KP-trees, after the authors of this procedure, in the QRAM. We state the main result and invite the interested reader to check the details in Kerenidis and Prakash (2017, 2020a).

**Theorem B.2** (Implementing Quantum Operators Using an Efficient Data Structure Kerenidis and Prakash (2017)). Let  $A \in \mathbb{R}^{n \times m}$ . There exists a data structure to store the matrix A with the following properties:

- 1. The size of the data structure is  $O(nm \log (nm))$ .
- 2. The time to update/store a new entry  $(i, j, A_{ij})$  is  $O(\log (nm))$ .
- Provided coherent quantum access to this structure there exists quantum algorithms that implement the mappings of Definition B.1 in time O(polylog(nm)).

This definition of quantum access makes a data normalization parameter appear in many quantum algorithms. For a matrix X, we call this normalization parameter  $\mu(X)$ . The smaller this parameter is, the more efficient the algorithms are. The appendices of Kerenidis and Prakash (2017) describe how to implement the data structure for  $\mu(X) = \|X\|_F$ . A subsequent manuscript by Kerenidis and Prakash (2020a) describes how to obtain other values of  $\mu$ , which we describe in the following definition.

**Definition B.3** ( $\mu(X)$  (*Kerenidis and Prakash*, 2020a)). Let  $X \in \mathbb{R}^{n \times d}$  be a matrix. We define  $\mu(X) = \min_{p \in [0,1]} (\|X\|_F, \sqrt{s_{2p}(X)s_{2(1-p)}(X^T)})$ , with  $s_q(X) = \max_i \|X_{i,\cdot}\|_q^q$ , for  $q \in [0,2]$ .

It is possible to probe the optimal  $\mu$  during the data loading preprocessing step. In practice, if the dataset is entirely available at loading time, one could measure  $\mu$  for different values of  $p \in [0,1]$  and compare it to the Frobenius norm, to know which data structure is more efficient. The simulation of the quantum access routine is not required by our evaluation framework, while the estimation of the best  $\mu$  is.

To retrieve data from quantum algorithms one has to consider some tomography procedures. The algorithms that we consider leverage the following tomography routine.

**Theorem B.4** (Tomography (Kerenidis and Prakash, 2020)). Given access to the mapping  $U_x \mid 0 \rangle \mapsto \mid x \rangle$  and its controlled version, for  $\delta > 0$ , there is an algorithm that produces an estimate  $\overline{x} \in \mathbb{R}^m$  with  $\|x\|_2 = 1$  with probability at least 1 - 1/poly(m) using U  $O(\frac{m \log m}{\delta^2})$  times such that  $\| \mid x \rangle - \overline{x} \|_2 \leq \delta$  and using U  $O(\frac{\log(d)}{\delta^2})$  such that  $\| \mid x \rangle - \overline{x} \|_{\infty} \leq \delta$ 

Now that we discussed input and output routines, we move on and present the main building blocks of the quantum algorithms considered in this work: phase estimation, amplitude estimation, and distance/inner products estimation. Each of this routine builds on the previous one.

**Theorem B.5** (Phase Estimation (Nielsen and Chuang, 2010)). Let U be a unitary operator with eigenvectors  $\begin{vmatrix} v_j \end{vmatrix}$  and eigenvalues  $e^{i\theta_j}$  for  $\theta_j \in [-\pi,\pi]$ , i.e. we have  $U \begin{vmatrix} v_j \end{vmatrix} = e^{i\theta_j} \begin{vmatrix} v_j \end{vmatrix} v_j$  for  $j \in [n]$ . For a precision parameter  $\epsilon > 0$ , there exists a quantum algorithm that runs in time  $O(\frac{T(U)\log(n)}{\epsilon})$  and with probability 1 - 1/poly(n) maps a state  $|\phi_i\rangle = \sum_{j \in [n]} \alpha_j \begin{vmatrix} \epsilon \\ v_j \end{pmatrix}$  to the state  $\sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in [n]} \alpha_j \begin{vmatrix} v_j \\ v_j \end{vmatrix} = \sum_{j \in$ 

This procedure can be made consistent, in the sense that multiple runs of the same algorithm return a phase with the same error (Ta-Shma, 2013; Kerenidis et al., 2020b). In our case study, we simulate this version of phase estimation, which is a consistent phase estimation. We model the error using both the theory behind phase estimation and the procedure that makes it consistent. Using phase estimation, we can build an amplitude estimation algorithm.

**Theorem B.6** (Amplitude Estimation (Brassard et al., 2002)). There is a quantum algorithm called amplitude estimation which takes as input one copy of a quantum state  $|\psi\rangle$ , a unitary transformation  $U=2|\psi\rangle\langle\psi|-I$ , a unitary transformation V=I-2P for some projector P, and an integer t. The algorithm outputs  $\tilde{a}$ , an estimate of  $a=\langle\psi|P|\psi|\psi|P|\psi\rangle$ , such that:

$$|\tilde{a}-a| \leq 2\pi \frac{\sqrt{a(1-a)}}{t} + \frac{\pi^2}{t^2}$$

with probability at least  $8/\pi^2$ , using U and V t times each. If a=0 then  $\tilde{a}=0$  with certainty, and if a=1 and t is even, then  $\tilde{a}=1$  with certainty.

Using amplitude estimation on a modified Hadamard test circuit, it is possible to estimate distances and inner products.

**Theorem B.7** (Distance and Inner Products Estimation (Kerenidis et al., 2019)). Assume for a matrix  $V \in \mathbb{R}^{n \times d}$  and a matrix  $C \in \mathbb{R}^{k \times d}$  that the following unitaries  $|i\rangle|0\rangle \mapsto |i\rangle|v_i\rangle$ , and  $|j\rangle|0\rangle \mapsto |j\rangle|c_j\rangle$  can be performed in time T and the norms of the vectors are known. For any  $\Delta > 0$  and  $\epsilon > 0$ , there exists a quantum algorithm that computes:  $|i\rangle|j\rangle|0\rangle \mapsto |i\rangle|j\rangle\left|\overline{d^2(v_i,c_j)}\right\rangle$  where  $|\overline{d^2(v_i,c_j)}-d^2(v_i,c_j)|\leqslant \epsilon$  w.p.  $\geq 1-2\Delta$  in time  $\widetilde{O}\left(\frac{\|v_i\|\|c_j\|^T\log(1/\Delta)}{\epsilon}\right)$ .

Leveraging phase estimation, it is possible to implement one of the main ingredient of the quantum PCA model extraction pipeline of Fig. 3: the binary search for  $\theta$ .

**Theorem B.8** (Quantum Binary Search for the Singular Value Threshold (Bellante et al., 2022)). Let there be quantum access to a matrix  $\mathbf{A} \in \mathbb{R}^{n \times m}$ . Let  $\eta, \epsilon$  be precision parameters, and  $\theta$  be a threshold for the smallest singular value to consider. Let  $p \in [0,1]$  be the factor score ratios sum to retain. There exists a quantum algorithm that runs in time  $\widetilde{O}\left(\frac{\mu(\mathbf{A})\log(\mu(\mathbf{A})/\epsilon)}{\epsilon\eta}\right)$  and outputs an estimate  $\theta$  such that  $\left|p - \sum_{i:\overline{\sigma}_i \geq \theta} \lambda^{(i)}\right| \leq \eta$ , where  $\left|\overline{\sigma}_i - \sigma_i\right| \leq \epsilon$ , or detects whether such  $\theta$  does not exists.

Similarly, with the aid of quantum tomography, it is possible to extract the top principal components and corresponding eigenvalues.

**Theorem B.9** (Top-k Singular Vectors Extraction Bellante et al. (2022)). Let there be efficient quantum access to a matrix  $\mathbf{A} \in \mathbb{R}^{n \times m}$ , with singular value decomposition  $\mathbf{A} = \sum_i^r \sigma_i \mathbf{u}_i \mathbf{v}_i^T$ . Let  $\delta > 0$  be a precision parameter for the singular vectors,  $\epsilon > 0$  a precision parameter for the singular values, and  $\theta > 0$  be a threshold such that  $\mathbf{A}$  has k singular values greater than  $\theta$ . Define  $p = \frac{\sum_{i: \overline{a}_i \geq \theta} \sigma_i^2}{\sum_j^r \sigma_j^2}$ . There exist quantum algorithms that estimate: The top k left singular vectors  $\mathbf{u}_i$  of  $\mathbf{A}$  with unit vectors  $\overline{\mathbf{u}}_i$  such that  $\|\mathbf{u}_i - \overline{\mathbf{u}}_i\|_2 \leq \delta$ 

<sup>&</sup>lt;sup>6</sup> We use  $\log(nm)$  instead of  $\log^2(nm)$  because the extra  $\log(nm)$  term in the original statement comes from the size of the system word.

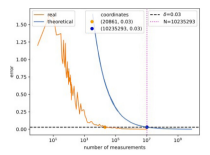


Fig. C.6. Theoretical bound (blue) on the number of measurements required for tomography and a numerical simulation (orange), on the first principal component of CIC-MalMem2022. The horizontal dashed line indicates the target  $\ell_2$  error (0.03) of a vector of length 55. The vertical dashed line represents the theoretical bound. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

with probability at least 1-1/poly(n), in time  $\widetilde{O}\left(\frac{\|A\|}{\theta}\frac{1}{\sqrt{p}}\frac{\mu(A)}{\epsilon}\frac{kn}{\delta^2}\right)$ ; The top k singular values  $\sigma_i$ , factor scores  $\lambda_i$ , and factor score ratios  $\lambda^{(i)}$  of A to precision  $\epsilon$ ,  $2\epsilon\sqrt{\lambda_i}$ , and  $\epsilon\frac{\sigma_i}{\|A\|_F^2}$  respectively, with probability at least 1-1/poly(m), in time  $\widetilde{O}\left(\frac{\|A\|}{\theta}\frac{1}{\sqrt{p}}\frac{\mu(A)k}{\epsilon}\right)$  or during any of the two procedures above.

Finally, building on the distance and inner product estimation routine, one can use a quantum version of k-means.

**Theorem B.10** (q-Means (Kerenidis et al., 2019)). Assume to have quantum access to a data matrix  $V \in \mathbb{R}^{n \times d}$ . For  $\delta > 0$ , the q-means algorithm with high probability outputs centroids  $\{\overline{\mu_j}\}_{j=1}^k$  that are  $\delta$ -close in  $\ell_2$  norm to the centroids of the classical k-means algorithm in time

$$\widetilde{O}\left(kd\frac{\eta}{\delta^2}\kappa(V)(\mu(V)+k\frac{\eta}{\delta})+k^2\frac{\eta^{1.5}}{\delta^2}\kappa(V)\mu(V)\right)$$

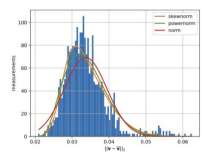
per iteration with  $1 \le \|v_i\|^2 \le \eta$  and a number of iterations proportional to the classical algorithm.

The simulation code is available on github $^7$  and more info can be found in Fioravanti (Fioravanti, 2022).

#### Appendix C. Vector state tomography

The number of measurements,  $N=\frac{36d\log d}{\delta^2}$  is the result of a probabilistic bound that guarantees a success probability greater than 1-1/poly(d). Maintaining this failure probability low is crucial in applications where tomography has to be repeated many times on different vectors – imagine its use in the prediction stage of a machine learning model – as its repeated use will eventually lead to an estimate that exceeds the target error. In this section, we exhibit a case where the number of samples  $N=\frac{36d\log d}{\delta^2}$  is larger than the effective number of samples needed to obtain a certain accuracy: i.e., with a number of measurements considerably lower than N we obtain an estimate  $\overline{x}$  with error  $\delta$ . We perform tomography over the first principal component of the dataset CIC-MalMem-2022 (Carrier et al., 2022). The vector has size d=55. This experiment is reported in Fig. C.6. The plot shows how many measurements (x-axis) are necessary to get a vector estimate with a specific error (y-axis), both following the theoretical bound (blue curve) and simulating the actual tomography procedure (orange curve).

To get an estimate with error 0.03 in  $\ell_2$  distance, we need about  $10^4$  measures instead of  $\approx 10^7$ , as suggested by the theoretical bound. To corroborate this finding, in Fig. C.7, we start from a fixed number of measurements (20.861) (which is the number of measurement sufficient



**Fig. C.7.** Distribution of error (x-axis) for tomography of the first principal component of CIC-MalMem2022 by repeating the experiment 1000 times with 20,861 samples. From this statistic with fit and plot three distributions (skew normal, power normal, and normal distribution).

**Table C.7**Comparison for classical © (PCA70) and quantum @ (QPCA70) principal components classifier with major and minor components over KDDCUP99.

α (%)	Recall (%)		Precision (%)		F1-score (%)		Accuracy (%)	
	c	q	c	q	c	q	c	q
1	98.62	98.68	97.26	96.99	97.94	97.83	98.75	98.68
2	98.66	98.73	95.96	95.58	97.29	97.13	98.35	98.24
4	98.80	98.90	91.61	91.42	95.07	95.01	96.91	96.88
6	98.84	98.99	88.88	88.25	93.60	93.31	95.78	95.73
8	98.93	99.27	86.11	85.84	92.08	92.07	94.88	94.85
10	99.59	99.87	83.25	83.14	90.69	90.74	93.85	93.87

to reach error 0.03 from Fig. C.6), and we repeat the tomography for 1000 times, plotting on the *y*-axis the frequency of the  $\ell_2$  error observed. As we can see, the error is roughly centered around 0.03.s

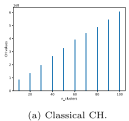
### Appendix D. Performance analysis with minor and major components over KDDCUP99

We corroborate the efficacy of the classifier discussed in Section 5.3 with another experiment. In this setting, we classify a sample as an attack if  $(T_1 > c_1 \text{ or } T_2 > c_2)$ , and as normal otherwise. We maintain the same error parameters reported in Section 5.3 to estimate both major and minor components (QPCA70 with 10 major and 7 minor components, with respect to 10 major and 6 minor of the classical case). To extract the minor components, we set the threshold to  $\theta =$  $\sqrt{0.20}$  as a parameter in the quantum least-q singular vectors extraction (Theorem B.9). With these error parameters, we extract a number of principal components very close to the classical model (QPCA70 with 10 major and 7 minor components, with respect to 10 major and 6 minor of the classical case). Also in this case, we match the classical performance, reported in Table C.7. When using both major and minor components, the recall is higher than the model that used only major components (as reported in Table 3), but precision is lower. This is expected as using two control summations in OR (checking both  $T_1$ and  $T_2$ , rather than  $T_1$  only) leads to a higher chance of classifying an observation as an anomaly, resulting in an increase in false positives and a decrease in false negatives (hence improving recall).

## Appendix E. Clustering with PCA-based dimensionality reduction and K-means over KDDCUP99

In this section, we demonstrate the feasibility of applying our methodology to the clustering task. We perform K-Means after applying PCA-based dimensionality reduction and compare the classical and quantum algorithm versions. We evaluate the clustering quality of the classical and quantum algorithm versions by computing the *Calinski-Harabasz (CH) index (Pedregosa et al., 2011)* on the resulting clusters (see Fig. E.8).

<sup>&</sup>lt;sup>7</sup> https://github.com/tommasofioravanti/sq-learn



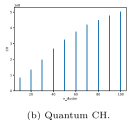


Fig. E.8. CH comparison between PCA and K-Means and q-PCA and q-Means, by varying the number of clusters  $n_k$ .

For this experiment, we use the KDDCUP99 dataset, projecting all the data into a new 1-dimension PCA space and applying the K-Means clustering. We vary the number of clusters  $n_k = [10, 20, 30, \dots, 100]$  and compute the CH index. Since the PCA model retains only the first principal component, we classically compute the percentage of variance retained by the first principal component, which corresponds to  $\approx 0.6$ . We use this value as input parameter p = 0.6 into the quantum binary search, with  $\epsilon_\theta = \epsilon = 5$  and  $\eta = 0.1$ . Then, we extract the top-k components with error  $\delta = 0.1$ . After retrieving a classical description of the first principal component, we use it to project the data into the q-PCA feature space. We apply the q-means algorithm over these 1-dimensional data with error  $\delta = 0.0005$  with  $n_k$  clusters. Once obtained a classical description of the clustering, we compute the CH score. As shown In Fig. E.8(a), we match the classical CH index values.

#### Further information on experiments.

For more detail on the wrapper approach and the validation process we used for hyperparameter optimization we refer the interested reader to Kohavi and John (1997), Akiba et al. (2019). Our hyperparameter tuning in Section 5.5 found that the number of quantiles that maximizes the F1 score is 751.

#### Data availability

The datasets are publicly available and the link to the repo containing the main code is included the manuscript.

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#### Further reading

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