A modular framework for randomness extraction based on Trevisan’s construction

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Informally, an extractor delivers perfect randomness from a source that may be far away from the uniform distribution, yet contains some randomness. This task is a crucial ingredient of any attempt to produce perfectly random numbers—required, for instance, by cryptographic protocols, numerical simulations, or randomised computations. Trevisan’s extractor raised considerable theoretical interest not only because of its data parsimony compared to other constructions, but particularly because it is secure against quantum adversaries, making it applicable to quantum key distribution.

We discuss a modular, extensible and high-performance implementation of the construction based on various building blocks that can be flexibly combined to satisfy the requirements of a wide range of scenarios. Besides quantitatively analysing the properties of many combinations in practical settings, we improve previous theoretical proofs, and give explicit results for non-asymptotic cases. The self-contained description does not assume familiarity with extractors.

I. INTRODUCTION

Random numbers are key ingredients for many purposes concerning communication or computation: secretly shared, perfectly random bit strings enable two parties to communicate in private using a one-time pad, without the possibility of a third party decrypting any of the messages they exchange. Stochastic algorithms used in numerical simulation or machine learning also rely on random numbers as part of their input. In all such applications, it is usually best to have uniform randomness available, that is, an observer should not have prior knowledge about the distribution of numbers, or, more specifically, the content of bit strings: Each string should be equally probable from his point of view. Some applications, such as encrypting a message with information-theoretic security, are even impossible if the randomness used to choose the key is not equivalent to a uniform distribution. Unfortunately, despite their usefulness and the need for them, uniformly distributed random bits are almost impossible to generate in practice. On the other hand, there are plenty of physical resources containing “some” randomness, for instance radioactive decay, thermal fluctuations, certain measurements on photons, or many others.

This contrast motivates the study of randomness extractors: Functions that map longer, slightly random bit strings shorter, perfectly random bit strings. They convert an initial distribution of random numbers (the source) that satisfies certain assumptions on “how random” it is into an almost uniform distribution over the output bit strings. As suggested by intuition, this is impossible in a completely deterministic way, and extractors indeed require a second source of randomness, the seed, that is usually assumed to be perfectly uniformly distributed.

The goal of this work is twofold: to implement a specific randomness extractor devised by Trevisan in 1999 as a practical companion to the abundant amount of theoretical literature on the subject, and to provide an overview and guidance on the topic to experimentalists who need to use extractors, but would not benefit from working through all fundamental publications. Trevisan’s construction has three particular advantages: For one, it is secure in the presence of quantum side information, as was shown by one of the authors in collaboration with others. This is especially important in the context of cryptography, where an adversary usually has some prior information about the initial distribution used as raw material to produce a secret key. With a quantum-proof extractor, it is possible to eliminate all these undesired correlations by turning the initial distribution into a uniform one—a task referred to as privacy amplification. With quantum key distribution (QKD) systems gradually transitioning from research labs into commercial applications, it is very important to implement this crucial protocol step, and given a bound on how random and how correlated with some (quantum-)memory a bit string is, the algorithm can indeed perform the task of producing truly random and uncorrelated bits with the help of a short seed of uniformly distributed bits.

Another crucial advantage of Trevisan’s construction is that the required seed length is only poly-logarithmic in the length of the input. This greatly outperforms randomness extractors based on (almost) universal hashing, which are currently most often used in quantum cryptographic applications, but require a seed whose size unfortunately scales with the length of the raw input (output) bits.

In addition, Trevisan’s construction is a strong extractor, which means that the seed is almost independent of the final output. This implies that randomness in the seed is not consumed in the process (compared to weak extractors) and can be used at a later time—or, as in
the case of privacy amplification, it can be obtained by the adversary without compromising the security of the QKD scheme.

Despite the considerable theoretical attention the field of extractors has received during the last decade, there is, to our knowledge, only a single publication, Ref. [7], that discusses a prototypical implementation of Trevisan’s construction. However, their work has some drawbacks: Compared to Ref. [7], our implementation offers greater flexibility as the operator can combine various different building blocks that make up the extractor, and so can specifically engineer an algorithm for his needs. Comparing the performance, our implementation exceeds the throughput of [7] by several orders of magnitude, and is for the first time able to scale to data sets of realistic size (exceeding the maximal amount considered in [7] by 10 orders of magnitude) for which the amount of extracted randomness actually exceeds the size of the initial seed, which marks the regime in which Trevisan’s construction prevails over two-universal hashing. Besides, the full source code of our implementation is available and can be inspected and used as basis for further research. We therefore hope that our implementation will be of use for applications in the context of quantum cryptography, for implementing random number generators, or as a testbed for developing new ideas about extractors.

In Section II, we give more proper descriptions and definitions of the involved concepts and constructs. In particular, we discuss the necessary notions of entropy and the distance of a distribution from uniform (relative to an overserver). However, no prior knowledge about randomness extractors is assumed. Section III contains the necessary technical details, and can be skipped upon first reading. Section IV is devoted to the implementation: It describes the software architecture and discusses some important technical details, explains how to add new components, and gives concise algorithmic descriptions of all components. In Section V, we present comprehensive performance measurements, and discuss which combinations of primitives are useful for which purpose. The appendices collect formal definitions, provide known extractor results with explicitly spelled out constants that are, in contrast to many discussions that rely on asymptotic notations, vital for an implementation, and give proofs for some new propositions developed in the paper.

II. OVERVIEW

A. What are extractors?

There are numerous possibilities to produce random numbers, and many of them rely on some random physical process, turning, for instance, thermal fluctuations into random bit strings. The laws of physics state that these processes produce distributions with a non-zero entropy, and hence are somewhat random. But the uniform distribution or maximal entropy case is most often not within reach: thermal fluctuations, for instance, require infinite temperatures to produce truly random bit strings. It is therefore necessary to have an algorithm that extracts random numbers from some given initial distribution satisfying a lower bound on its entropy, turning them into uniformly distributed ones. By shrinking the bit string (i.e., reducing the support of its distribution), we increase its randomness until it achieves its maximum. It is easy to see that such a task is impossible for any deterministic routine [2]. But assuming that we have two distributions (seed and source) over bit strings at our disposal, promised to be uncorrelated and fulfilling a lower bound on their entropy, the task comes into reach. Such algorithms are called randomness extractors, and their general structure is shown in Figure 1. The additional randomness is usually taken to be uniform, and is called the seed. A natural aim is to seek algorithms that minimise the required size of the seed, or in other words, the amount of additional randomness. Extractors depend on several parameters, specifying source, seed, and output. This section explains the different parameters and how they are quantified, and discusses their connection. In the second part, we briefly outline Trevisan’s construction.

2 For simplicity we also treat the case of a uniform seed in this work, but some variations of Trevisan’s extractor still apply when the additional randomness just fulfills a lower bound on its entropy [4], and so the methods and code that we have developed can also be adapted to this setting.

1 The sources are available under the terms of the GNU General Public License (GPL), version 2—see www.gnu.org. Essentially, this means that the code can be used and modified free of charge for research (or even commercial) work, provided that any improvements to the code are made available under similar terms.
First, let us consider how to quantify the amount of randomness contained in the source. As per the seminal works of Boltzmann, Shannon and von Neumann, the amount of randomness contained in some distribution of numbers is best quantified by its entropy, traditionally given as \( \sum p_x \log p_x \). Here, \( x \) ranges over all output values, and \( p_x \) is the probability to observe outcome \( x \). This notion of entropy originates from statistical mechanics, where we deal with large numbers of independent entities that are usually also identically distributed. In contrast to that, we are interested in a single run of our extractor, and not in statements about the output distribution obtained from many instances of the extractor applied to many independent copies of the initial distribution. Consequently, we have to alter the notion of entropy.

Intuitively, the amount of randomness in some distribution is quantified by the ability to predict the observed values. This leads to the definition of the guessing probability \( p_{\text{guess}}(X) \) as the probability of correctly guessing the value of the random variable \( X \). It is given by \( p_{\text{guess}}(X) = \max_x p_x \) — the optimal strategy is to guess the most probable value. The bigger \( p_{\text{guess}} \), the less random the source is. This is quantified by the min-entropy, defined by \( H_{\text{min}}(X) = - \log \max_x p_x \).

The definition does so far not consider the possible presence of side information. In a more complex setting, there might be some side information \( E \) correlated to the source \( X \), and the task becomes to extract uniform randomness from \( X \) that is independent of \( E \). In a cryptographic context, \( E \) represents the adversary’s information about the source. Clearly, if \( E \) is a one-to-one copy of \( X \), this task is impossible, even if \( X \) is perfectly uniform. The notions of guessing probability and entropy consequently need to be extended such that they measure the randomness of the source conditioned on \( E \). If the side information is classical, then extractors proven sound in the absence of side information can be used with only a small adjustment of the parameters. However, this changes dramatically if the observer is allowed to use the power of quantum mechanics.

To guess the value of \( X \), a player holding a quantum state in a system \( E \) may measure this system, and make a guess based on the observed outcome. For every value \( X = x \), his quantum memory is in some conditional state \( \rho_x \), and his task reduces to distinguishing the different states \( \rho_x \). Mathematically, such a measurement is specified by a positive operator-valued measure \( \{ E_x \} \). Thus, the probability to correctly guess the value taken by \( X \) is given by \( p_{\text{guess}}(X|E) = \sum_x p_x \text{tr} \rho_x E_x \). The corresponding entropic quantity, the conditional min-entropy, is given by \( H_{\text{min}}(X|E) = - \log \sum_x p_x \text{tr} \rho_x E_x \), where we take \( \{ E_x \} \) to be the optimal measurement.

Having specified the quantification of randomness, we need to define what we mean by an “almost uniform” distribution over the output \( Z = \text{Ext}(X, Y) \), where \( X \) is the source and \( Y \) the seed. Again intuitively, we would like to assure that a player holding some side information cannot do better than with a random guess, that is, the probability that he guesses correctly should be close to \( \frac{1}{2} \), if the output is a bit string of length \( m \). Mathematically, this is expressed by requiring that the joint state of the output and the side information \( \rho_{ZE} \) is close to a product state of a perfectly uniform output, \( \tau \) — the fully mixed state — and the side information \( \rho_E \), that is, we want \( \rho_{ZE} \approx \tau \otimes \rho_E \). The distance between these states is usually denoted by \( \varepsilon \), and referred to as the error of the extractor. Colloquially, an error of \( \varepsilon \) corresponds to a probability of at most \( \frac{1}{2} + \varepsilon \) that the output can be guessed correctly, and a probability of at most \( \frac{1}{2} + \varepsilon \) that any single bit can be guessed.

We are now able to define extractors in more detail. We assume that the input are bit strings of length \( n \) and that the distribution has a conditional min-entropy of at least \( k \). For processing each input string, \( d \) randomly distributed bits may be used. The output should consist of bit strings having length \( m \), and the distribution of outcomes should be \( \varepsilon \)-close to uniform and independent from the side information. We call a deterministic function taking as input the source and the seed and achieving these goals a quantum-proof \((k,\varepsilon)-\text{extractor}\). The output length of such an extractor is \( m \). Naturally, we would like to have \( m \) as close to \( k \) as possible, which means that most of the entropy has been extracted. The value \( k - m \) is therefore called the entropy loss. The extractor is called strong if the output is also close to independent of the value of the seed, or equivalently, the output of the extractor is a pair of bit strings, the first being the value of the random bits used as seed, and the second being

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3 See Lemma [A3] for an exact statement.

4 We use the trace distance to measure how close two states are, see Appendix [A] for an exact definition.

5 See Definition [A2] for a formal definition.
the output. This is exactly the setting needed for the privacy amplification step in quantum key distribution protocols, as both Alice and Bob need to use the same value for the seed, in order to produce a correlated bit string. It is thus assumed that the bit values for the seed are uniformly distributed, but known to the adversary, since they are publicly announced by one of the parties.

The most commonly used (at least in theoretical considerations) strong extractor in quantum key distribution protocols is based on two-universal hash functions \([5, 6]\). A family is a collection of functions that map longer bit strings to shorter bit strings. Over a random choice of the function from the set, two-universality requires that it is extremely unlikely for different bit strings to be mapped to the same output. While universal hashing is optimal in the entropy loss\(^6\) of the required seed length (the size of the function family: as many bits as necessary to randomly select one member) scales as a multiple of \(n\), the input data length (or, in the case of almost universal hashing \([6]\), as a multiple of \(m\), the output length).

It is important to emphasise that strong extractors provide security just based on an entropic assumption, namely the amount of (conditional) min-entropy of the initial distribution. In contrast, pseudo-random number generators are based on complexity theoretic assumptions. For instance, the presumed existence of functions that are hard to invert on average in polynomial time can be turned into an algorithm taking a short random seed and producing an output distribution that “looks” like the uniform distribution to any algorithm running in polynomial time (see Ref. \([10]\) for further information and formal definitions). While such generators greatly outperform our current implementation\(^7\), they require much stronger assumptions and give rise to weaker promises on the output distribution.

After this general discussion on extractors and related issues, we now describe Trevisan’s construction in more detail.

### B. Trevisan’s Construction

Trevisan’s seminal contribution originates in the insight that a certain class of error-correcting codes (ECC), called list-decodable codes \([13]\), can be re-interpreted as extractors. In fact, the codes are one-bit extractors, and deliver a single perfectly random bit from a larger reservoir of slightly random bits. Since an error correcting code is a deterministic mapping from shorter into longer bit strings to make them more robust against the influence of errors acting on the encoded data, the connection between ECCs and bit extractors is not immediately obvious. Trevisan’s first observation was that if we randomly select a position of an ECC’s output string, the corresponding bit is uniformly distributed, provided that the initial distribution has enough min-entropy. If the code outputs bit strings of length \(n = \text{poly}(n)\), a logarithmic long seed of random bits is needed, since exactly \(\log n\) bits are necessary to specify a position of an \(n\)-bit string.

Of course, we are interested in much longer outputs than just a single bit. The second observation of Trevisan states that outputs of many uses of the one-bit extractor can be concatenated so that the output is still uniformly distributed, and that we do not need to choose a completely new set of random seed bits for every use of the one-bit extractor. This is achieved using a construction of Nisan and Wigderson \([14]\), the Nisan-Wigderson pseudo-random generator. The basic idea is that the initial choice of random bits taken from the seed is divided into sets of random bits with small overlap. For example, 100 random bits are divided into 15 sets, each consisting of 10 bits. If the overlap is not too large, there are not too many correlations induced by seeding the elements of each set into the one-bit extractors and concatenating the output bits. The randomness available in the initial distribution can then be used to cope with these additional correlations. Dividing the original seed bits into smaller sets is done using an algorithm called weak design. The complete process is summarised in Figure \(2\).

It turns out that there are many examples of one-bit extractors and weak designs that fulfill the requirements needed for the above procedure to work. Trevisan’s construction is therefore not really a single algorithm, but rather a recipe to combine different one-bit extractors and weak designs to generate a quantum-proof strong extractor. The exact choice of either building block (we also refer to them as primitives in the following) depends very much on the application and on the parameter regime of interest. Consequently, we decided to implement different possible choices and let the operator decide which ones to use. We now present two exemplary use cases that do especially highlight the need to prioritise between speed, entropy loss, and the assumptions on the initial randomness.

Suppose first that we have at our disposal a fast source providing very good random numbers, or equivalently, having a very high entropy. Ideally, we would like to extract all randomness, but since producing new random numbers is fast, we can allow a substantial entropy loss, concentrating on performance instead. In this case, the combination of the GF\((p)\)-weak design with the XOR one-bit extractor is the best choice, achieving a through-

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\(^6\) An extractor will always have an entropy loss \(\Delta \geq 2 \log 1/\varepsilon + O(1)\), where \(\varepsilon\) is the error of the extractor \([9]\).

\(^7\) Practical implementations of pseudo-random number generators, among them the variant used in the Linux kernel \([11]\), rely on cryptographic hash functions like SHA-512 \([15]\). Since these functions, in turn, are used in numerous computing scenarios that extend well beyond cryptography, many recent CPUs offer special-purpose machine instructions that allow for particularly efficient implementations. This makes it practically impossible for an implementation of Trevisan’s construction to beat the throughput of cryptographic hash algorithms that are, besides, much simpler from an algorithmic point of view.
A. Trevisan’s extractor

1. Description

As briefly sketched in the previous section, Trevisan’s construction consists in applying multiple times the same one-bit extractor to the input string, using different weakly correlated seeds for each run. The seeds are chosen as substrings of some longer seed $y \in \{0,1\}^d$. Let $\{S_i\}_i$ be a family of sets such that for all $i$, $|S_i| = t$ and $S_i \subset [d] = \{1,\ldots,d\}$. Then $y_{S_i}$ — the string formed by the bits of $y$ at the positions given by the elements $j \in S_i$ — is a string of length $t$. For a given one-bit extractor $C : \{0,1\}^n \times \{0,1\}^t \rightarrow \{0,1\}$, and such a family of sets $\{S_i\}_{i=1}^n$, Trevisan’s extractor is defined as the concatenation of the output bits of $C$ when used with the seeds $y_{S_i}$, namely

$$\text{Ext}(x, y) := C(x, y_{S_1}) \cdots C(x, y_{S_n}).$$  \hfill (1)

The performance of the extractor naturally depends on the performance of the one-bit extractor, but also on the independence of the seeds used for each run of the one-bit extractor. Intuitively, the smaller the cardinality of the intersections of the sets $\{S_i\}$, the more randomness we can extract form the source, but the larger the seed. The exact condition is given in the following definition.

**Definition III.1** (weak design [15]). A family of sets $S_1, \ldots, S_m \subset [d]$ is a weak $(m, t, r, d)$-design if

1. For all $i$, $|S_i| = t$.
2. For all $i$, $\sum_{j=1}^{i-1} 2^{|S_i \cap S_j|} \leq rm$.

In the following, we refer to the parameter $r$ as the overlap of the weak design.

As an example, if we use a quantum-proof $(k,\varepsilon)$-strong extractor as one-bit extractor and a weak $(m, t, r, d)$-design, the construction given by [15] is a quantum-proof $(k+rm, m\varepsilon)$-strong extractor (see Lemma [B.8]). Thus, if $r = 1$, Trevisan’s extractor has roughly the same entropy loss as the underlying one-bit extractor. Note also that the error $\varepsilon$ of the one-bit extractor is the error per bit for Trevisan’s construction.

2. Constructions overview

We always denote the input length by $n$, and the output length by $m$. We choose $\varepsilon$ in such a way that it

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8 The second condition of the weak design was originally defined as $\sum_{j=1}^{i-1} 2^{|S_i \cap S_j|} \leq r(m-1)$. We prefer to use the version of [15], since it simplifies the notation without changing the design constructions.
corresponds to the error per bit for the final construction. We use $d$ to describe the seed length of Trevisan’s extractor and $t$ for the seed length of the underlying one-bit extractor. $r$ denotes the overlap of the weak design, and $k$ the min-entropy required in the source, which is often expressed as $k = \alpha n$.

In the following, we briefly summarize the constructions described in Sections III.B and III.C. We take the input length $n$, output length $m$, and error per bit $\varepsilon$ to be fixed, and calculate the seed length $d$ and entropy needed in the source $k$ as functions of these three parameters.

a. Weak designs: In Section III.B we describe two weak designs, the first was originally proposed by Nisan and Wigderson [17], and has parameters $d = t^2$ and $r = 2e$ for any prime power $t$ and any $m$. This means that the seed of the final construction is the square of the seed of the one-bit extractor, and the entropy loss induced by the seed of the final construction is the square of the seed length.

In the source $k = \gamma n + rm + 6 \log \left(\frac{1 + \sqrt{2}}{\varepsilon} + \log \frac{4}{3}\right)$, $t = \frac{2\log 2}{h^{-1}(\gamma)} \log n \log \left(2 + \sqrt{2}\right)$, $d = t^2$ or $at^2$, where $\gamma$ is a free parameter that influences the amount of extracted randomness and the length of the initial seed (details in Section III.C), and $a$ is given by Eq. (2).

b. XOR-code: The XOR-code is a one-bit extractor, which simply computes the XOR of a substring of the input. With the two different weak designs, we find that the randomness and seed needed are $k = \gamma n + rm + 6 \log \left(\frac{1 + \sqrt{2}}{\varepsilon} + \log \frac{4}{3}\right)$, $t = \frac{2\log 2}{h^{-1}(\gamma)} \log n \log \left(2 + \sqrt{2}\right)$, $d = t^2$ or $at^2$, where $\gamma$ is a free parameter that influences the amount of extracted randomness and the length of the initial seed (details in Section III.C), and $a$ is given by Eq. (2).

c. Lu’s construction: This one-bit extractor selects a random substring of the input by performing a walk on an expander graph, and then hashes the result to one bit by taking the parity of the bitwise product with a random string. With the two different weak designs, we find that the randomness and seed needed are $k = \gamma n + rm + 6 \log \left(\frac{1 + \sqrt{2}}{\varepsilon} + \log \frac{4}{3}\right)$, $t = \log n + 3c(\ell - 1) + \ell$, $c = \left\lceil \frac{-\log w}{2\log 5\sqrt{2}/8} \right\rceil$, $\ell = \left\lceil \frac{8\log \varepsilon - 8\log(2 + \sqrt{2})}{\log(1 - \nu + w)} \right\rceil$, $d = t^2$ or $at^2$, where $\nu \leq 1/2$ is a free parameter, $a$ is given by Eq. (2), and $w$ is the solution to the equation $w \log w = (1 - \nu + w) \log(1 - \nu + w)$.

d. Polynomial hashing: This constructions uses almost universal hash functions. With the two different weak designs, we find that the randomness and seed needed are $k = rm + 4 \log \left(\frac{1}{\varepsilon} + 6\right)$, $t = 2 \left\lceil \log n + 2 \log \left(\frac{2}{\varepsilon}\right) \right\rceil$, $d = t^2$ or $at^2$, where $a$ is given by Eq. (2).

B. Weak designs

The weak design construction we use (see Section III.B for a description) is originally from Nisan and Wigderson [17], who proved that it is a standard design — a notion stronger than weak designs, originally used by Trevisan [3], but which Raz et al. [15] showed to be unnecessary. Hartman and Raz [16] proved that this construction is a weak $(m, t, r, d)$-design with overlap $r = e^2$ for a prime $t$, $d = t^2$, and $m$ a power of $t$. Ma and Tan [18] improved Hartman and Raz’s analysis, and showed that $r = e$ for any prime power $t$ and any $m$ which is a multiple of a power of $t$. However, for a practical implementation, we need a construction that is valid for any $m$. We prove in Appendix C that this construction is a weak $(m, t, r, d)$-design for any prime power $t$, any $m$, and $r = 2e$.

As mentioned in Section III.A, a larger overlap leads to a larger entropy loss. In Section III.B, we adapt an iterative construction of the basic design from Ma and Tan [18], to construct a new design with $r = 1$. We prove in Appendix C that this construction is correct.

1. Basic construction

In this section we describe a weak design construction, that is, we define a family of sets that satisfy the conditions of Definition III.C.

\[ w < \nu \text{ can actually be chosen freely. The above value minimizes the walks on the expander graph.} \]

9 Hartman and Raz [16, Corollary 2] show that there exist a $d = O(t^2)$ and $r = O(1)$ such that for any $m > t^{\log t}$ the construction is a $(m, t, r, d)$-design, however the restriction $m > t^{\log t}$ and constants in the $O$-notation which depend on $m$ make this unusable in practice. Ma and Tan [18] conjecture that the basic construction is a weak $(m, t, e, t^2)$-design for any $m$, and use this in their implementation. To make up for the lack of proof, they simply count the intersections between the sets $S_i$ after generating the design, to make sure that the overlap is bounded by $e$. [10]
This construction makes use of polynomials over a finite field \( GF(t) \). Every set \( S_p \) is indexed by one such polynomial \( p : GF(t) \to GF(t) \). To construct a weak \((m, t, r, d)\)-design we need \( m \) sets, and therefore \( m \) such polynomials, which we take in increasing order of their coefficients. For example, if \( m = 6 \) and \( t = 2 \), the polynomials are \( \sum_{i=0}^{m} \alpha_i x^i \), with the coefficients \( (\alpha_2, \alpha_1, \alpha_0) \) taken in the following order: \((0, 0, 0), (0, 0, 1), (0, 1, 0), (0, 1, 1), (1, 0, 0), (1, 0, 1)\). In general, the \( n \)-th polynomial is given by \( p(x) = \sum_{i=0}^{m} \alpha_i x^i \), with \( \alpha_i = (n-1)/t^i \) mod \( t \) and \( c = \lceil \log m \rceil - 1 \).

The elements of the set \( S_p \) are all the pairs of values \( S_p := \{(x, p(x)) : x \in GF(t)\} \). Each set thus has \(|S_p| = t\) elements, and \( S_p \subset [d] \) holds for \( d = t^2 \), where we map \([d]\) to \([t] \times [t]\) in the obvious way. We prove in Appendix C.1 that for all \( m \) and \( p \), this construction has \( \sum_{(q < p)} 2^{S} \| \leq 2em \), where by \( \{q < p\} \) we mean the set of all polynomials that come before \( p \).

2. Reducing the overlap

Note that any weak design can be viewed as a binary \((m \times d)\)-matrix \( W \), where the value \( w_{ij} = 1 \) if \( j \in S_i \). To construct a weak design with \( r = 1 \), we will use the construction from Section II.B repeatedly with different values \( m \), but the same \( t \), obtaining designs \( W_{B,0}, \ldots, W_{B,ℓ} \). We then construct a new design \( W \) by placing these in its diagonal, that is, 

\[
W = \begin{pmatrix}
W_{B,0} & & \\
& \ddots & \\
& & W_{B,ℓ}
\end{pmatrix}.
\]

Let \( m \) and \( ℓ \) be fixed, and let \( r' = 2r \) be the parameter from the basic construction. The number of calls to the basic construction is given by 

\[
ℓ := \max \left\{ 1, \left\lfloor \frac{\log(m-r') - \log(t-r')}{\log r' - \log(t-1)} \right\rfloor \right\}.
\]

And each design \( W_{B,ℓ} \) is constructed with \( m \), sets, defined as follows:

\[
n_i := \left(1 - \frac{1}{r'}\right)^i \left(\frac{m}{r'} - 1\right), \quad \text{for } 0 \leq i \leq ℓ - 1,
\]

\[
m_i := \sum_{j=0}^{i} n_j - \sum_{j=0}^{i-1} m_j, \quad \text{for } 0 \leq i \leq ℓ - 1,
\]

\[
m_ℓ := m - \sum_{j=0}^{ℓ-1} m_j.
\]

The weak design \( W \) thus has \( d = (ℓ+1)t^2 \). In Appendix C.2 we prove that this construction has \( r = 1 \).

Figure 3 discusses the parameter behaviour of the block weak design.

C. One-bit extractors

1. XOR-code

This extractor computes the XOR for \( ℓ \) random positions of the input, it is thus an \( ℓ \)-local extractor (see Appendix A for a precise definition). This construction is efficient to compute, but requires a seed of length \( t \in O(\log n \log 1/ε) \), where \( n \) is the input length and \( ε \) the error of the construction, instead of the optimal \( O(\log n + \log 1/ε) \). It also has an entropy loss linear in the input length.

Lemma III.2 (XOR-code \cite[Theorem 41]{BrzSpil20}). For any \( ε > 0 \), \( n \in \mathbb{N} \) and \( ℓ \in [n] \), the function

\[
C_{n,ε,ℓ} : \{0,1\}^n \times [n]^ℓ \to \{0,1\}
\]

\[
(x, i_1, \ldots, i_ℓ) \mapsto \bigoplus_{j=1}^{ℓ} x_{i_j}
\]

\cite[Theorem 41]{BrzSpil20} actually proves that this construction is a \( δ \)-approximately \((ε, L)\)-list-decodable code. But such a code is an \((ε, L2(δ)n)\)-list-decodable code, which in turn is a classical-proof extractor by Lemma B.3.\footnote{\cite[Theorem 41]{BrzSpil20} also proves this construction is an \( ε \)-injective \((δn)\)-tilde-decodable code, which in turn is a classical-proof extractor by Lemma B.3.}
is a classical-proof \( l \)-local \((k, \varepsilon)\)-strong extractor with \( k = h\left(\frac{\ln 2}{\ln(1+\sqrt{2})}\right)n + 3\log \frac{1}{\varepsilon} + \log \frac{1}{\delta} \) and seed length \( t = \ell \log n \), where \( h(p) = -p \log p - (1-p) \log(1-p) \) is the binary entropy function.

By Lemma 3.7, this construction is a quantum-proof \((k, (1+\sqrt{2})\sqrt{\varepsilon})\)-strong extractor. And by Lemma 3.8 if we use this in Trevisan’s construction, the final extractor is a quantum-proof \((k + r_m, m(1 + \sqrt{2})\sqrt{\varepsilon})\)-strong extractor.

Let our source have min-entropy \( H_{\min}(X|E) = \alpha n \). We want the entropy loss induced by this one-bit extractor to be roughly \( \gamma n \), and need to find the appropriate \( \ell \) for the desired value of \( \gamma \) since \( \gamma(\ell, \varepsilon) = h\left(\frac{\ln 2}{\ln(1+\sqrt{2})}\right) \) for some \( \gamma < \alpha \). Solving for \( \ell \), we find \( \ell(\gamma, \varepsilon) = \frac{\ln 2}{h^{-1}(\gamma)} \log \frac{2}{\varepsilon} \).

This implies that \( \gamma \) directly influences the length of the seed, which we discuss below. Since the inverse binary logarithm \( h^{-1}(\cdot) \) is not analytically available, we need to resort to numerical techniques to determine the appropriate value of \( \ell \) for a given \( \gamma \). It is convenient to distinguish the experimental entropy deficiency \( \alpha \) from the loss induced by the extraction procedure by introducing a parameter \( \mu \) such that \( \gamma = \mu \alpha \).

For \( \varepsilon = \frac{\gamma^2}{(1+\sqrt{2})^2} \), Trevisan’s construction is a quantum \((k, \varepsilon^\prime m)\)-strong extractor with \( k = \gamma n + rm + 6 \log \frac{1+\sqrt{2}}{\varepsilon} + \log \frac{1}{\delta} \). The seed of the one-bit extractor has length \( t = \ell \log n = \frac{\ln 2}{h^{-1}(\gamma)} \log n \log \frac{2(1+\sqrt{2})}{\varepsilon} \), and the seed of the complete construction has length \( d \).

Especially the choice of \( \mu \) influences the behaviour of the XOR extractor. Figures 4, 5, and 6 depict and discuss the effect of the various chosen and inferred parameters.

2. Lu’s construction

Lu [20] shows how to construct a local one-bit extractor, i.e., an extractor for which each bit of the output only depends on a subset of the input bits. He then uses his one-bit extractor in Trevisan’s construction. Here, we adapt the parameters of his construction to build a quantum-proof extractor.

Lu’s extractor proceeds in two steps. The first consists in selecting a substring of the input; the second hashes this string to one bit [12]. To select the substring of the input, he performs a random walk on a \( g \)-regular graph—a graph in which every vertex is connected to exactly \( g \) other vertices.

Recall that a graph \( G \) is uniquely identified by its vertices and edges, and is consequently specified by \( G = (V, E) \), where \( V \) is the vertex set and \( E \) the edge set. An alternative representation of more importance in our context is the adjacency matrix. For a graph with \( n \) vertices,

\[\text{XOR extractor: Scaling behaviour}\]

this is an \( n \times n \) matrix in which the entry \( a_{ij} \) denotes the number of edges from vertex \( i \) to vertex \( j \). The diagonal is typically filled with ones; since the graphs considered here are undirected (i.e., the direction of edges is not taken into account, only the fact that two vertices are connected), the adjacency matrix is symmetric.

The eigenvalues of the adjacency matrix are referred to as eigenvalues of the graph. For our purpose, the ratio between the second largest and largest eigenvalue plays an important role, and is labelled as \( \lambda \). Graphs with a small \( \lambda \) are called expander graphs, and are common objects in pseudo-randomness generation, see Ref. 22 for a review.

For an input string of length \( n \), we choose a graph with \( n \) vertices, so that each vertex corresponds to a bit position of the string. Let \( (v_1, \ldots, v_\ell) \) be the vertices visited during a walk of \( \ell \) steps. We select the \( \ell \) corresponding bits of the input \( x \), that is, \( (x_{v_1}, \ldots, x_{v_\ell}) \), and then hash it by computing the parity of the bitwise product of this string with a random seed \( \beta \in \{0,1\}^\ell \). The output is thus \( z = \bigoplus_{i=1}^{\ell} \beta_i x_{v_i} \).

Lu [20] proves that the concatenation of the output bits \( z \) for all possible seeds is a \((\delta, L)\)-list decodable code.

\[12\] This type of construction is sometimes referred to as sample-then-extract [21], although Lu [20] simply describes it as a local list-decodable code.

\[13\] This hash function is also used in Section III C 3.
FIG. 5. Ratio between output size and seed length for various input sizes. The parameters $\mu = 0.05$ and $\varepsilon = 10^{-7}$ are fixed for this computation. A ratio of 1, indicated by a dashed red line, denotes the parameter regime where the amounts of extracted bits and the seed spent coincide; for values exceeding this threshold, the particular advantages of Trevisan’s construction over two-universal hashing prevail because the ratio is better than what can be achieved with the latter approach. Recall, though, that the seed acts as a catalyst that can be included in the final randomness since the extracted bits are independent of the seed.

The initial source entropy $\alpha$ accounts for a variation of about one order of magnitude of the extraction threshold. As a rule of thumb, the break-even point is at input sizes of roughly $10^9$ bits, which amounts to approximately $2^{30}$ bytes (roughly 1 GiB) of data.

The inset shows the number of extracted bits less the seed spent.

With

$$L = \frac{2 h (\nu) n}{26^2},$$

(5)

for a $\nu \leq 1/2$ given by

$$\nu = 1 + \lambda^2 - \delta^2.$$

(6)

Since $\delta^2 < 1$, $\frac{2 h (\nu) n}{26^2}$ can only be satisfied if $\lambda^2 < \nu$. This can be obtained by taking as expander graph $G$ a given construction $G_0$ to the power $c$. $G$ is defined as the graph with adjacency matrix $A = A_0^c$, where $A_0$ is the adjacency matrix of $G_0$. We then have $\lambda = \lambda_0^c$. A random walk of length $\ell$ on $G_0^c$ is equivalent to a random walk of length $c \ell$ on $G_0$, in which only the first of every $c$ steps is remembered, and the others deleted [23].

To construct the regular expander graph $G_0$, we employ an algorithm reviewed in Ref. [22]. Let us only summarise the essential facts here:

- The construction is restricted to degree $g = 8$, and the ratio between the second-largest and largest eigenvalue can be shown to be $\lambda = 5\sqrt{2}/8 \approx 0.884$.

- It is possible to compute the graph for all dimensions (i.e., number of nodes) that can be expressed as $\ell^2$ for $\ell \in \mathbb{N}$. This restriction is much more relaxed than for other constructions, and does not pose any problems in real applications. Formally, the vertex set of the graph is defined on $\mathbb{Z}_\ell \times \mathbb{Z}_\ell$. Each vertex $(x, y) \in \mathbb{Z}_\ell \times \mathbb{Z}_\ell$ is connected to the vertices $(x \pm 2y, y)$, $(x \pm (y + 1), y)$, $(x, y \pm 2x)$, and $(x, y \pm (2x + 1))$, which uniquely defines the edges. Notice that the arithmetic must be performed modulo $\ell$, so the computationally (comparatively) cheap additions and multiplications are unfortunately accompanied by an expensive modulo division [23].

- The complete graph does not need to be computed in advance, but can be constructed during the random walk, and using a constant amount of space.

An obvious optimisation possibility that is available because the multiplicative factor 2 is small is to compute the modulo division not unconditionally, but only when the intermediate result really exceeds $\ell$. 

FIG. 6. Break-even points (i.e., minimal input length for which the amount of extracted randomness exceeds the required seed size) for varying values of $\mu$ and $\varepsilon$. The parameter $\alpha$ is fixed to 0.8. As a rule of thumb, $\mu = 0.05$ is close to the optimal value irregardless of the error parameter $\varepsilon$. 

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For a given \( \nu \), we choose \( c \) and \( \ell \) which minimize the number of steps \( c\ell \). By setting \( w := \lambda^{2\nu} \) and taking the derivative of \( c\ell \) with respect to \( w \), we find that minimum is obtained for the \( w \) which is the solution of the equation

\[
w \log w = (1 - \nu + w) \log(1 - \nu + w).
\]

The number of steps on the expander graph is then given by \( c = \left\lfloor \frac{\log w}{2 \log \lambda_0} \right\rfloor \) and \( \ell = \left\lfloor \frac{4 \log \delta}{\log(1 - \nu + w)} \right\rfloor \).

The walk on the \( g \)-regular graph requires \( nh(n) \) bits of seed to choose the first vertex, and \( c(\ell - 1) \log g \) bits for the direction of the walk for each following step. The final hashing uses \( \ell \) bits of seed, for a total of \( t = nh(n) + c(\ell - 1) \log g + \ell \).

From Lemma B.3 and [6], Lu’s one-bit extractor is a classical-proof \((h(\nu)n + 3 \log \frac{1}{\delta} - 2, 2\delta)\)-strong extractor. By Lemma B.5 it is quantum-proof \((h(\nu)n + 3 \log \frac{1}{\delta} - 2, (2 + \sqrt{2})\sqrt{\delta})\). And from Lemma B.8 when used with a weak \((m, t, r, d)\)-design, Trevisan’s construction is a quantum-proof \((k, m\varepsilon)\)-strong extractor with \( k = h(\nu)n + rm + 6 \log 2 + \sqrt{2} - 2 \).

Unfortunately, Lu’s construction is not useful in a practical setting owing to its unfortunate parameter scaling: The number of random walk steps increases considerably with decreasing parameter \( \nu \), see Figure 4. However, as Figure 8 shows, small values of \( \nu \) are required for even tiny extraction fractions. Overall, this makes the construction reach parameter realms where it is preferable over two-universal hashing functions (namely, when the length of the extracted bits exceeds the amount of initial seed) only rarely, as Figure 9 shows.

### 3. Polynomial hashing

Renner [5] proved that universal hash functions are good extractors. Tomamichel et al. [6] showed that the same holds for \( \delta \)-almost universal \((\delta-AU)\) hash functions, given that \( \delta \) is small enough. For the range of \( \delta \) that build good extractors, almost universal hashing requires a seed of length \( \Omega(m + \log n) \), where \( n \) is the input and \( m \) the output length. This seed is too large for many applications; however in the case of one-bit extractors, this reduces to \( \Omega(\log n) \), and is achievable with the construction we describe here.

This construction is in fact the concatenation of two hash functions, and uses a seed of length \( 2\ell \), where \( \ell \) will be specified later. The first is known as polynomial hashing — or alternatively as a Reed-Solomon code, because the concatenation of the hashes for all seeds corresponds to the encoding of the input with a Reed-Solomon code. We partition the input string \( x \in \{0, 1\}^n \) in blocks \( x = (x_1, \ldots, x_\ell) \), each of length \( \ell \) (if necessary, we pad the last string \( x_\ell \) with \( 0s \)). We view each block as an element of a field \( x_i \in GF(2^\ell) \), and evaluate the polynomial

\[
p_\alpha(x) = \sum_{i=1}^s x_i x^{\alpha^{s-i}},
\]

---

15 See Appendix B.3 for a definition of (almost) universal hash.
where $\alpha \in \text{GF}(2^\ell)$ is the first half of the seed. This family is $\frac{\alpha}{2^\ell\cdot2}$-AU$_2$. [23]

Since the $\delta$ of polynomial hashing is too large (relative to the output length) to build an extractor, we combine it with another hash function—sometimes referred to as a Hadamard code, as the concatenation of the outputs over all seeds corresponds to the Hadamard encoding. This hash function computes the parity of the bitwise product of $p_a(x)$ and the second half of the seed, $\beta \in \{0, 1\}^\ell$. The output is thus $z = \bigoplus_{i=1}^{\ell} \beta_i p_a(x)_i$. Since this hash function is $\frac{1}{2}$-AU$_2$, by [23] Theorem 5.4 the combination of the two is $\delta$-AU$_2$ with $\delta = \frac{1}{2} + \frac{s}{2\ell}$.

Choosing $\ell = \lceil \log n + 2 \log 1/\varepsilon' \rceil, s = \lceil n/\ell \rceil$ we get

$$\delta = \frac{1}{2} + s - \frac{1}{2\ell} < \frac{1}{2} + \frac{n}{2\ell} \leq \frac{1}{2} + \varepsilon^2.$$  

From Theorem [B.7] this is a quantum-proof $(4 \log \frac{1}{2} + 2, 2\varepsilon')$-strong extractor. And plugging this in Trevisan’s construction with a $(m, 2\ell, r, d)$-design and $\varepsilon = 2\varepsilon'$, we get from Lemma [B.8] a quantum-proof $(4 \log \frac{1}{2} + 6 + rm, m\varepsilon)$-strong extractor. The seed of the one-bit extractor has length $l = 2\ell = 2(\log n + 2 \log 2/\varepsilon')$, and the seed of the complete construction has length $d$.

Figure 10 discusses the parameters of the polynomial hashing extractor.

![Figure 9](image.png)

**FIG. 9.** Comparison of seed size and output size for Lu’s construction. The interesting regime is only reached in very rare cases.

![Figure 10](image.png)

**FIG. 10.** Polynomial hashing parameter overview (calculations are for $r = 2e$). The parameters are easy to evaluate because there is no dependence on $\alpha$, and there is also no extraction factor $\mu$—the extractor works equally well for high- and low-entropy sources. The required seed is consistently small (shown in the bottom inset); it increases linearly as $\varepsilon$ decreases exponentially.

The degree of the polynomial that needs to be evaluated is the crucial factor. Even for small inputs like $n = 10^8$, corresponding to roughly 1 MiB of data, the degree is $\approx 10000$. Since the polynomial needs to be evaluated for every extracted bit, this makes the polynomial hashing extractor an unsuitable choice for performance intensive scenarios. The top inset shows the regime in which the extractor delivers more bits than initially invested for the seed. It outperforms two-universal hashing for a very wide range of parameters.

IV. IMPLEMENTATION

A. Implementation Architecture

We now turn our attention to describing the implementation of the Trevisan extraction framework by first outlining the software architecture, that is, the high-level conceptual point of view, followed by a discussion of some important implementation details and notes on how to add new primitives to the infrastructure. While many important details are still omitted for the sake of brevity, the full source code is available at [https://github.com/wolfgangmaurerer/libtrevisan](https://github.com/wolfgangmaurerer/libtrevisan) for inspection and modification. Besides instructions on how to build the code, the website also contains detailed information on how to use the program, which we will not discuss here any further.
1. Architecture

The architecture was designed to satisfy two particular constraints: Correctness and maximum throughput. To achieve the latter, we use C++\(^{16}\) to implement all performance-critical parts, since the language is statically compiled and does not require any intermediate layers that add runtime penalties to interpreted or byte-compiled languages like, for instance, Matlab, but still allows us to maintain a clean and extensible design based on modern software engineering techniques\(^{20}\). The implementation is portable across a wide range of machines from laptops to high performance computing (HPC) machines, and also provides opportunities to benefit from low-level capabilities of recent CPUs, for instance to accelerate bit-level manipulations. We have tested the code on Linux and MacOS machines.

To ensure correctness of the calculations, we base the implementation on independent libraries (NTL\(^{27}\) and OpenSSL\(^{28}\) for working with finite fields of arbitrary size) that can be selected at compile time\(^{14}\). Checking that both variants arrive at the same results for identical parameter sets increases the faith in the reliability of the calculations. Another means to ensure code correctness is given by a large number of invariants and sanity checks that are spread all across the implementation. To not compromise the performance goals, it is possible to deactivate the checks at compile time so that they incur no runtime penalty.

Another major design decision is the focus on multi-core machines: Nowadays, machines with only a single core are a rare exception, and algorithms that are limited to only one thread of execution voluntarily sacrifice a large fraction of the available computational power, which is obviously not desirable in a high-performance setting. We use the threading building blocks library\(^{29}\) as basis for the implementation, which allows for fine-tuning the distribution of work across the system resources in a precise manner. We also employ a mostly lock-free architecture (see, e.g., Ref.\(^{30}\) for a review) that avoids any computation stalls due to the need for synchronised communication between computation elements.

The code also contains parts that are not performance-critical, for instance calculating the parameters from given user settings. This is conveniently done in very high-level languages that allow for working in abstract terms without having to consider any details of the underlying machine architecture. To this end, we have integrated the possibility to call code written in the R language (using the techniques provided by Ref.\(^{31}\); see\(^{32}\) for an overview about R), which enjoys widespread use in statistical data processing and machine learning.

It is also possible to compute the weak design ahead of time, store it on disk and re-use it for multiple runs of the extractor — since computing the weak design is a deterministic operation that does not require any randomness, this is admissible to do. In matrix representation, a weak design for output length \(m\) and a total seed length \(d\) is an element \(\mathbb{F}_2^{t \times d}\). Each row contains \(t\) ones and \(d-t\) zeroes, so the matrix fill for the standard design is \(\frac{mt}{m\left(d-t\right)} \approx \frac{1}{t}\).

A total seed of 50 KiBit, for instance, amounts to a fill of about 0.5%, which exceeds the threshold for typical sparse matrix techniques to pay off\(^{33}\). We found the data transfer times from the underlying block device to be larger than the time required to compute the weak design on the fly, albeit this may change with the availability of high-speed storage. For the block weak design, the situation is more favourable since only the basic design needs to be stored, and the remaining elements can be reconstructed with very little computational effort.

Finally, we emphasise that the code can either be used in stand-alone mode (also including a dry-run mode for parameter estimation), or as a library as part of a larger project.

2. Implementation details

Weak designs and one-bit extractors are implemented as C++ classes derived from mixed interface/implementation-type base classes. Trevisan’s algorithm solely operates on the base class objects using dynamic polymorphism, and does not require any knowledge about the internal structure of the primitives.

The source code contains full information on how to implement and integrate new primitives, so we only summarise briefly what methods need to be provided.

Weak designs need to be derived from \texttt{class weakdes}, and must implement

- \texttt{compute_S1(uint64_t i, vector indices)} — compute the \(i\)th index set, and store the results in \texttt{indices}.
- \texttt{compute_d()} — compute the required amount of initial seed.
- \texttt{get_r()} — report the overlap \(r\) to the higher-level algorithms.

Optionally, the function \texttt{set_params(uint64_t t, uint64_t m)} can, but need not be implemented to initialise the parameters required for all weak designs.

---

\(^{16}\) We rely on numerous features of the new language standard C++11, so at the time of writing, only sufficiently new compilers are able to build the code.

\(^{17}\) NTL cannot be used in scenarios with high performance requirements since it is restricted to running on one single core per design, which does not agree well with contemporary machine architectures. It can only be used in a single primitive that requires operations on \(\text{GF}(2^t)\) because the library operates with a single, global irreducible polynomial, which makes it effectively impossible to operate on fields of different dimensions simultaneously.
Determining \( d \) from \( t \) seems straightforward, but is accompanied by constraints—the GF\((2^d)\) based weak design, for instance, only works for values of \( t \) that can be represented as a power of 2, so the design typically needs to choose larger values (resulting in more initial seed) than requested.

One-bit extractors need to be derived from class `bitext`, and must implement

- `num_random_bits()` — compute the amount \( t \) of initial seed bits required for every extracted bit.
- `compute_k()` — determine the minimal source entropy required by the extractor for the parameter set under consideration.
- `extract(void *initial_rand)` — extract one bit using the provided subset of the initial randomness.

There are also generic functions to assign global randomness and other generic parameters to the 1-bit extractor. They can, but need not be provided by an implementation.

On the lower layers, the implementation was designed to use elementary machine arithmetic (as opposed to software-based multi-precision arithmetic) whenever possible; this is an obvious precondition for an implementation with good performance. In all performance critical operations, logarithms are not computed using floating point, but with integer operations since usually only floor operations are required, which the code assumes to be the default setting.

The code uses a fixed-width integer data type with 64 bits to represent potentially large quantities like the number of input bits. It is important to note that the width of the index data type sets an upper bound on the amount of randomness that can be handled by the code, namely to \( 2^{w-3} \) bytes (for \( w = 32 \) respectively \( w = 64 \)), which corresponds to \( 2^w \) bits (the datum is used as an index into a bit field, and this field need not be representable by a machine quantity). Since contemporary 64-bit machines cannot handle more than \( 2^{48} \) bytes owing to virtual address space management limits [11], the choice does not introduce any additional limits. To process large amounts of randomness (multiple gigabytes), 64 bit machines and a 64-bit kernel running on the machine are required, which the code assumes to be the default setting.

### B. Algorithms

In the following, we give a concise description of all algorithms in a form that is helpful for actual implementations—in some contrast to the previously given descriptions that focus more on mathematical clarity, we provide recipes in a pseudo-formal language that is close enough to many contemporary imperative and object-oriented programming languages, yet still sufficiently abstract to avoid hiding the algorithmic core behind technical side-work. Although each algorithm can be captured with very few statements, we remark that a practical implementation needs to account for many non-trivial technical issues; our reference implementation published as a part of this paper comprises about 5000 lines of source code.

#### 1. Trevisan’s extractor

The Trevisan algorithm is independent of the type of weak design and bit extractor used; only the inferred parameters depend on the specific properties of the components:

```plaintext
1: procedure Trevisan(\( \mathcal{WD} \), \( \mathcal{Ext} \), \( n, m, \mu, \alpha, \varepsilon, \varphi^0, \varphi^d \))
2: \( t \leftarrow \mathcal{Ext}.InputSize(n, m, \mu, \alpha, \varepsilon) \)
3: \( d \leftarrow \mathcal{WD}.InputSize(t) \)
4: Save space for \( m \) bits in \( \varphi^0 \)
5: Save space for \( t \) numbers \( \in [d] \) in \( S \)
6: for \( i \leftarrow 0, m − 1 \) do ▷ Data parallel
7: \( S \leftarrow \mathcal{WD}.computeS(i) \)
8: \( b \leftarrow 0 \)
9: for \( j \leftarrow 0, t − 1 \) do ▷ Indices refer to bits
10: \( b_j \leftarrow \varphi^0_{S_j} \)
11: end for
12: \( \varphi^0 \leftarrow \mathcal{Ext}.extract(b, \varphi^d) \)
13: end for
14: return \( \varphi^0 \)
15: end procedure
```

The components \( \mathcal{WD} \) and \( \mathcal{Ext} \) may impose boundary conditions on the parameters; for instance, the single-bit seed length \( t \) must be a power of a prime number for the weak designs implemented in this paper.

#### 2. Weak Designs

##### a. Construction of Hartman and Raz

The weak design of Hartman and Raz is based on evaluating polynomials over finite field; recall from Section [11][A2] that the dimension of the field needs to be a power of a prime number. We have implemented two variants: One based on the extension field \( \mathbb{F} = \text{GF}(2^d) \), and one based on the prime field \( \mathbb{F} = \text{GF}(p) \). The bit extractors can require arbitrary values of \( t \) that are not necessarily compatible with the constraints of the weak design. In this case, \( t \) needs to be increased to the next possible value \( t' \) that can be provided by the weak design. Consequently, we need to distinguish between \( t \), which represents the value that can be provided by the weak design, and \( t^{req} \), which is the value originally requested by the bit extractor. It necessarily holds that \( t \geq t^{req} \).

The basic algorithm for both finite fields is as follows (indices in square brackets denote bit selections):

```plaintext
1: procedure \( \text{HR}\).ComputeS(\( \mathbb{F}, i, m, t \))
2: \( c \leftarrow \left\lceil \frac{\log m}{\log t} \right\rceil + 1 \)
```
3: for $j \leftarrow 0, c$ do \(\triangleright\) Prepare polynomial coefficients
4: \(\alpha_j \leftarrow [j \cdot \text{nb}(t), j + \text{nb}(t) - 1] \mod t\)
5: end for

6: for $a = 0, a < 2^m$ do
7: \(b \leftarrow \sum_j \alpha_j a^j\)
8: \(S^a_t[\log(t), 2 \cdot \log(t) - 1] \leftarrow b\)
9: \(S^a_t \leftarrow S^a_t \cdot a\)
10: \(S^a_t \leftarrow S^a_t \mod |F|\)
11: end for

12: return $S$
13: end procedure

For a field of prime dimension $p$, all calculations are performed modulo $p$. Notice, though, that it is not sufficient to simply divide by $p$ after any multiplication (or addition/subtraction) has been performed, because this can easily lead to intermediate results that exceed the maximal bit width available in hardware. Multiplying two 40-bit numbers, for instance, can result in an 80-bit value, which exceeds the word size of 32 and 64 bit machines. A naïve solution could fall back to using arbitrary-precision software arithmetic, which is unfortunately much slower than native machine hardware arithmetic. Consequently, we use have made sure to use algorithms that avoid intermediate overflows and can work with multiplicands of up to 61 bits, which is sufficient for our purposes. See the source code or Ref. [34] for details.

For the extension field $GF(2^m)$, it is not sufficient to perform a simple division of arithmetic results by a scalar to satisfy the constraints of the finite field. Instead, all elements of the field are formally interpreted as polynomials over the binary field, and arithmetic operations are performed modulo an irreducible polynomial that needs to be constructed dependent on the field order. It can be shown (see, e.g., Ref. [27]) that for every field order, an irreducible polynomial of order 3 or 5 exists, so calculations can be optimised for these cases.

b. Block Weak Design The block weak design is based on a basic design whose matrix representation is re-used multiple times as part of the total weak design—once the matrix representation of the basic design is known, it is possible to construct the complete design by placing sub-matrices of the basic design matrix on the diagonal of a larger matrix. One possible implementation could thus use sparse matrix techniques to store the basic design in memory, and derive all other blocks from this representation.

When the basic design is not represented by a matrix, but as vectors of indices, it is possible to compute the content of $W_{B,j}^k$ from the basic design row $W_{B,0}^k$ by adding $j \cdot t^2$ to all values of the set $S$ corresponding to the matrix row. Since it is possible to re-arrange the rows of $W$ without changing the properties of the weak design, we use a suitable permutation (derived from the data in Eq. [4], see the source code for details) of the rows of $W$ such that all rows that originate from the same row of the basic design are adjacent to each other, which allows us to cache calls to the basic construction. Since the design is traversed from row to row in the Trevisan algorithm, the permuted row order minimises calls to the basic construction.

1: procedure BWD.computeS($WD, i, i^c, S^c, t$)
2: \(\text{Infer } j, k \text{ from } i\)
3: if $k \neq i^c$ then
4: \(i^c \leftarrow k\)
5: \(S \leftarrow WD.\text{computeS}(i^c)\)
6: for $\zeta \leftarrow 0, t - 1$ do \(\triangleright\) fill cache
7: \(S^c_\zeta \leftarrow S^c_{\zeta}\)
8: end for
9: else
10: for $\zeta \leftarrow 0, t - 1$ do
11: \(S^c_\zeta \leftarrow S^c_{\zeta} + j \cdot t^2\)
12: end for
13: end if
14: return $S$
15: end procedure

3. 1-Bit extractors

Finally, we discuss the algorithms used for the 1-bit extractors implemented as part of this paper.

a. XOR Code An implementation of the XOR code requires to derive the parameter $l$ from the experimental parameters; since this can be achieved by a standard numerical optimisation, we will not discuss a formal algorithm here, but refer the reader to the source code for the details. The algorithm itself is compact:

1: procedure XOR.extract($\varphi^d, \varphi^s$)
2: \(r \leftarrow 0\)
3: for $i \leftarrow 0, l - 1$ do
4: \(\zeta \leftarrow \varphi^d[i \cdot \text{nb}(n - 1), (i + 1) \cdot \text{nb}(n - 1) - 1]\)
5: \(r \leftarrow r \oplus \varphi^s[\zeta]\)
6: end for
7: return $r$
8: end procedure

b. Polynomial Hashing The algorithm to perform polynomial hashing based on a concatenation of a Reed-Solomon and a Hadamard code is as follows:

1: procedure RSH.extract($\varphi^d, \varphi^s, n, \varepsilon$)
2: \(\zeta \leftarrow 0, l \leftarrow \lceil \log n + 2 \log 2/\varepsilon \rceil\)
3: \(s \leftarrow \lceil n/l \rceil\)
4: Pick irreducible polynomial for $GF(2^t)$
5: for $i \leftarrow 0, s - 1$ do \(\triangleright\) Determine coefficients
6: \(c_i \leftarrow \varphi^d[i \cdot l, (i + 1) \cdot l - 1]\)
7: end for
8: \(\alpha \leftarrow \varphi^d[0 : l - 1] \quad \triangleright\) Reed-Solomon step
9: \(r \leftarrow \sum_s c_i \alpha^{s-i} \quad \triangleright\) Computed over $GF(2^t)$
10: \(b \leftarrow 0 \quad \triangleright\) Hadamard step
Most of the implementation complexity for the Lu expander stems from the need to select subsets of bit strings. To simplify distributing the initial randomness provided by the weak design into three components as shown above, the actual implementation assumes that the contributions start on indices that are evenly divisible by the bit width of the data type used to represent edges. This simplifies the implementation, but implies that a slightly larger amount of randomness than theoretically possible is required, albeit the increase is only by a negligible additive factor.

V. RUNTIME COMPARISON

Owing to the many aspects—throughput, scalability, weak design versus extractor performance, parameter ranges, machine characteristics, among others—involved in determining code performance, and because of the large number of combinations of primitives, it is neither possible nor reasonable to present measurements for all cases (since the full sources are available, measurements for a particular case of interest can be easily conducted by interested parties). Instead, we focus on a selection of measurements that describe cases of typical experimental interest. We use two machines to run the tests; detailed technical specifications are shown in Table I.

| Machine         | CPU                     | # CPUs | Cores/CPU | Threads/core | Σ Threads | RAM [GiB] | Kernel   | Clock Speed |
|-----------------|-------------------------|--------|-----------|--------------|-----------|-----------|----------|-------------|
| Laptop          | Intel Core i5           | 6      | 1         | 2            | 4         | 48        | Linux 3.0 | 1.6 Ghz     |
|                 | AMD Opteron             | 4      | 2         | 4            | 4         | 32        | Linux 3.0 | 1.9 Ghz     |

*a Pairs of two CPUs share one socket*

TABLE I. Machines deployed in the benchmark measurements.

The measurement results are shown in Figures 13, 14, and 15 refer to the captions for a detailed discussion of the results.
FIG. 11. Scaling behaviour of RSH with a block design for varying input lengths. For a small number of CPUs, performance degrades considerably with increasing input length, as expected for a non-local extactor. Good throughput (more than 100 kbit/s) is only obtained for very small input sizes ($2^{12}$ is only 4KiBit of data!) for which the required amount of initial seed drastically exceeds the extracted amount of randomness.

With many cores, the achieved speed-up does initially not compensate the overhead for setting up and performing parallel operations, so the throughput increases to a local maximum, and then decreases as expected with larger input lengths. Consequently, it is not just sufficient to add more CPUs for a given scenario to increase throughput; practical book-keeping tasks and technical aspects can easily dominate the actual problem. In particular, this implies that purely technical improvements like porting the processing to massively parallel approaches like GPU computing will not automatically resolve all performance needs; a proper choice of primitives for given requirements is essential, which is only possible with a framework that allows for flexibly combining these primitives.

FIG. 12. Throughput comparison of our results (obtained on a laptop, represented by boxplots) with the results obtained by Ma et al. (represented by triangles) for the combination of primitives supported by their implementation. Since the code of [7] seems to be limited to running on one CPU core, we have also included an artificially constrained measurement measurement for the code discussed in this paper. Generally, our framework is 2–3 orders of magnitude faster in terms of throughput, and allows for dealing with inputs that surpass Ref. [7] by many orders of magnitude.

APPLICATIONS

We have presented a modular, scaleable implementation of Trevisan’s construction for randomness extraction, together with detailed parameter derivations and improved mathematical proofs. We have shown that the feasibility or non-feasibility of Trevisan’s scheme is not mainly a question of computational complexity issues, but does depend on the particular choice of primitives used as components of the algorithm; different scenarios require different constituents. Although our measurements indicate that there exist use cases that require theoretical improvements to make Trevisan’s construction applicable (mostly because short-seed extractors all suffer from a low extraction rate), the implementation can, for instance, satisfy the needs of all current quantum key distribution schemes. The authors hope that the public availability of the source code, together with the extensible architecture, will spawn contributions from other researchers to turn future theoretical progress into practical results.

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Appendix A: Extractor definitions

An extractor $\text{Ext} : \{0,1\}^n \times \{0,1\}^d \rightarrow \{0,1\}^m$ is a function which takes a weak source of randomness $X$ and a uniformly random, short seed $Y$, and produces some output $\text{Ext}(X,Y)$, which is almost uniform. The extractor is said to be strong, if the output is approximately independent of the seed.

The distance from uniform is measured by the trace distance, defined as $d(\rho, \sigma) := \frac{1}{2}\|\rho - \sigma\|_{\text{tr}}$, where $\| \cdot \|_{\text{tr}}$ denotes the trace norm given by $\| A \|_{\text{tr}} := \text{tr} \sqrt{A^\dagger A}$.

**Definition A.1** (strong extractor [19].) A function $\text{Ext} : \{0,1\}^n \times \{0,1\}^d \rightarrow \{0,1\}^m$ is a $(k,\varepsilon)$-strong extractor, if for all distributions $X$ with min-entropy $H_{\text{min}}(X) \geq k$ and a uniform seed $Y$, we have\(^{18}\)

$$\frac{1}{2}\|\rho_{\text{Ext}(X,Y)} - \tau_U \otimes \rho_Y\|_{\text{tr}} \leq \varepsilon,$$

where $\tau_U$ is the fully mixed state on a system of dimension $2^m$.

When (quantum) side information $E$ about the source $X$ is present, the randomness of the source is measured relative to this side information. We also require the output of the extractor to be close to uniform and independent from $E$.

**Definition A.2** (quantum-proof strong extractor [18], Section 2.6). A function $\text{Ext} : \{0,1\}^n \times \{0,1\}^d \rightarrow \{0,1\}^m$ is a quantum-proof (or simply quantum) $(k,\varepsilon)$-strong extractor, if for all states $\rho_{XE}$ classical on $X$ with

\(^{18}\) A more standard classical notation would be $\frac{1}{2}\|\text{Ext}(X,Y) \circ Y - U \circ Y\| \leq \varepsilon$, where the distance metric is the variational distance. However, since classical random variables can be represented by quantum states diagonal in the computational basis, and the trace distance reduces to the variational distance, we use the quantum notation for compatibility with the rest of this work.
Scaling behaviour: Block(GF(p))+RSH on 48 core Opteron (n=216, 40 repetitions)

FIG. 15. Throughput scaling of the block(GF(p))/RSH primitive combination for an increasing amount of CPUs. The speedup is well below one even for a moderate number of involved cores, and sees a further slow-down in the many-core case. Nonetheless, data rates that are reasonable for practical application are obtained, making the primitive combination a viable choice to post-process the output of slow devices for which it is important to not sacrifice valuable entropy, as is the case for the faster XOR extractor. While the per-core measurement in Figure 14 is more interesting from a scalability point of view, this figure provides guidance to what resources are necessary to satisfy given experimental constraints.

\[ H_{\text{min}}(X|E)_\rho \geq k, \] and for a uniform seed \( Y \), we have
\[
\frac{1}{2} ||p_{\text{Ext}(X,Y)} \otimes \rho_Y \otimes \rho_E||_\text{tr} \leq \epsilon,
\]
where \( \tau_U \) is the fully mixed state on a system of dimension \( 2^n \).

The function \( \text{Ext} \) is a classical-proof \((k, \epsilon)\)-strong extractor with uniform seed if the same holds with the system \( E \) restricted to classical states.

Note that any conventional extractor (Definition A.1) is classical-proof with slightly weaker parameters.

**Lemma A.3** (\cite{55} Section 2.5, \cite{59} Proposition 1). Any \((k, \epsilon)\)-strong extractor is a classical-proof \((k+\log \frac{1}{\epsilon}, 2\epsilon)\)-strong extractor.

In the extractor constructions described in Section II, we are particularly interested in extractors which only need to process a few bits of the input for every bit of output. These extractors are called local, and defined as follows.

**Definition A.4** (\(\ell\)-local extractor \cite{21}). An extractor \( \text{Ext} : \{0,1\}^n \times \{0,1\}^d \rightarrow \{0,1\}^m \) is \(\ell\)-locally computable (or \(\ell\)-local), if for every \( y \in \{0,1\}^d \), the function \( x \mapsto \text{Ext}(x,y) \) depends on only \( \ell \) bits of its input, where the bit locations are determined by \( y \).

This notion of local extractors applies equally to extractors with and without (quantum) side information.

**Appendix B: Known extractor results**

The next sections contain many known theorems on extractors, which we need to derive the parameters of the constructions from Section II.

1. List-decodable codes

A standard error correcting code guarantees that if the error is small, any string can be uniquely decoded. A list-decodable code guarantees that for a larger (but bounded) error, any string can be decoded to a list of possible messages.

**Definition B.1** (list-decodable code \cite{40}). A code \( C : \{0,1\}^n \rightarrow \{0,1\}^m \) is said to be \((\epsilon, L)\)-list-decodable if every Hamming ball of relative radius \( 1/2 - \epsilon \) in \( \{0,1\}^n \) contains at most \( L \) codewords.

List-decodable error correcting codes are known to be 1-bit extractors \cite{20, 21}. This has been rewritten out explicitly in \cite{4}.

**Lemma B.2** (\cite{4} Theorem D.3). Let \( C : \{0,1\}^n \rightarrow \{0,1\}^m \) be an \((\epsilon, L)\)-list-decodable code. Then the function
\[
C' : \{0,1\}^n \times [n] \rightarrow \{0,1\}
\]
\[
(x,y) \mapsto C(x)_y,
\]
is a \((\log L + \log \frac{1}{\epsilon}, 2\epsilon)\)-strong extractor.

As noted in a footnote of \cite{4}, this lemma can be strengthened to classical-proof extractors.

**Lemma B.3.** Let \( C : \{0,1\}^n \rightarrow \{0,1\}^m \) be an \((\epsilon, L)\)-list-decodable code. Then the function
\[
C' : \{0,1\}^n \times [n] \rightarrow \{0,1\}
\]
\[
(x,y) \mapsto C(x)_y,
\]
is a classical-proof \((\log L + \log \frac{1}{\epsilon}, 2\epsilon)\)-strong extractor.

\[19\] In the arXiv version, this theorem is numbered C.3
2. One-bit extractors

König and Terhal [39] show that any one-bit extractor is quantum-proof.

Theorem B.4 ([39 Theorem III.1]). Let $C : \{0,1\}^n \times \{0,1\}^t \to \{0,1\}$ be a $(k,\varepsilon)$-strong extractor. Then $C$ is a quantum-proof $(k + \log 1/\varepsilon, 3\sqrt{\varepsilon})$-strong extractor.

If we however have a construction which has already been shown to be a classical-proof $(k,\varepsilon)$-strong extractor, then Theorem B.4 can be refined as follows.

Lemma B.5 (Implicit in [39]). Let $C : \{0,1\}^n \times \{0,1\}^t \to \{0,1\}$ be a classical-proof $(k,\varepsilon)$-strong extractor. Then $C$ is a quantum-proof $(k, (1 + \sqrt{2}/\varepsilon)\sqrt{\varepsilon})$-strong extractor.

3. Universal hashing

A family of hash functions is almost universal, if the probability of a collision is low.

Definition B.6 ([25]). A family of hash functions $\{h : X \to Z\}$ is said to be $\delta$-almost universal$_2$ ($\delta$-AU$_2$), if for any $x, x' \in X$ with $x \neq x'$,
\[
\Pr [h(x) = h(x')] \leq \delta,
\]
where the hash functions are chosen uniformly at random.
The family is said to be universal$_2$, if it is $\delta$-AU$_2$ with $\delta = \frac{1}{|Z|}$.

Tomamichel et al. [6] show that for such a family of hash functions $\{h_y\}_y$, the corresponding extractor — defined as $\text{Ext}(x,y) := h_y(x)$ — is quantum-proof if $\delta$ is small enough.

Theorem B.7 ([6 Theorem 7]). If a family of hash functions $\{h : \{0,1\}^n \to \{0,1\}^m\}$ is $\delta$-AU$_2$, for $\delta = \frac{1+\varepsilon^2}{2^m}$, then chosen uniformly at random, they build a quantum-proof $(m + 4\log \frac{1}{\varepsilon} + 1, 2\varepsilon)$-strong extractor.

4. Trevisan’s extractor

In [4 Theorem 4.6], De et al. show that if a $(k,\varepsilon)$-strong one-bit extractor is used in Trevisan’s construction, the final extractor is a quantum-proof $(k + rm + \log 1/\varepsilon, 3m\sqrt{\varepsilon})$-strong extractor, where $m$ is the output length and $r$ is a parameter of the weak design.

That theorem is the combination of the following implicit lemma and Lemma A.3.

Lemma B.8 (Implicit in [4]). Let $C : \{0,1\}^n \times \{0,1\}^t \to \{0,1\}$ be a quantum-proof $(k,\varepsilon)$-strong extractor with uniform seed and $S_1, \ldots, S_m \subset [d]$ a weak $(m,t,r,d)$-design. Then Trevisan’s extractor, $\text{Ext}_C : \{0,1\}^n \times \{0,1\}^d \to \{0,1\}^m$, is a quantum-proof $(k + rm, mn\varepsilon)$-strong extractor.

If we use a one-bit extractor which is known to be quantum proof, we get better parameters from Lemma B.8 than [4 Theorem 4.6].

Appendix C: Weak design proofs

1. Basic construction

Lemma C.1. The weak design construction described in Section III B 3 has $r < 2e$.

Proof. Ma and Tan [18] prove that if $m \in [e^c, e^{c+1}]$ and $e^c$ divides $m$, then the weak design has $r < e$. The lemma is thus immediate for $m = k t^c$ and any integer $1 \leq k \leq t$.

Let $k t^c < m < (k+1)t^c$ for some integer $1 \leq k < t$.
Since the construction for $m$ is the same as the construction for $m' = (k+1)t^c$ with the last sets $S_p$ dropped, the overlap can only decrease. Thus
\[
\sum_{q<p} 2^{|S_q \cap S_p|} < (k+1)e^{c} t^c < \frac{k+1}{k} e m \leq 2em.
\]

2. Reducing the overlap

Lemma C.2. The weak design construction described in Section III B 3 has $r = 1$.

Proof. For simplicity, we number the sets of the weak design $W$ with two indices $(i,j)$, where $0 \leq i \leq \ell$ and $1 \leq j \leq m_i$, and label the corresponding set of the basic weak design $S_i$. We need to show that the second condition of Definition III.1 holds for $r = 1$, namely that for all $(i,j)$,
\[
\sum_{(g,h) < (i,j)} 2^{|S_g \cap S_h|} \leq m_i
\]
where $\{(g,h) : (g,h) < (i,j)\} := \bigcup_{g<i} \{(g,h) : h < m_g\} \cup \{(i,h) : h < j\}$.

Note that (C1) implies that for all $0 \leq k \leq \ell - 1$,
\[
\sum_{j \leq k} n_j \leq \sum_{j \leq k} m_j < \sum_{j \leq k} n_{j+1},
\]
from which we get
\[
m_k < \sum_{j \leq k} n_{j} - \sum_{j \leq k-1} m_{j} + 1 \leq n_k + 1.
\]
Furthermore, from the sum of a geometric series, we have
\[
\sum_{j \leq k-1} n_j + r' n_{k} = \frac{1 - (1 - \frac{1}{r'})^k}{1 - (1 - \frac{1}{r'})} n_0 + r' \left(1 - \frac{1}{r'}\right)^k n_0
\]
\[
= r' n_0 = m - r'.
\]
For any two sets $S_{i,j}$ and $S_{g,h}$ with $i \neq g$, we have $|S_{g,h} \cap S_{i,j}| = 0$. Thus for any set $S_{i,j}$ with $i \leq \ell - 1$, we have

$$
\sum_{(g,h) < (i,j)} g|S_{g,h} \cap S_{i,j}| = \sum_{g<i,h \leq m_g} 1 + \sum_{h<j} 2|S_{h} \cap S_{j}| \\
\leq \sum_{g<i} m_g + r'm_i \\
< \sum_{g<i} n_g + 1 + r'(n_i + 1) \\
= m + 1,
$$

where we used (C1) and (C2) in the second from the last line, and (C3) in the last line. Since the LHS of the above inequality is an integer, and the inequality is strict, we must have

$$
\sum_{(g,h) < (i,j)} g|S_{g,h} \cap S_{i,j}| \leq m.
$$

Finally, for the case of $S_{\ell,j}$, note that $\ell$ was chosen such that $m_\ell \leq t$. This can be seen as follows.

$$
m_\ell = m - \sum_{j \leq \ell - 1} m_j \leq m - \sum_{j \leq \ell - 1} n_j \\
= m - \frac{1 - (1 - \frac{1}{n})^\ell}{1 - (1 - \frac{1}{n})} (m - 1) \\
= r' + \left(1 - \frac{1}{r'}\right) (m - r').
$$

By plugging (3) in this, we get $m_\ell \leq t$. Since $t$ is the size of the finite field, the polynomial used to generate the elements of $S_{i,j}$ has all coefficients 0, except the constant term which is $j$. We thus have $S'_t = \{(x,j)\}_{x \in GF(t)}$, and so the sets $\{S_{\ell,j}\}_{j \in GF(t)}$ have no intersection. Hence

$$
\sum_{(g,h) < (i,j)} 2|S_{g,h} \cap S_{i,j}| \leq \sum_{g \leq \ell} m_\ell = m.
$$

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