On the Randomness Complexity of Property Testing^{*}

Oded Goldreich[†] Department of Computer Science Weizmann Institute of Science Rehovot, ISRAEL. oded.goldreich@weizmann.ac.il Or Sheffet[‡] Department of Computer Science Carnegie Mellon University Pittsburgh, PA 15213, USA. osheffet@andrew.cmu.edu

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Abstract

We initiate a general study of the randomness complexity of property testing, aimed at reducing the randomness complexity of testers without (significantly) increasing their query complexity. One concrete motivation for this study is provided by the observation that the product of the randomness and query complexity of a tester determine the actual query complexity of implementing a version of this tester that utilizes a weak source of randomness (through a randomness-extractor). We present rather generic upper- and lower-bounds on the randomness complexity of property testing and study in depth the special case of testing bipartiteness in two standard property testing models.

Keywords: Property Testing, Randomness Complexity, Weak Sources of Randomness, Randomness Extractors, Sampling.

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[‡]Work done while Or was a graduate student at the Weizmann Institute of Science.

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1 Introduction

Property testing (initiated by Rubinfeld and Sudan [24] and Goldreich, Goldwasser, and Ron [14]) is concerned with a relaxed type of decision problems. Specifically, for a fixed property (resp., a set) Π , the task is to distinguish between objects that have property Π (resp., are in Π) and objects that are "far" from having property Π (resp., are "far" from any object in Π). The focus of property testing is on sublinear-time algorithms, which in particular cannot examine the entire object. Instead, these algorithms, called *testers*, may obtain bits in the representation of the object by issuing adequate queries. Indeed, in this case, the query complexity of testers becomes a measure of central interest.

For natural properties, testers of sublinear query-complexity must be randomized (see details in Section 3.1). This is a qualitative assertion, and the corresponding quantitative question arises naturally: for any fixed property Π and a sublinear function q, what is the randomness-complexity of testers for Π that have query-complexity q?

In addition to the natural appeal of the foregoing question, there are concrete reasons to care about it. Firstly, the randomness-complexity of a tester determines the length of PCPs that are constructed on top of this tester. Indeed, this was the motivation for the interest of Goldreich and Sudan [17] and Ben-Sasson, Sudan, Vadhan, and Wigderson [7] in reducing the randomness complexity of low-degree testing. Secondly, the randomness-complexity of a tester affects the time and query complexities of implementing a version of this tester while utilizing a weak source of randomness. This motivation is further discussed in Section 1.2.

Indeed, the randomness-complexity of testers was considered in some prior work, starting in [17]. This subject is the pivot of [7] and is the main topic studied by Shpilka and Wigderson [28]. However, these works refer to specific (algebraic) tasks (i.e., testing low-degree polynomials and group homomorphisms). In contrast, our focus in this paper is either on general properties (see Section 1.4) or on specific combinatorial properties (see Section 1.3).

1.1 The Perspective of Average-Estimation

Property testing is a vast generalization of the task of estimating the average value of a function. Specifically, consider the task of distinguishing between functions $f : \{0,1\}^n \to \{0,1\}$ having average value exceeding 0.5 and functions that are ϵ -far from having this property (i.e., functions having average value below $0.5 - \epsilon$). Clearly, this task can be solved by a randomized algorithm that queries the function at $O(1/\epsilon^2)$ (random) points. This query-complexity is optimal and any algorithm achieving it, called a *sampler*, must be randomized (see Canetti, Even, and Goldreich [9]). Furthermore, a quantitative study of the randomness-complexity of samplers in terms of their querycomplexity was also carried out in [9]. The current paper may be viewed as extending this study to the domain of general property testing.

Note that estimating the average value of a function corresponds to very restricted properties of functions. In particular, these properties are *symmetric* (i.e., are invariant under any relabeling of the inputs to the function). In contrast, most of the study of property testing refers to properties that are not symmetric (e.g., being a low-degree polynomial, monotonicity, representing a graph that has a certain graph property, etc). Furthermore, while all symmetric properties of Boolean functions are easily testable by straightforward sampling, this cannot be said about property testing in general (nor about the numerous special cases that were studied in the last decade (see surveys of Fischer [10] and Ron [23])).

1.2 A Concrete Motivation: Using Weak Sources of Randomness

In the context of traditional randomized algorithms, a concrete motivation for minimizing the randomness-complexity is provided by the exponential effect of the randomness-complexity on the time-complexity of a possible derandomization. In contrast, in the context of property testing, derandomization is typically infeasible, because (as noted above) deterministic testers cannot have sublinear query complexity. Instead, a different motivation (advocated by the first author [12]), becomes very relevant in this context.

We refer to the effect of the randomness-complexity on the overhead involved in implementing the tester when using only weak sources of randomness (rather than perfect ones). Specifically, we refer to the paradigm of implementing randomized algorithms by using (a single sample from) such a weak source, and trying all possible seeds to an adequate randomness extractor (see below). We shall see that the overhead created by this method is determined by the randomness-complexity of the original algorithm.

Loosely speaking, a randomness extractor is a function $E : \{0,1\}^n \times \{0,1\}^s \to \{0,1\}^r$ that uses an s-bit long random seed in order to transform an n-bit long (outcome of a) weak source of randomness into an r-bit long string that is almost uniformly distributed in $\{0,1\}^r$. Specifically, we consider arbitrary weak sources that are restricted (only) in the sense that, for a parameter k, no string appears as the source outcome with probability that exceeds 2^{-k} . Such sources are called (n, k)-sources (and k is called the min-entropy). Now, E is called a (k, ϵ) -extractor if for any (n, k)source X it holds that $E(X, U_s)$ is ϵ -close to U_r , where U_m denotes the uniform distribution over m-bit strings (and the term 'close' refers to the statistical distance between the two distributions). For further details about (k, ϵ) -extractors, the interested reader is referred to Shaltiel's survey [25].

Next, we recall the standard paradigm of implementing randomized algorithms while using sources of weak randomness. Suppose that the algorithm A has time-complexity t and randomnesscomplexity r < t. Recall that, typically, the analysis of algorithm A refers to what happens when A obtains its randomness from a perfect random source (i.e., for each possible input α , we consider the behavior of $A(\alpha, U_r)$, where $A(\alpha, \omega)$ denotes the output of A on input α when given randomness ω). Now, suppose that we have at our disposal only a weak source of randomness; specifically, a (n,k)-source for $n \gg k \gg r$ (e.g., n = 10k and k = 2r). Then, using a (k, ϵ) extractor $E: \{0,1\}^n \times \{0,1\}^s \to \{0,1\}^r$, we can transform the *n*-bit long outcome of the weak source into 2^s strings, each of length r, and use the resulting 2^s strings (which are "random on the average") in 2^s corresponding invocations of the algorithm A. That is, upon obtaining the outcome $x \in \{0,1\}^n$ from the source, we invoke the algorithm A for 2^s times such that in the ith invocation we provide A with randomness E(x, i). The results of these 2^s invocations are processed in the natural manner. For example, if A is a decision procedure, then we output the majority vote obtained in the 2^s invocations (i.e., when given the input α , we output the majority vote of the sequence $\langle A(\alpha, E(x, i)) \rangle_{i=1,\dots,2^s}$). An analysis of the error probability of this procedure is provided in Section 2.4.

Let us consider the cost of the foregoing implementation. We assume, for simplicity, that the running-time of the randomness extractor is significantly smaller than the running-time of A. Then, algorithm A can be implemented using a weak source, while incurring an overhead factor of 2^s . Thus, we focus on providing lower and upper bounds on the aforementioned overhead (i.e., 2^s) as a function of r (the number of random bits used by the original tester). Recalling that $s > \log_2(n-k)$ and n > k > r - s must hold (cf. [25]), it follows that for $k = n - \Omega(n)$ the overhead factor (i.e., 2^s) is *lower bounded* by $\Omega(n)$, which is $\Omega(r)$. On the other hand, for $k = n^{\Omega(1)}$, efficient randomness-extractors using $s = (1+o(1)) \log_2 n$ (and providing $r = k^{1-o(1)}$) are known (see [25, 26]). It follows

that (the overhead factor of) 2^s is upper bounded by $n^{1+o(1)}$ (i.e., is almost linear in n), even when utilizing the randomness in the source in an almost optimal manner (i.e., extracting $r = k^{1-o(1)}$ almost random bits from any (n, k)-source). We comment that in the most natural case of weak sources, that is, sources of constant min-entropy rate (i.e., $k = \Omega(n)$), the extraction rate can be improved to linear (i.e., $r = \Omega(k)$); see [25, 29]. Thus, for $k = \Omega(n)$ (resp., $k = n^{\Omega(1)}$), the overhead factor (i.e., 2^s) is upper bounded by a function that is almost linear in r (resp., polynomial in r).

To summarize, we have established our claim that the time-complexity of implementing randomized algorithms when using weak sources of randomness is related to the randomness-complexity of these algorithms. The same applies to the query complexity of testers. Specifically, for (n, k)sources satisfying $k = \Omega(n)$ (resp., satisfying $k = n^{\Omega(1)}$), the query-complexity of implementing a tester is almost linear in $r \cdot q$ (resp., is poly $(r) \cdot q$), where q is the query-complexity of the original tester that uses a perfect source of (r bits of) randomness.

1.3 Specific Algorithms

The motivation discussed in Section 1.2 is best illustrated by our results regarding testing bipartiteness in the bounded-degree model (as initiated by Goldreich and Ron [15]). Specifically, fixing a degree bound d, the task is to distinguish (N-vertex) bipartite graphs of maximum degree d from (N-vertex) graphs of maximum degree d that are ϵ -far from bipartite (for some parameter ϵ), where ϵ -far means that $\epsilon \cdot dN$ edges have to be omitted from the graph in order to yield a bipartite graph. We note that no deterministic algorithm of o(N) time-complexity can solve this promise problem (see Section 3.1.1). Yet, there exists a probabilistic algorithm of time-complexity $\tilde{O}(\sqrt{N}\text{poly}(1/\epsilon))$ that solves this problem correctly (with probability 2/3). This algorithm makes $q \stackrel{\text{def}}{=} \tilde{O}(\sqrt{N}\text{poly}(1/\epsilon))$ incidence-queries to the graph, and (as described in the work Goldreich and Ron [16]) has randomness-complexity $r > q > \sqrt{N}$ (yet $r < q \cdot \log_2 N$).¹

Let us now turn to the question of implementing the foregoing tester in a setting where we have access only to a weak source of randomness. In this case, the implementation calls for invoking the original tester $\tilde{O}(r)$ times, which yields a total running time of $\tilde{O}(r) \cdot \tilde{O}(q) > q^2 > N$ (and the same bound holds for its query-complexity). But in such a case we better use the standard (deterministic) decision procedure for bipartiteness!

Fortunately, a randomness-efficient implementation of the original tester of [16] is possible. This implementation (presented in Section 4.2) has randomness-complexity $r' = \text{poly}(\epsilon^{-1} \log N)$ (rather than $r = \text{poly}(\epsilon^{-1} \log N) \cdot \sqrt{N}$). Thus, the cost of the implementation that uses a weak source of randomness is $\tilde{O}(r' \cdot q) = \tilde{O}(\sqrt{N} \text{poly}(1/\epsilon))$, which matches the original bound (up to differences hidden in the $\tilde{O}()$ and poly() notation).

The randomness-efficient implementation of the [16]-tester presented in Section 4.2 is based on pin-pointing the "random features" used in the original analysis, and providing an alternative (randomness-efficient) implementation that satisfies the same features. In general, such features typically include various "hitting" and "sampling" conditions (see Section 2.2 and Goldreich's survey [11]). In such cases, using randomness-efficient hitters and samplers may yield a significant saving in the randomness-complexity of the underlying tester. While this approach suffices in many cases, in other cases a more significant modification of the original tester yields better results. This is indeed the case with respect to the randomness-efficient tester presented in Section 4.1.

In Section 4.1 we consider testers for graph properties in the adjacency matrix model (as initiated by Goldreich, Goldwasser, and Ron [14]). Specifically, we consider the task of testing bipartiteness.

¹We comment that $\Omega(\sqrt{N})$ is a lower-bound on the query-complexity of any property tester of bipartiteness (in the bounded-degree model; see [15]).

We recall that the tester presented in [14] works by selecting a random set of $\tilde{O}(\epsilon^{-2})$ vertices and inspecting the (corresponding) induced subgraph. In fact, as shown in [14], it suffices to make $\tilde{O}(\epsilon^{-3})$ queries. A randomness-efficient implementation of the "random features" used in the original analysis, allows reducing the randomness-complexity to $\tilde{O}(\epsilon^{-1}) + O(\log N)$, where N denotes the number of vertices. In contrast, using an alternative approach, we present a tester of randomness-complexity $O(\log(1/\epsilon)) \cdot \log N$, while maintaining a query-complexity bound of $\tilde{O}(\epsilon^{-3})$. The latter randomness-efficient tester is the main technical contribution of this work. In the next paragraph, we provide an extremely high-level description of the principles underlying its design.

The original tester works by first selecting a random sample of $t = \tilde{O}(\epsilon^{-1})$ vertices, and the analysis refers to 2^t candidate 2-colorings that are induced by all possible 2-partitions of this sample. The tester then selects an auxiliary sample of $\tilde{O}(t/\epsilon)$ vertex-pairs and checks whether this sample rules out all these 2^t candidate 2-colorings. The analysis boils down to showing that if the graph is ϵ -far from bipartite then, with high probability, all these candidate 2-colorings are ruled out. This is shown by applying a union bound on this set of 2^t candidate 2-colorings, which means that each candidate has to be ruled out with probability at least $1 - 2^{-t}$. Thus, the randomness complexity of any implementation of this tester must exceed t. Seeking to achieve randomness-complexity that is linearly related to log t, we perform a preliminary step aimed at obtaining a single 2-partition of the initial t-vertex sample that induces a single candidate 2-coloring, which will be checked as in the original tester. The preliminary step obtains such a 2-partition by collecting constraints on the mutual placements of pair of vertices. These constraints are found using the same mechanism that underlies the checking of candidates in the original tester. The punch-line is that here we are dealing with $\binom{t}{2}$ (rather than 2^t) events, which allows us to work with an error probability of $t^{-2}/O(1)$ (rather than $2^{-t}/O(1)$) per each event.

Thus, Sections 4.1 and 4.2 represent two approaches to reducing the randomness-complexity of testers: Section 4.2 represents the approach of merely providing randomness-efficient implementation of some random features used in the analysis of the original tester. In contrast, Section 4.1 represents the approach of redesigning the tester (while, indeed, benefiting from ideas that underly the design of the original tester).

1.4 Generic Bounds

In contrast to the specific algorithms described in Section 1.3, we now consider quite generic lowerand upper-bounds on the randomness-complexity of property testers as a function of their querycomplexity. These bounds (as well as the rest of our study) refer to testers with constant error probability. We stress that these results do not refer to the time-complexity of the testers, which makes the lower-bounds stronger (and the upper-bound weaker).

Lower bounds. We show that, for a wide class of properties of functions defined over a domain of size D, the randomness-complexity of testing with q queries is at least $\log_2(D/q) - O(1)$. The aforementioned class includes all known testers (see details below). Needless to say, the dependence on the query-complexity is essential, because deterministic testers of query-complexity D exist for any property. Furthermore, the randomness-complexity of any tester can be decreased by an additive term of t while increasing the query complexity by a factor of 2^t . The lower-bound asserts that for natural property testers (where $q \ll D$), the randomness-complexity should "compensate" for not scanning the entire domain; that is, $2^r \cdot q = \Omega(D)$, where r denotes the randomnesscomplexity of the tester (and q its query-complexity).

The lower-bounds established in Section 3.1 apply to two general and natural classes of prop-

erties. In particular, these lower-bounds apply to testing low-degree polynomials (cf., e.g., Blum, Luby, and Rubinfeld [8] and Rubinfeld and Sudan [24]), locally-testable codes (cf., e.g., Goldreich and Sudan [17]), testing graph properties (in both the adjacency matrix and incidence-list models, see Goldreich, Goldwasser, and Ron [14] and Goldreich and Ron [15], resp.), testing monotonicity (cf., e.g., Goldreich, Goldwasser, Lehman, Ron, and Samorodnitsky [13]), and testing of clustering (cf., e.g., Alon, Dar, Parnas, and Ron [2]).

Upper bounds. The upper-bound established in Section 3.2 refers to any property and assert that the randomness-complexity of any tester may be reduced to $\log_2 D + \log_2 \log_2 R + O(1)$, where R is the size of the range of the functions we refer to (and D is the size of their domain).

Note that the gap between the lower and upper bounds is $\log_2 q + \log_2 \log_2 R + O(1)$. We note that in the special case of evaluating the average of Boolean functions by query-optimal samplers, the gap can be reduced to a constant by using the improved lower-bound of Radhakrishnan and Ta-Shma [22] (which implies that the randomness-complexity of any sampler is at least $\log_2(D/q) + 2\log_2(1/\epsilon) - O(1)$, while query-optimal samplers have $q = \Theta(\epsilon^{-2})$ (see [9])). See further discussion in Section 3.1.3.

1.5 Organization

In Section 2 we review some basic tools (e.g., randomness-efficient hitters) that are used in this work. Our generic results are presented in Section 3, where Section 3.1 provides lower bounds and Section 3.2 provides upper bounds. The specific testers for the case of bipartiteness are presented in Section 4, where the Section 4.1 refers to the adjacency matrix model and Section 4.2 refers to the bounded-degree model.

2 Preliminaries

In this section we review some basic tools that are used in this work. Specifically, Section 2.2 reviews the basic definitions and results reagrding randomness-efficient hitters, which are used extensively in Section 4. In addition, 4-wise independent sequences are reviewed in Section 2.1 (and used in Section 4.2), whereas randomness-efficient error-reduction is reviewed in Section 2.3 (and used in Section 3.2). We believe that some readers can afford skipping the current section.

Notation: The notation \tilde{O} represents an upper bound that is almost linear in the argument; that is, $\tilde{O}(x)$ means an upper bound of the form $O(\operatorname{poly}(\log x) \cdot x)$. Similarly, $\tilde{\Theta}$ represents a \tilde{O} upper bound that is tight up to a polylogarithmic factor; that is, $\tilde{\Theta}(x)$ means an upper bound of $\tilde{O}(x)$ that is matched by a lower bound of the form $\Omega(x/\operatorname{poly}(\log x))$.

We often use the phrase "with high probability" without specifying the error bound, which is typically a sufficiently small constant. In all cases, the meaning of this phrase should be clear from the context.

2.1 Pairwise and 4-wise independent sequences

Let S be a finite set and $t \leq \ell$ be integers. A distribution over S^{ℓ} is called *t*-wise independent if its restriction to any t coordinates is uniformly distributed over S^{t} . That is, a sequence of (possibly dependent) random variables $(X_1, ..., X_{\ell})$, each distributed over S, is called *t*-wise independent if

for every $i_1 < i_2 < \cdots < i_t$ (in $[\ell]$) and for every $(v_1, ..., v_t) \in S^t$ it holds that $\Pr[(\forall j \in [k]) X_{i_j} = v_j] = |S|^{-t}$. In case t = 2, we call the sequence pairwise independent.

In the following construction (due to [1]), we assume that |S| is a power of 2, and identify S with the corresponding finite field. Let $\alpha_1, ..., \alpha_\ell$ be (fixed and) distinct elements of this field, and consider the distribution generated by uniformly and independently selecting $s_0, s_1, ..., s_{t-1} \in S$, and outputting the sequence $(r_1, ..., r_\ell)$, where $r_i = \sum_{j=0}^{t-1} \alpha_i^j s_j$. Then, this sequence is t-wise independent. Note that this sequence is generated using $t \log_2 |S|$ random bits.

2.2 Randomness-Efficient Hitters

The hitting problem is a one-sided version of the (Boolean) sampling problem (see, e.g., [11]). Given parameters n (length), ϵ (density) and δ (error), and oracle access to any function $f : \{0, 1\}^n \to \{0, 1\}$ such that $|\{x : f(x) = 1\}| \ge \epsilon 2^n$, the task is to find a string that is mapped to 1.

Definition 2.1 (hitter): A hitter is a randomized algorithm that on input parameters n, ϵ and δ , and oracle access to any function $f: \{0, 1\}^n \to \{0, 1\}$ such that $|f^{-1}(1)| \ge \epsilon 2^n$, satisfies

$$\Pr[\operatorname{hitter}^{f}(n,\epsilon,\delta) \in f^{-1}(1)] > 1-\delta$$

When ϵ and δ are fixed, we say that the resulting algorithm is a hitter for sets of density ϵ with error probability δ .

We shall also say that such a hitter hits any set of density ϵ with probability (at least) $1 - \delta$.

We briefly recall a few known results (and refer the interested reader to [11] for details). For any constant $\delta > 0$, using a pairwise-independent sequence of length $O(1/\epsilon)$, we obtain a hitter for sets of density ϵ with error probability δ . Thus, this hitter has query-complexity $O(1/\epsilon)$ and randomness-complexity 2n. An alternative hitter based on the neighborhood of a random vertex in an expander graph has query-complexity $O(1/\epsilon)$ and randomness-complexity n. Combining any of these hitters with a random walk (of length $O(\log(1/\delta))$) on an expander graph, we obtain a hitter for sets of density ϵ and any desired error probability δ such that this hitter has query-complexity $O(\epsilon^{-1}\log(1/\delta))$ and randomness-complexity $r + O(\log(1/\delta))$, where $r \in \{n, 2n\}$ depending on the basic hitter we use.

Note that each of the foregoing hitters generates a sequence of candidate strings in $\{0, 1\}^n$, and uses queries to f merely for the selection of one of these strings. In the subsequent text, we actually refer only to the sample-generating part of these hitters.

2.3 Randomness-Efficient Error-Reduction

Error-reductions are closely related to oblivious samplers (see, e.g., [25] or [11]). Intuitively, given a probabilistic decision procedure of (two-sided) error probability $\epsilon < 1/2$, we wish to obtain a probabilistic decision procedure of (two-sided) error probability $\delta < \epsilon$. Representing the foregoing procedure (coupled with a generic input) by a Boolean function f (which maps the procedure's randomness to its decision), we obtain the following definition.

Definition 2.2 (error reduction): An error reduction is a randomized oracle machine, denoted M, that on input parameters n and $\delta < \epsilon < 1/2$, and oracle access to any function $f: \{0, 1\}^n \to \{0, 1\}$, satisfies the following condition. If for some $\sigma \in \{0, 1\}$ it holds that $|f^{-1}(\sigma)| \ge (1 - \epsilon) \cdot 2^n$, then

$$\Pr[M^f(n,\epsilon,\delta)=\sigma] > 1-\delta$$

When ϵ and δ are fixed, we say that the resulting machine reduces error ϵ to error δ .

In Section 3.2 we shall use a randomness-efficient error-reduction that reduces error 2/5 to error 1/3 by making a constant number of oracle calls and using *n* random bits. This error-reduction works by selecting a random vertex in a bounded-degree expander graph (of size 2^n) and querying *f* on all the neighbours of this vertex. For details, the interested reader is referred to to [11].

2.4 Analysis of the standard use of extractors

In continuation to Section 1.2, we prove the following claim.

Claim 2.3 Let A be a randomized decision procedure of randomness-complexity r and error probability p, and $E: \{0,1\}^n \times \{0,1\}^s \to \{0,1\}^r$ be an (k,ϵ) -extractor. Consider the algorithm A' that, on input α , obtains a single sample x from an (n,k)-source and rules according to the majority value in $\langle A(\alpha, E(x,i)) \rangle_{i=1,\dots,2^s}$. Then, A' has error probability at most $2(p+\epsilon)$. Furthermore, if E is actually a $(k-t,\epsilon)$ -extractor and $p+\epsilon < 1/2$ then A' has error probability 2^{-t} .

Proof: The analysis of the foregoing implementation is based on the fact that "on the average" the 2^s strings extracted from the source approximate a perfect *r*-bit long source (i.e., a random setting of the *s*-bit seed yields an almost uniformly distributed *r*-bit string). Specifically, by definition, if X is a (n, k)-source then $E(X, U_s)$ is ϵ -close to U_r . It follows that the probability that $A(\alpha, E(X, U_s))$ errs is at most $p + \epsilon$. By Markov Inequality, the probability that the majority of the values in $\langle A(\alpha, E(X, i)) \rangle_{i=1,\dots,2^s}$ are wrong is at most $2(p + \epsilon)$. The main part of the claim follows.

Towards the furthermore clause, fixing any α , we call a string $x \in \{0, 1\}^n$ bad if the probability that $A(\alpha, E(x, U_s))$ is wrong is at least 1/2. Using the hypothesis that E is $(k - t, \epsilon)$ -extractor it follows that there are at most 2^{k-t} bad strings (otherwise, defining X' to be uniformly distributed on the set of bad strings, we reach a contradiction to the hypothesis (because $E(X', U_s)$ is not ϵ -close to U_r)). Hence, the outcome of a (n, k)-source is bad with probability at most 2^{-t} and the claim follows.

Comment. We note that randomized procedures with one-sided error probability p can be implemented using a weak random source as long as $p + \epsilon < 1$. An important case is of search problems for which the randomized algorithm finds a correct solution with probability 1 - p and halts without solution otherwise. When implementing such an algorithm, we may output any solution obtained in any of the invocations of the original algorithm, which means that we "rule by or" rather than "ruling by majority".

3 Generic Bounds

We consider testing properties of functions from D to R; that is, all functions considered here have domain D and range R. Fixing a set of such functions, denoted Π , and a proximity parameter, denoted $\epsilon > 0$, we focus on the task of ϵ -testing Π as arises from the following definition.

Definition 3.1 (testers): A randomized oracle machine T is called an ϵ -tester for Π if the following two conditions hold:

- 1. For every $f \in \Pi$ it holds that $\Pr[T^f = 1] \geq 2/3$.
- 2. For every f that is ϵ -far from Π it holds that $\Pr[T^f = 1] \leq 1/3$, where f is ϵ -far from Π if for every $g \in \Pi$ it holds that $\Pr_{x \in D}[f(x) \neq g(x)] > \epsilon$.

In case the first condition holds with probability 1, we say that T has one-sided error. A tester is called non-adaptive if it determines its queries based solely on its internal coin-tosses (and independently of the answers to prior queries).

The query and randomness complexities of T are defined in the natural manner.

Note that we have defined property testers for finite properties and for a fixed value of the proximity parameter ϵ . The definition may be extended to infinite properties and varying ϵ , by providing the tester with |D|, |R| and ϵ as inputs (and assuming D = [|D|]). Occasionally, we shall assume that $\epsilon \geq |D|^{-1}$; otherwise, ϵ -testers coincide with standard decision procedures.

3.1 Lower Bounds

We provide lower-bounds on the randomness complexity of testing two general classes of properties.

3.1.1 Strongly evasive properties

The first class that we consider consists of properties that are "strongly evasive" in the sense that the values (of some function) at any set that contains a constant fraction of the domain are consistent both with some function that has the property and with some other function that is far from having the property.

Definition 3.2 (strongly evasive): For fixed parameters ϵ and ρ , the property Π is called strongly evasive (with respect to parameters ϵ and ρ) if there exists a function $f: D \to R$ such that for every $D' \subset D$ of density ρ (i.e., $|D|' = \rho \cdot |D|$), there exists $f_1 \in \Pi$ and $f_0: D \to R$ that is ϵ -far from Π such that for every $x \in D'$ it holds that $f_1(x) = f_0(x) = f(x)$.

Many natural properties are strongly evalve (with respect to various pairs of parameters); see examples below. We mention that Definition 3.2 is incomparable to the standard definition of evalveness (cf., e.g., [20]): On one hand, strong evalveness has a non-deterministic flavor (i.e., it refers to all choices of D' after f is fixed) and furthermore it refers to the relaxation of property testing (i.e., f_0 is far from Π rather than only not in Π). On the other hand, we shall focus on constant values of $\rho < 1$, whereas standard evalveness refers to $\rho = 1 - |D|^{-1}$.

We show that testing any strongly evasive property requires randomness complexity that is logarithmic in the ratio of the domain size over the query complexity. This result can be easily proved by extending a similar result regarding samplers (proved by Canetti, Even, and Goldreich [9]).

Theorem 3.3 Let Π be strongly evasive with respect to ϵ and ρ . Then any ϵ -tester for Π that has query complexity q, must have randomness complexity greater than $\log_2(\rho|D|/q)$.

Proof: Let T be an arbitrary ϵ -tester of query-complexity q and randomness-complexity r, and f be a function witnessing the fact that Π is strongly evasive (i.e., for every set of density ρ of the domain, there exists $f_1 \in \Pi$ and $f_0 : D \to R$ that is ϵ -far from Π such that f, f_1 and f_0 agree on all elements in this set). For every $\omega \in \{0, 1\}^r$, we consider the set of queries made by T when the outcome of T's coin-tosses equals ω and T is given oracle access to f. Denoting the latter set by Q_{ω} , we let $D' = \bigcup_{\omega \in \{0,1\}^r} Q_{\omega}$. (Indeed, like the Q_{ω} 's, the set D' depends on f.) Clearly, $|D'| \leq 2^r \cdot q$. The theorem follows by proving that $|D'| > \rho \cdot |D|$.

Suppose towards the contradiction that $|D'| \leq \rho \cdot |D|$. Then, by our choice of the function f, there exists $f_1 \in \Pi$ and $f_0 : D \to R$ that is ϵ -far from Π such that for every $x \in D'$ it holds that $f_1(x) = f_0(x) = f(x)$. It follows that T^{f_1} and T^{f_0} behaves exactly as T^f (because all these functions agree on D'), which yields a contradiction (because T must accept f_1 with probability at least 2/3 and accept f_0 with probability at most 1/3).

Some applications. Many graph properties are strongly evasive, but since such properties will be at the focus of Section 3.1.2, we mention first a few examples that refer to different types of properties.

1. Multi-variate polynomial. For every m and d, we consider the set of m-variate polynomial of total degree d over a finite field F. To see that this set of functions over F^m is strongly evasive consider the all-zero function, f, and let $f_1 = f$. Then, for every D' of density 1/2, let $f_0(x) = 0$ if $x \in D'$ and $f_0(x) = 1$ otherwise. Assuming |F| > 4d (and using the Schwartz-Zippel Lemma), it follows that f_0 is 1/4-far from any degree d polynomial (because any non-zero polynomial of degree d may evaluate to zero on at most a d/|F| fraction of its domain). Invoking Theorem 3.3, we conclude that 1/4-testing the set of m-variate polynomials of total degree d over F, while using q queries, requires randomness at least $\log_2(|F^m|/2q) = m \log_2 |F| - \log_2 q - 1$.

We mention that, for every constant $\epsilon > 0$, the low-degree ϵ -tester of Ben-Sasson, Sudan, Vadhan, and Wigderson [7] uses $q = O(d \log |F|)$ queries and $m \log_2 |F| + \log_2(m \log |F|) + O(1)$ random bits.

Codes of linear distance. A binary code C ⊂ {0,1}ⁿ of distance d = Ω(n), is viewed as a set of functions of the form f : [n] → {0,1}, where each function corresponds to a codeword. To see that this set is strongly evasive consider any codeword f, and let f₁ = f. Then, for every D' of density 1 - (d/2n), let f₀(x) = f(x) if x ∈ D' and f₀(x) = 1 - f(x) otherwise. Clearly, f₀ is (d/2n)-far from any codeword. Invoking Theorem 3.3 (and using d ≤ n), we conclude that d/2n-testing the set of n-bit long codewords of C, while using q queries, requires randomness at least log₂(n/2q).

We mention that the codeword tests (for codes) that are obtained from typical PCP constructions (by following the transformation of Goldreich and Sudan [17]) achieve such (minimal) randomness complexity.

3. Monotone functions. A function f: {0,1}ⁿ → {0,1} is said to be monotone if f(x) ≤ f(y) for every x ≺ y, where ≺ denotes the natural partial order among strings (i.e., x₁ ··· x_n ≺ y₁ ··· y_n if x_i ≤ y_i for every i and x_i < y_i for some i). To see that the set of monotone functions is strongly evasive consider the all-one function f, and let f₁ = f. Then, for every D' of density 1/4, let f₀(σz) = f(σz) if {0z, 1z} ∩ D' ≠ Ø and f₀(σz) = 1 − σ otherwise. Note that if {0z, 1z} ∩ D' = Ø then f₀ must be modified at either 0z or 1z in order to obtain a monotone function. Thus, f₀ is 1/4-far from being monotone. Invoking Theorem 3.3, we conclude that 1/4-testing the set of monotone functions over {0,1}ⁿ, while using q queries, requires randomness at least log₂(2ⁿ/2q) = n − log₂q − 1.

We mention that the ϵ -tester for monotonicity of Goldreich, Goldwasser, Lehman, Ron, and Samorodnitsky [13] uses $O(n/\epsilon)$ queries and $n + \log_2 n$ random bits.

Turning back to graph properties, we focus on the bounded incidence lists model (of [15]), because the results of Section 3.1.2 do not apply to it. We mention a few properties of bounded-degree graphs that are strongly evasive in the (bounded) incidence lists model. Examples include connectivity and being Eulerian (or Hamiltonian), which can be demonstrated to be strongly evasive by starting with the *N*-cycle (and omitting edges). Additional examples such as planarity and bipartiteness can be demonstrated to be strongly evasive by starting with the empty graph (and adding edges). By invoking Theorem 3.3, we conclude that, in all these cases (which refer to a constant degree bound), $\Omega(1)$ -testing the set of N-vertex graphs that have the corresponding property by using q queries requires randomness at least $\log_2(N/q) - O(1)$. This lower bound is tight in some cases (e.g., connectivity (see the second author's thesis [27])) but not in others (e.g., bipartiteness (see Section 4.2)).

3.1.2 Relabeling-invariant properties

The second class that we consider consists of properties that are invariant under some "nice" relabeling of D, where a set of relabelings (or permutation) S is considered nice if a random $\pi \in S$ maps each element in D to the uniform distribution over D (i.e., for every $x, y \in D$ it holds that $\Pr_{\pi \in S}[\pi(x) = y] = |D|^{-1}$). We comment that a similar notion was considered by Kaufman and Sudan [19].

Definition 3.4 (invariant properties): Let S_D be a set of permutations over D. We say that the property Π is S_D -invariant if for every $f: D \to R$ and every $\pi \in S_D$ it holds that $f \in \Pi$ if and only $(f \circ \pi) \in \Pi$, where $(f \circ \pi)(x) = f(\pi(x))$.

We consider only sets S_D that correspond to a transitive group of permutations over D; that is, S_D is permutation group and for every $x, y \in D$ there exists a permutation $\pi \in S_D$ such that $\pi(x) = y$. Needless to say, the set of all permutations is a transitive group of permutations, but so are also many other permutation groups (e.g., the group of all cyclic permutations). Note that, for any transitive group S_D of permutations over D, it holds that $\Pr_{\pi \in S_D}[\pi(x) = y] = 1/|D|$, for every $x, y \in D$. (To see this, consider any $x, y, z \in S_D$, let $p_{x,y} = \Pr_{\pi \in S_D}[\pi(x) = y]$, and, using $\phi \in S_D$ such that $\phi(z) = y$, note that $p_{x,y} = \Pr_{\pi \in S_D}[(\phi \circ \pi)(x) = y] = \Pr_{\pi \in S_D}[\pi(x) = z] = p_{x,z}$.)

Theorem 3.5 Let S_D be a transitive group of permutations over D, and Π be a non-empty and S_D -invariant property of functions from D to R. Suppose that, for some $\sigma \in R$, the all- σ function is 2ϵ -far from Π . Then any non-adaptive ϵ -tester for Π that has query complexity q, must have randomness complexity at least $\log_2(|D|/q) - 1$.

Proof: Like the proof of Theorem 3.3, the current proof is based on deriving a contradiction from the hypothesis that the tester never examines most of the function (i.e., $|D'| \ll |D|$). The difference is in the way that this contradiction is derived, since we can no longer take the straightforward route offered by strong evasiveness.

Let T be an ϵ -tester for Π , and denote its query-complexity and randomness-complexity by q and r respectively. Since T is non-adaptive, its queries are oblivious of the oracle. For every $\omega \in \{0,1\}^r$, we denote by Q_{ω} the set of queries made by T when the outcome of its coin-tosses equals ω , and let $D' = \bigcup_{\omega \in \{0,1\}^r} Q_{\omega}$. Again, $|D'| \leq 2^r \cdot q$, and the theorem follows by proving that |D'| > |D|/2.

Let $f: D \to R$ be a function in Π with the maximum number of σ values, among all functions in Π . By the hypothesis, $|\{x \in D : f(x) \neq \sigma\}| > 2\epsilon |D|$. Suppose, for a moment, that $|\{x \in D \setminus D' : f(x) \neq \sigma\}| \ge \epsilon |D|$, and let h be defined such that h(x) = f(x) if $x \in D'$ and $h(x) = \sigma$ otherwise. Then (by the maximality of f), h is ϵ -far from Π . However, T^h behaves exactly as T^f (because hand f agree on D'), which yields a contradiction because T must accept f with probability at least 2/3 and accept h with probability at most 1/3.

It is left to prove that if $|D \setminus D'| \ge |D|/2$ then $|\{x \in D \setminus D' : f(x) \ne \sigma\}| \ge \epsilon |D|$. This does not necessarily hold, but we shall show that it holds when replacing f by another function in Π that also has a maximum number of σ values. Here we use the hypothesis that Π is an S_D -invariant

property, where S_D is a transitive group of permutations over D. Specifically, consider a random permutation $\pi \in S_D$, and let $f' = (f \circ \pi) \in \Pi$. Then, $f' \in \Pi$ and $|\{x \in D : f'(x) \neq \sigma\}| > 2\epsilon |D|$. On the other hand, since S_D is a transitive group of permutations over D, for every $x, y \in D$ it holds that $\Pr_{\pi \in S_D}[\pi(x) = y] = 1/|D|$. It follows that, for a random permutation $\pi \in S_D$, the expected size of $\{x \in D \setminus D' : f'(x) \neq \sigma\}$ equals

$$|D\setminus D'|\cdot rac{|D\setminus f^{-1}(\sigma)|}{|D|}\geq \epsilon |D|\,,$$

where the inequality is due to the hypotheses $|D \setminus D'| \ge |D|/2$ and $|D \setminus f^{-1}(\sigma)| > 2\epsilon |D|$. Thus, there exists a $f' \in \Pi$ such that $|\{x \in D \setminus D' : f'(x) \ne \sigma\}| \ge \epsilon |D|$, and the theorem follows.

Main application. As hinted in Section 3.1.1, the most appealing application of Theorem 3.5 is to testing graph properties in the adjacency matrix model (initiated by Goldreich, Goldwasser, and Ron [14]). In this model, N-vertex graphs are represented by Boolean functions defined over $[N] \times [N]$. For technical reasons, here (but not elsewhere) we represent such graphs as Boolean functions defined over the set of the $\binom{N}{2}$ (unordered) vertex-pairs, which is actually more natural (as well as non-redundant). (Using the set of $N^2 - N$ ordered pairs of non-identical elements would have worked too.) Note that the set of all permutations over [N] induces a transitive group of permutations over these pairs, where the permutation $\pi : [N] \to [N]$ induces a permutation that maps pairs of the form $\{i, j\}$ to $\{\pi(i), \pi(j)\}$. Indeed, any graph property is invariant under this group, and Theorem 3.5 can be applied whenever either the empty graph or the complete graph is far from the property. We note that all the (non-trivial) graph properties considered in [14, Sec. 6-9] fall into the latter category (and that the testers of [14] are all non-adaptive).

Corollary 3.6 (testing graph properties in the adjacency matrix model): Let Π be a graph property and suppose that either the empty graph or the complete graph is 2ϵ -far from Π . Then, any nonadaptive ϵ -tester for Π that has query complexity q, must have randomness complexity at least $2\log_2 N - \log_2 q - O(1)$.

Note that q adaptive Boolean queries can always be replaced by 2^q non-adaptive Boolean queries. We warn, however, that the more query-efficient transformation that replaces q adaptive (adjacency matrix) queries by $2q^2$ non-adaptive queries (see [3, 18]) is inapplicable here, because this transformation does not preserve the randomness-complexity.

Other applications. We note that any property that refers to sets of objects (e.g., sets of points as in Alon, Dar, Parnas, and Ron [2]) is invariant under the group of all permutations. Another application domain consists of matrix-properties that are preserved under row and column permutations.

Generalizations. Theorem 3.5 can be generalized to properties that are S_D -invariant under a set of permutations that is "sufficiently mixing" in the sense that a permutation selected uniformly in S_D maps each element of the domain to a distribution that has high min-entropy. For example, for a parameter $\alpha \ge 1$, it suffices that for every $x \in D$ and $y \in D$ it holds that $\Pr_{\pi \in S_D}[\pi(x) = y] \le \alpha/|D|$. In this case, we shall prove that $|D'| > |D|/2\alpha$, and a lower-bound of $\log_2(|D|/q) - \log_2(2\alpha)$ on the randomness-complexity follows. A different generalization is obtained by replacing σ with a set of values $S \subset R$ and referring to properties for which every function $f: D \to S$ is 2ϵ -far from the property.

3.1.3 Discussion

Although Theorems 3.3 and 3.5 are incomparable, most applications of Theorem 3.5 can be obtained also by using Theorem 3.3. Still, in some cases, it is easier to see that the conditions of Theorem 3.5 are met. For example, this is the case when the invariance of the property is obvious from the setting (e.g., as in the case of any graph property in the adjacency matrix model).

Both Theorems 3.3 and 3.5 yield a lower-bound of the form $\log_2(|D|/q) - O(1)$, which is independent of the proximity parameter ϵ . We believe that, for a wide range of parameters, the right lower-bound should be $\log_2(|D|/q) + \Omega(\log(1/\epsilon)) - O(1)$. Furthermore, in some cases where $q = \Omega(\epsilon^{-2})$, one may hope to obtain a $\log_2 |D| - O(1)$ lower-bound. Indeed, this is the case for average-estimation (see [22, 30]), which in turn is a special case of property testing. Specifically, in this case a lower-bound of $\log_2(D/q) + 2\log_2(1/\epsilon) - O(1)$ holds [22], whereas $q = O(\epsilon^{-2})$ holds when using query-efficient testers. (Note that $q = \Omega(\epsilon^{-2})$ must hold [9], and so this lower bound cannot be improved above $\log_2 |D| - O(1)$; indeed, the lower bound of [22] is tight (up to an additive constant) for any $q \ll |D|$.)

3.2 Upper Bounds

We start with a totally generic bound, and later focus on testing graph properties.

3.2.1 A generic bound

Recall that we refer to properties of functions from D to R. The following result can be easily proved by extending a similar result regarding samplers (presented in [9]), which in turn is proved using well-known techniques (cf., e.g., Newman [21]).

Theorem 3.7 If Π has an ϵ -tester that makes q queries then it has an ϵ -tester that makes O(q) queries and tosses $\log_2 |D| + \log_2 \log_2 |R| + O(1)$ coins. Furthermore, one-sided error and/or non-adaptivity are preserved.

For Boolean functions we get an upper-bound of $\log_2 |D| + O(1)$, which differs from the lowerbounds presented in Section 3.1 by an additive term of $\log_2 q + O(1)$. Indeed, the conjecture at the end of Section 3.1.3 shrinks the gap to a constant.

Proof: Let T be a tester as in the hypothesis, and suppose that it tosses r coins. Consider an 2^r -by- $|R|^{|D|}$ matrix in which the rows correspond to r-bit strings (representing possible outcomes of T's coin tosses) and the columns correspond to possible functions such that the entry (ω, f) equals the verdict of $T^f(\omega)$ (i.e., when T uses randomness ω and has oracle access to the function f). Note that the average values in any column that corresponds to a function in Π (resp., a function that is ϵ -far from Π) is at least 2/3 (resp., at most 1/3).

Using the probabilistic method (see [6]), we will show that there exists a multi-set Ω of $O(|D| \log |R|)$ rows such that, for each column, the average of this column *taken only over the rows in* Ω is 1/15-close to the average over the entire column. Using this set Ω , we consider the oracle machine that, when given access to any function f, selects uniformly $\omega \in \Omega$ and emulates $T^{f}(\omega)$. This machine accepts every $f \in \Pi$ with probability at least (2/3) - (1/15) = 3/5, rejects every f that is ϵ -far from Π with probability at least 3/5, and its randomness complexity is $\log_2 |\Omega| = \log_2 |D| + \log_2 \log_2 |R| + O(1)$. Using a randomness-efficient error-reduction (see Section 2.3), we obtain the desired tester. (Specifically, we reduce the error probability from 2/5 to 1/3, while increasing the number of queries by a multiplicative constant and maintaining the number of coin tosses.)

The probabilistic argument proceeds via a union bound over all possible $|R|^{|D|}$ functions. Fixing any function f, we consider the probability that, for a uniformly distributed multi-set Ω of size s, the following bad event occurs:

$$\left|2^{-r} \cdot \sum_{\omega \in \{0,1\}^r} T^f(\omega) - s^{-1} \cdot \sum_{\omega \in \Omega} T^f(\omega)\right| > \frac{1}{15}$$

$$\tag{1}$$

Using Chernoff bound, the probability that the bad event in Eq. (1) holds is at most $\exp(-\Omega(s))$. Thus, for $s = O(|D| \log |R|)$, we conclude that there exists a multi-set of size s such that, for every f, the bad event in Eq. (1) does not hold. The theorem follows.

Corollary. Applying Theorem 3.7 to testers of graph properties in the adjacency matrix model (of [14]), we conclude that if a property of N-vertex graphs is ϵ -testable using q queries then it has an ϵ -tester that makes O(q) queries and tosses $2\log_2 N + O(1)$ coins. We further discuss this model in Section 3.2.2.

3.2.2 Bounds for canonical testers of graph properties

The proof of Theorem 3.7 shows that for every tester T (of randomness complexity r) there exists a small set of coin-sequences Ω_T ($\subset \{0, 1\}^r$) that is essentially as good as the original set of coinsequences used by this tester (i.e., $\{0, 1\}^r$). This raises the question of whether there may exists a universal set Ω that is good for all testers (of randomness complexity r). Needless to say, the latter formulation is too general and is doomed to yield a negative answer (e.g., by considering, for any Ω , a pathological tester that behaves badly when fed with any sequence in Ω). Still such universal sets may exist for naturally restricted classes of testers.

One adequate class of testers was suggested by Goldreich and Trevisan [18], and it refers to testing graph properties in the adjacency matrix model. A canonical ϵ -tester for a property Π of *N*-vertex graphs is determined by an integer *k* and a property Π' of *k*-vertex graphs. Such a tester, sometimes referred to as *k*-canonical, selects uniformly a set of *k* vertices in the input graph *G* and accepts *G* if and only if the corresponding induced (*k*-vertex) subgraph has the property Π' . It was shown in [18] that if Π is ϵ -testable with query complexity *q* then Π has a *k*-canonical ϵ -tester with k = O(q). Thus, it is natural to consider the notion of a "universal set" of *k*-subsets of [*N*] that is good for all *k*-canonical testers.

Definition 3.8 A multi-set $\Omega \subseteq \{S \subset [N] : |S| = k\}$ is called (ϵ, k) -universal if for every property Π of N-vertex graphs and for every k-canonical ϵ -tester for Π , denoted T, the following holds:

- 1. For every G that has property Π , it holds that $\Pr_{\omega \in \Omega}[T^G(\omega) = 1] \ge 3/5$, where $T^G(\omega)$ denotes the execution of T when given the coin-sequence ω and oracle access to G.
- 2. For every G that is ϵ -far from property Π , it holds that $\Pr_{\omega \in \Omega}[T^G(\omega) = 1] \leq 2/5$.

Using an (ϵ, k) -universal set, we can reduce the randomness complexity of any k-canonical ϵ -tester T by selecting uniformly $\omega \in \Omega$ and emulating $T(\omega)$. The residual oracle machine, denoted T', is essentially an ϵ -tester for the same property, except that T' may err with probability at most 2/5 (rather than 1/3). Needless to say, T' has randomness complexity $\log_2 |\Omega|$ and query complexity $\binom{k}{2}$. Furthermore, T' preserves the possible one-sided error of T.

Clearly, the set of all k-subsets is (ϵ, k) -universal, because using this set coincides with the definition of a k-canonical ϵ -tester. We seek (ϵ, k) -universal sets that are much smaller; specifically,

by prior results we may hope to have (ϵ, k) -universal sets of size $O(N^2)$. By extending the proof of Theorem 3.7, we can prove the following result.

Theorem 3.9 There exist (ϵ, k) -universal sets (of subsets of [N]) having size $O(2^{k^2} + N^2)$.

The randomness complexity of the derived ϵ -tester is $\max(k^2, 2\log_2 N) + O(1)$, which is typically smaller than the randomness complexity of the k-canonical ϵ -tester (i.e., $k\log_2 N$). For $k = o(\sqrt{\log N})$, which holds whenever k only depends on ϵ (and ϵ is constant) as in [14, 3, 4], we get randomness-complexity $2\log_2 N + O(1)$, which is optimal since the domain size is N^2 .

Proof: The key observation is that a k-canonical tester is determined by the property Π' that it decides (for the induced k-vertex subgraph), while Π' can be described by $K = 2^{\binom{k}{2}} < 2^{k^2}$ bits which determine for each k-vertex graph whether it is in Π' . Thus, when applying a union bound as in the proof of Theorem 3.7, the number of k-canonical testers that we need to consider is less than 2^K . Hence, it suffices to have $2^K \cdot 2^{N^2} \cdot \exp(-\Omega(s)) < 1$, where 2^K upper-bounds the number of testers, 2^{N^2} upper-bounds the number of N-vertex graphs, and $\exp(-\Omega(s))$ upper-bounds the probability that a multi-set of size s is bad (as in Eq. (1)) with respect to a fixed tester and a fixed graph. Using $s = O(K + N^2)$, the claim follows.

Open problems. Can the upper-bound of Theorem 3.9 be improved; in particular, do there exist (ϵ, k) -universal sets (of subsets of [N]) having size $O(\text{poly}(k) \cdot N^2)$ or even $O(N^2)$? Can universal sets of small size (e.g., as in Theorem 3.9) be efficiently constructed?

Extension. Theorem 3.9 extends to any class of non-adaptive testers (for any property of functions from D to R) whose final decision only depends on the oracle answers. The point is that each such tester that makes q queries can be described by a function $f : R^q \to \{0, 1\}$, and thus the number of such testers is $2^{|R|^q}$. Hence, the size of the corresponding "universal set" is $O(|R|^q + |D| \log |R|)$.

4 Specific Algorithms: The Case of Bipartiteness

In this section we demonstrate two approaches to reducing the randomness-complexity of testers. Section 4.2 demonstrates the approach of merely providing a randomness-efficient implementation of some random features that are used in the analysis of the original tester. In contrast, Section 4.1 demonstrates the approach of redesigning the tester (while, indeed, benefiting from ideas that underly the design of the original tester).

In both sections we consider testing graph properties, but in two different standard models: In Section 4.1 we refer to the adjacency matrix model (introduced in Goldreich, Goldwasser, and Ron [14]), while in Section 4.2 we refer to the bounded-degree model (introduced in Goldreich and Ron [15]). In both sections, we focus on the problem of testing bipartiteness. Further details and additional testers are provided in the second author's thesis [27]. We make extensive use of randomness-efficient hitters as defined and discussed in Section 2.2.

4.1 In the Adjacency Matrix Model

In the adjacency matrix model an N-vertex graph G = (V, E) is represented by the Boolean function $g : [N] \times [N] \to \{0, 1\}$ such that g(u, v) = 1 if and only if u and v are adjacent in G (i.e., $\{u, v\} \in E$). In this section we present a randomness-efficient bipartite tester for graphs in the adjacency matrix

model. This tester is strongly influenced by the tester of Goldreich, Goldwasser, and Ron [14], but differs from it in significant ways. Still, it is instructive to start with a description of the tester of [14], hereafter referred to as the GGR tester.

4.1.1 The GGR tester

Essentially, the GGR tester selects a random set of $\tilde{\Theta}(\epsilon^{-2})$ vertices, inspects the subgraph of G induces by this set, and accepts if and only if this induced subgraph is bipartite. The analysis in [14] actually refers to the following description, which also has a lower query-complexity.

Algorithm 4.1 On input parameters N and ϵ , and oracle access to an adjacency predicate of an N-vertex graph, G = (V, E), proceed as follows:

- 1. Uniformly select a sample U of $\tilde{\Theta}(\epsilon^{-1})$ vertices.
- 2. Uniformly select a sample S of $\widetilde{\Theta}(\epsilon^{-2})$ vertex-pairs.
- 3. For each $u \in U$ and $(v_1, v_2) \in S$, check whether $\{u, v_1\}, \{u, v_2\}$ and $\{v_1, v_2\}$ are edges.
- 4. Accept if and only if the subgraph viewed in Step 3 is bipartite.

Clearly, this algorithm never rejects a bipartite graph, and thus its analysis focuses on the case that G is ϵ -far from being bipartite. One key observation is that each 2-partition, (U_1, U_2) , of U induces a 2-partition of the entire graph in which all neighbors of U_1 are on one side and all the other vertices are on the other side. A pair of vertices (v_1, v_2) detects that the latter partition is not a valid 2-coloring of G if there exists $u_1, u_2 \in U_1$ (resp., $u_1, u_2 \in U_2$) such that $\{u_1, v_1\}, \{v_1, v_2\}$ and $\{v_2, u_2\}$ are all edges of G. In such a case, we call the pair (v_1, v_2) a witness against (U_1, U_2) . The analysis in [14] shows that if G is ϵ -far from being bipartite then, with high probability, for every 2-partition of U there exists a pair in S that is a witness against this 2-partition. Let us briefly recall how this is done.

The first step is proving that, with high probability (say, with probability at least 5/6), the set U dominates all but an $\epsilon/8$ fraction of the vertices of G that have degree at least $\epsilon N/8$, where a set U dominates a vertex v if v is adjacent to some vertex in U. This step is quite straightforward. The next step is proving that this implies that for every 2-partition of U there exists at least $\epsilon N^2/2$ (ordered) vertex-pairs that are each a witness against this 2-partition. The implication is proved by confronting the following two facts:

- 1. Since G is ϵ -far from being bipartite, the 2-partition of V induced by any 2-partition of U has at least ϵN^2 (ordered) vertex-pairs that reside on the same side of the partition and yet are connected by an edge.
- 2. The number of (ordered) vertex-pairs (v_1, v_2) such that $\{v_1, v_2\} \in E$ but either v_1 or v_2 is not dominated by U is at most $\epsilon N^2/2$, because each low-degree vertex contributes at most $\epsilon N/4$ such (ordered) pairs and there are at most $\epsilon N/8$ high-degree vertices that are not dominated by U.

Having established the existence of at least $\epsilon N^2/2$ vertex-pairs that constitute a witness against any fixed 2-partition of U, it is clear that each random pair of vertices will be a witness with probability at least $\epsilon/2$, and selecting enough random pairs will do the job. The point, however, is that we need to rule out each of the $2^{|U|}$ possible 2-partitions of U. Thus, the number of selected pairs is set such that the probability that we do not find a witness against any specific 2-partition is smaller than $2^{-|U|}$. Indeed, setting $|S| = O(|U|/\epsilon)$ will do. This completes our review of [14].

As stated in Section 1.3, the foregoing approach supports a randomness-efficient implementation (of Algorithm 4.1). Specifically, U needs to be selected so that sets of density $\epsilon/8$ are avoided with probability at most $\epsilon/48$, while S is selected such that sets of density $\epsilon/8$ are avoided with probability at most $2^{-|U|}/6$. This yields randomness-complexity $\tilde{O}(\epsilon^{-1}) + O(\log N)$. The problem with the foregoing approach is that it is impossible to implement it using randomness-complexity below |U|, which in turn is $\Omega(\epsilon^{-1})$. Recall, however, that our aim is to obtain randomness-complexity that is linearly related to $O(\log(1/\epsilon))$.

4.1.2 A warm-up: randomness-efficient tester of query complexity $\widetilde{O}(\epsilon^{-4})$

A closer look at the foregoing argument reveals that a pair (v_1, v_2) such that $\{u_1, v_1\}, \{v_1, v_2\}$ and $\{v_2, u_2\}$ are all edges of G is not merely a witness against a specific 2-partition of U that places u_1 and u_2 on the same side. It is actually a witness against any 2-partition of U that places u_1 and u_2 on the same side. Viewed from a different perspective, such a pair (v_1, v_2) imposes a constraint on the "relevant" 2-partition of U; the constraint being that u_1 and u_2 should not be placed on the same side. It will be useful to consider the graph of these constraints, which has the vertex-set U and edges between each pair of vertices to which such a constraint is applied (i.e., there is an edge between u_1 and u_2 if there exists a pair $(v_1, v_2) \in V \times V$ that imposes a constraint on the pair (u_1, u_2)). Indeed, the 2-partitions of U that satisfy the set of these constraints are exactly the 2-colorings of this auxiliary graph.

The foregoing perspective suggests that it may be useful to try to accumulate constraints. At the very extreme, the graph of constraints will not be bipartite, which definitely allows us to reject (because it indicates that there are witnesses against each 2-partition of U).² Discarding this case, we consider another extreme case in which the graph of constraints is connected, leaving us with a single allowed 2-partition of U (i.e., a single 2-coloring of the constraint graph), which can be checked as in Algorithm 4.1. The point, however, is that in this case it will suffice to set $|S| = O(\epsilon^{-1})$ and more importantly to have a sample that rules out the remaining partition with constant probability (rather than with probability $2^{-|U|}$). This opens the door to a randomness-efficient implementation.

But what if the graph of constraints that we found is not connected? Unless this event is due to sheer lack of luck, it indicates that there are few pairs in $V \times V$ that impose constraints regarding vertex-pairs in $U \times U$ that are in different connected components of the constraint graph. This implies that, for every 2-partition of U that is consistent with the constraint graph (i.e., every 2-coloring of this graph), there are many pairs in $V \times V$ that constitute a witness against the 2-partition of some of the connected components. That is, each such pair imposes a constraint that refers to vertices that reside in the same connected component, and furthermore this constraint contradicts the constraints that are already present regarding this connected component.

Needless to say, for the foregoing to work, we should determine adequate thresholds for the notion of "few pairs in $V \times V$ that impose a constraint regarding vertex-pairs" (in $U \times U$). Let us start by spelling out the notion of imposing (or rather forcing) a constraint. We say that the pair $(v_1, v_2) \in V \times V$ constrains the pair $(u_1, u_2) \in U \times U$ if $\{u_1, v_1\}, \{v_1, v_2\}$ and $\{v_2, u_2\}$ are all edges of G. Next, we say that a pair $(u_1, u_2) \in U \times U$ is ρ -constrained if there are at least $\rho \cdot N^2$ vertex-pairs in $V \times V$ that constrain (u_1, u_2) . Leaving ρ unspecified for a moment, we make the following observations:

²We note that it follows from [5] that this case holds with high probability provided that U is selected uniformly among all $\tilde{O}(1/\epsilon)$ -size subsets. However, we cannot afford to select U in this manner.

- 1. Using a sample of $O(\rho^{-1} \cdot \log |U|)$ vertex-pairs in $V \times V$, with high probability, it holds that for every ρ -constrained pair $(u_1, u_2) \in U \times U$, the sample contains a pair that constrains (u_1, u_2) . This holds even if the sample is generated using a randomness-efficient hitter (which hits any set of density ρ with probability at least $1 - (|U|^{-2}/10)$, using randomness-complexity $O(\log |V| + \log |U|) = O(\log |V|)$). The point is that there are at most $|U|^2$ relevant pairs (i.e., pairs that are ρ -constrained), and we may apply a Union Bound as long as we fail on each such pair with probability at most $|U|^{-2}/10$ (or so).
- 2. Consider the graph $G_{U,\rho}$ consisting of the vertex-set U and edges corresponding to the ρ constrained pairs of vertices. Then, the number of vertex-pairs in $V \times V$ that constrain some
 pair of vertices (in U) that does not belong to the same connected component of $G_{U,\rho}$ is at
 most $|U|^2 \cdot \rho N^2$.

Recall that if G is ϵ -far from bipartite and U is good (i.e., U dominates almost all highdegree vertices) then, for every 2-partition of U, there are at least $\epsilon N^2/2$ pairs that constrain some pair of vertices that are on the same side of this 2-partition. It follows that at least $((\epsilon/2) - |U|^2 \rho) \cdot N^2$ of these pairs constrain pairs that are in the same connected component of $G_{U,\rho}$. Setting $\rho = \epsilon/(4|U|^2)$, we need to hit a set of density $\epsilon/4$, which is easy to do using a randomness-efficient hitter.

This analysis leads to an algorithm that resembles Algorithm 4.1, except that it uses a secondary sample S that has different features than in the original version. In Algorithm 4.1 the set S had to hit any fixed set of density $\epsilon/2$ with probability at least $1 - 2^{-|U|}$. Here the set S needs to hit any fixed set of density $\rho = \epsilon/(4|U|^2) < \epsilon^3$ with probability at least $1 - (|U|^{-2}/10)$. Thus, while in Algorithm 4.1 we used $|S| = O(|U|/\epsilon)$ but generating the set S required at least |U| random bits, here $|S| = \tilde{O}(|U|^2/\epsilon) = \tilde{O}(\epsilon^{-3})$ but generating the set S can be done using $O(\log N)$ random bits. (The set U is generated with the same aim as in Algorithm 4.1; that is, hitting a set of density ϵ with probability at least $1 - \epsilon^{-1}$. Such a set can be generated using $O(\log N)$ random bits.)

Thus, we obtain a (computationally efficient) ϵ -tester with randomness-complexity $O(\log N)$ and query-complexity $O(|U| \cdot |S|) = \tilde{O}(\epsilon^{-4})$. Our aim in the next section is to reduce the querycomplexity to $\tilde{O}(\epsilon^{-3})$ while essentially maintaining the randomness-complexity.

4.1.3 The actual algorithm: randomness-efficient tester of query complexity $\widetilde{O}(\epsilon^{-3})$

The query-complexity bottleneck in Section 4.1.2 is due to the size of S, which in turn needs to hit sets of density $\rho = O(\epsilon^3)$. Our improvement will follow by using a larger value of the threshold ρ (essentially $\rho = O(\epsilon^2)$). Recall that in Section 4.1.2 we used $\rho = O(\epsilon^3)$ in order to bound the total number of pairs that constrain pairs that are not ρ -constrained. Thus, using $\rho = O(\epsilon^3)$ seems inherent to an analysis that refers to each pair separately, and indeed we shall deviate from that paradigm in this section.

The planned deviation is quite natural. After all, we not not care about having specific edges in our constraint graph, but rather care about the connected components of that graph. For example, looking at any vertex $u \in U$, any pair in $V \times V$ that constrains any pair (u, u'), where $u' \in U \setminus \{u\}$, increases the connected component in which u resides. That is, let $\gamma(u_1, u_2)$ denote the fraction of vertex-pairs in $V \times V$ that constrain (u_1, u_2) , and recall that a pair (u_1, u_2) was called ρ -constrained if $\gamma(u_1, u_2) \geq \rho$. Thus, we (tentatively) say that $u \in U$ is ρ -constrained if $\sum_{u' \in U \setminus \{u\}} \gamma(u, u') \geq \rho$. Let us now see what happens.

1. Using a sample of $O(\rho^{-1} \cdot \log |U|)$ vertex-pairs in $V \times V$, with high probability, it holds that for every ρ -constrained vertex $u \in U$, the sample contains a pair that constraints (u, u'), for some

 $u' \in U \setminus \{u\}$. Again, this holds even if the sample is generated using a randomness-efficient hitter.

2. The number of vertex-pairs in $V \times V$ that constrain some pair of vertices $(u_1, u_2) \in U \times U$ such that either u_1 or u_2 is not ρ -constrained is at most $2|U| \cdot \rho N^2$. This means that we can ignore such vertex-pairs (in $V \times V$) even when setting $\rho = O(\epsilon/|U|)$ or so.

Thus, taking a sample S' as in Item 1, will result in having a constraint graph $G_{U,S'}$ in which each ρ -constrained vertex resides in non-singleton connected components. In particular, the number of non-singleton connected components is at most |U|/2.

Note, however, that unlike in Section 4.1.2, the foregoing facts do not yield an upper-bound on the number of vertex-pairs in $V \times V$ that constrain some pair of vertices (in U) that does not belong to the same connected component of $G_{U,S'}$. Loosely speaking, we shall iterate the same process on the non-singleton connected components of $G_{U,S'}$, while recalling that the only vertices that form singleton connected components in $G_{U,S'}$ are not ρ -constrained (and thus can be ignored). This suggests an iterative process, which will halt after at most $\log_2 |U|$ iterations in a situation analogous to having no ρ -constrained vertices. At this point we may proceed with a final sample of pairs that, with high probability, will yield a constraint that conflicts with the existing ones.

Clarifying the foregoing iterative process requires generalizing the notion of ρ -constrained vertices such that it will apply to the connected components determined in the previous iteration. Consider a partition of U, denoted $\overline{U} = (U^{(0)}, U^{(1)}, ..., U^{(k)})$, where $U^{(0)}$ may be empty and k may equal 0, but for every $i \in [k]$ it holds that $U^{(i)} \neq \emptyset$. In the first iteration, we use $\overline{U} = (\emptyset, \{u_1\}, ..., \{u_t\})$, where $U = \{u_1, ..., u_t\}$. In later iterations, $U^{(1)}, ..., U^{(k)}$ will correspond to connected components of the current constraint graph and $U^{(0)}$ will contain vertices that were cast aside at some point.

Definition 4.2 (being constrained w.r.t a partition): For $i \in \{0, 1, ..., k\}$, we say that $u \in U^{(i)}$ is ρ -constrained w.r.t \overline{U} if $\sum_{u' \in U'} \gamma(u, u') \geq \rho$, where $U' = \bigcup_{j \in [k] \setminus \{i\}} U^{(j)}$. Recall that $\gamma(u_1, u_2)$ denote the fraction of vertex-pairs in $V \times V$ that constrain (u_1, u_2) , where the pair $(v_1, v_2) \in V \times V$ constrains the pair $(u_1, u_2) \in U \times U$ if $\{u_1, v_1\}, \{v_1, v_2\}$ and $\{v_2, u_2\}$ are all edges of G.

We stress that the foregoing sum does not include vertices in either $U^{(0)}$ or $U^{(i)}$. Our analysis will refer to the following algorithm, which can be implemented within randomness-complexity $O(\log(1/\epsilon)) \cdot \log_2 N$ and query-complexity $\tilde{O}(\epsilon^{-3})$.

Algorithm 4.3 (The bipartite tester, revised):

- 1. Select a sample U of $\tilde{O}(\epsilon^{-1})$ vertices by using a hitter that hits any set of density $\epsilon/8$ with probability at least $1 (\epsilon/100)$.
- 2. For $i = 1, ..., \ell + 1$, where $\ell = \log_2 |U|$, select a sample S_i of $\tilde{O}(\epsilon^{-2})$ vertex-pairs by using a hitter that hits any set of density $\rho = \epsilon / \tilde{O}(|U|)$ with probability at least $1 \tilde{O}(|U|)^{-1}$. (This hitter has randomness-complexity $O(\log N + \log |U|) = O(\log N)$.) Let $S = \bigcup_{i=1}^{\ell+1} S_i$.
- 3. For each $u \in U$ and $(v_1, v_2) \in S$, check whether $\{u, v_1\}, \{u, v_2\}$ and $\{v_1, v_2\}$ are edges.
- 4. Accept if and only if the subgraph viewed in Step 3 is bipartite.

Needless to say, the peculiar way in which S is selected is aimed to support the analysis.

Lemma 4.4 If G is ϵ -far from being bipartite then Algorithm 4.3 rejects with probability at least 2/3.

Proof: We may assume that U is good in the sense that it dominates all but $\epsilon N/8$ of the vertices that have degree at least $\epsilon N/8$. As argued above (and shown in [14]), there are at most $\epsilon N^2/2$ vertex pairs that have an endpoint that is not dominated by $U = \{u_1, ..., u_t\}$. Starting with $\overline{U} = (\emptyset, \{u_1\}, ..., \{u_t\})$, we shall proceed in iterations proving that in each iteration one of the following two events occur:

- 1. There are $\Omega(\epsilon N^2)$ vertex pairs that form constraints that contradict the existing constraints. In this case, with very high probability, the algorithm will select such a pair and will reject (because the subgraph that it sees is not 2-colorable).
- 2. There exist ρ -constrained vertices with respect to the current partition $\overline{U} = (U^{(0)}, U^{(1)}, ..., U^{(k)})$, where $U^{(1)}, ..., U^{(k)}$ are connected components of the current constraint graph and $U^{(0)}$ contains vertices that were cast aside in previous iterations. We shall also show that ρ -constrained (w.r.t \overline{U}) vertices cannot be in $U^{(0)}$. In this case, with very high probability, the algorithm will find new constraints and in particular it will find such a constraint between every ρ -constrained (w.r.t \overline{U}) vertex and some vertex that is in one of the other k connected components.

We shall shortly take a closer look at what happens in the second case (i.e., Case 2) and prove that indeed at least one of the foregoing cases must hold. But before doing so, we note that the second case (i.e., Case 2) becomes impossible once we reach a situation in which k = 1, at which point the algorithm must reject due to the first case (i.e., Case 1).

Let us first take a closer look at what happens in Case 2. Suppose that $u \in U^{(i)}$ is ρ -constrained w.r.t the current \overline{U} . Then by the foregoing, due to a newly found constraint, vertex u gets connected to some vertex in $\bigcup_{j \in [k] \setminus \{i\}} U^{(j)}$. This means that each $U^{(i)}$ $(i \neq 0)$ that contains some ρ -constrained vertex gets merged to some $U^{(j)}$ $(j \neq 0$ and $j \neq i$). We will not add any constraint that refers to vertices that were cast aside (i.e., those in $U^{(0)}$). Thus, vertices that were cast aside in the past (since they were not ρ -constrained w.r.t a previous partition) will remain in $U^{(0)}$, and indeed they are also not ρ -constrained w.r.t any later partition. (This is the case because in a later partition, some components get merged and some move to $U^{(0)}$, which can only decrease the "constrains count" towards being ρ -constrained.) For $i \neq 0$, if $U^{(i)}$ was not merged with any other $U^{(j)}$ $(j \neq 0$ and $j \neq i$) then it contains no ρ -constrained vertex, and we cast it aside (i.e., move it to the new $U^{(0)}$). Thus, in each iteration, the number of connected components not cast aside (i.e., k) shrinks by a factor of at least two (because each such connected component merges with at least one such other connected component).

We now prove that at least one of the two aforementioned conditions must hold. Looking at the current partition \overline{U} , we first note that if one of the connected components (including those contained in $U^{(0)}$) is not bipartite then we already have a set of constraints that is self-contradictory (i.e., does not allow a 2-coloring of the subgraph we have seen so far). This situation is a special case of Case 1, and indeed in this sub-case the algorithm rejects. Disposing of this sub-case, we now consider an arbitrary 2-coloring of the constraint graph, and the 2-partition that it induces on the rest of G (i.e., we put on the first side all the vertices that are *dominated by some vertex of* U that was colored by the second color). Then, there are at least ϵN^2 vertex-pairs that are adjacent and were put on the same side, and at least $\epsilon N^2/2$ of these vertex-pairs have both its vertices dominated by U. Each such (v_1, v_2) is of one of the following two types.

(i) The vertex-pair (v_1, v_2) constrains a pair of vertices (u_1, u_2) where both vertices are in the same connected component of the constraint graph. As showed next, such a pair imposes a constraint that contradicts the constraints of the current graph. Thus, this pair contributes to the pairs counted in Case 1.

To see that the said constraint contradicts the constraints of the current graph, recall that since (v_1, v_2) constrains the pair $(u_1, u_2) \in U \times U$ it holds that the edges $\{u_1, v_1\}, \{v_1, v_2\},$ and $\{v_2, u_2\}$ form an odd-length path between u_1 and u_2 . On the other hand, v_1 and v_2 were placed on the same side of the 2-partition of V, which implies that u_1 and u_2 were assigned the same color by a 2-coloring of the current constraint graph. Since u_1 and u_2 are in the same connected component of that graph, it follows that they are connected by an even-length path (which reflects an even-length path in G). Thus, the new set of constraints form an odd-length cycle.

(ii) The vertex-pair (v_1, v_2) constrains a pair of vertices (u_1, u_2) that belong to different connected component of the constraint graph. As showed next, the existence of more than $\epsilon N^2/4$ such pairs implies Case 2 (i.e., the existence of ρ -constrained vertices, which in particular are not in $U^{(0)}$).

We first recall that a vertex in $U^{(0)}$ can not be ρ -constrained with respect to the current partition, because it is not ρ -constrained with respect to some previous partition and because the previous partition allows more pairs to be counted.

As for the main claim, note that each pair of the current type is counted towards determining whether u_1 (resp., u_2) is ρ -constrained with respect to the current partition. The total "pair count" of each vertex that is not ρ -constrained is smaller than ρN^2 . Thus, for $\rho = \epsilon/(4|U|)$, there are less than $|U| \cdot \rho N^2 = \epsilon N^2/4$ pairs of the current type that refer to vertices that are not ρ -constrained. It follows if there are more than $\epsilon N^2/4$ pairs of the current type, then ρ -constrained vertices must exist, which imply that Case 2 holds.

We conclude that either there are more than $\epsilon N^2/4$ vertices of type (ii), which imply that Case 2 holds, or there are more than $\epsilon N^2/4$ vertices of type (i), which imply that Case 1 holds.

Recall that if Case 2 holds then the number of non-discarded connected components (i.e., k) shrinks by a factor of at least 2. Thus, after $\log_2 |U|$ iterations, the current partition must satisfy $k \leq 1$, and thus Case 2 cannot hold in the next iteration. The lemma follows.

Conclusion. Using Algorithm 4.3 and its analysis as provided by Lemma 4.4, we obtain:

Theorem 4.5 There exists a bipartite tester (in the adjacency matrix model) of time-complexity $\operatorname{Poly}(\epsilon^{-1} \cdot \log N)$, query-complexity $\widetilde{O}(\epsilon^{-3})$ and randomness-complexity $O(\log(1/\epsilon)) \cdot \log_2 N$. Furthermore, as Algorithm 4.1, this tester always accepts a bipartite graph, and in case of rejection it provides a witness of length $\widetilde{O}(\epsilon^{-2}) \cdot \log_2 N$ (that the graph is not bipartite).

Theorem 4.5 improves over the randomness-efficient implementation of Algorithm 4.1 (which has randomness-complexity $\tilde{O}(\epsilon^{-1}) + O(\log N)$) whenever $\epsilon < 1/\tilde{O}(\log N)$.

Open problem. Needless to say, we are aware of the bipartite tester of Alon and Krivelevich [5], which has better query-complexity than the GGR tester (as well as our tester). Specifically, the query-complexity of the tester of [5] is $\tilde{O}(\epsilon^{-2})$ rather than $\tilde{O}(\epsilon^{-3})$. Theorem 3.7 implies that the tester of [5] has a randomness-efficient implementation, but it does not provide an explicit one. We conjecture that there exists a randomness-efficient bipartite tester that has query-complexity $\tilde{O}(\epsilon^{-2})$ and time-complexity poly($\epsilon^{-1} \log N$).

4.2 In the Bounded-Degree Model

The bounded-degree model refers to a fixed degree bound, denoted d. An N-vertex graph G = (V, E) (of maximum degree d) is represented in this model by a function $g : [N] \times [d] \rightarrow \{0, 1, ..., N\}$ such that $g(v, i) = u \in [N]$ if u is the i^{th} neighbor of v and g(v, i) = 0 if v has less than i neighbors. In this section we provide a randomness-efficient implementation of the bipartite tester of Goldreich and Ron [16], which refers to the bounded-degree model. Thus, we start with a description of that tester.

Algorithm 4.6 (The bipartite tester of [16]): On input parameters N, d, ϵ , and oracle access to an incidence function for an N-vertex graph, G = (V, E), of degree bound d, repeat $T \stackrel{\text{def}}{=} \Theta(\frac{1}{\epsilon})$ times:

- 1. Uniformly select a ("start") vertex s in V.
- 2. (Try to find an odd-length cycle through vertex s):
 - (a) Perform $K \stackrel{\text{def}}{=} \operatorname{poly}((\log N)/\epsilon) \cdot \sqrt{N}$ random walks starting from vertex s, where each walk is of length $L \stackrel{\text{def}}{=} \operatorname{poly}((\log N)/\epsilon)$.
 - (b) Let R_0 (respectively, R_1) denote the set of vertices that were reached from vertex s in an even (respectively, odd) number of steps in any of these walks.
 - (c) If $R_0 \cap R_1$ is not empty then reject.

If the algorithm did not reject in any one of the above T iterations, then it accepts.

Clearly, this algorithm never rejects a bipartite graph. Indeed, the analysis of [16] focuses on the case that the graph G is ϵ -far from bipartite, and shows that the algorithm will reject G with high probability. The rather involved analysis breaks down to two complementary facts that refer to a notion of a good start vertex. Loosely speaking, a start vertex is called good if, when the tester selects it in Step 1, the probability that the tester finds an odd-length cycle in Step 2 is somewhat small (say, below 1/10). This is indeed the definition used in [16, Sec. 4], but the analysis there actually refers to a technical condition that is stated in [16, Lem. 4.5] and refers to what happens when taking two independent random walks from the start vertex. Thus, here we call a start vertex good if it does not satisfy the said condition (stated in [16, Lem. 4.5]), and it is not good if it satisfies this condition.

Most of [16, Sec. 4] is devoted to establishing the fact that if G is ϵ -far from bipartite then an $\Omega(\epsilon)$ fraction of the vertices are not good. It is crucial for us that this technically involved analysis (provided in [16, Sec. 4.2–4.4]) does not refer at all to the algorithm; it rather refers to the definition of a good vertex, which (as stressed above) refers to a mental experiment in which one takes two independent random walks from this vertex. Thus, this analysis remains intact regardless of how we chose to implement Algorithm 4.6.

The complementary fact regarding good vertices is that when the tester selects a vertex that is not good (in Step 1), the probability that it finds an odd-length cycle in Step 2 is not too small (say, at least 1/10). Indeed, this fact refers to Algorithm 4.6 itself, but its rather simple proof (provided in [16, Sec. 4.5]) only presumes that the K random walks are distributed in a 4-wise independent manner. Specifically, the analysis in [16, Sec. 4.5] defines a random variable for each pair of walks such that this random variable represents the event of finding an odd-length cycle via the corresponding two walks. Then, Chebyshev's Inequality is applied while relying on the expectation and variance of the sum of these random variables. As one may guess, the said expectation and variance are computed by only relying on the expectation of the individual random variables and the co-variances of all possible pairs of random variables. Thus, the analysis remains valid as long as the said expectation and co-variance maintain their value, which is definitely the case provided that each pair of random variables maintains its behavior. Noting that each pair of random variables refers to at most four different random walks, we establish our claim that the analysis of [16] only presumes that the K random walks are distributed in a 4-wise independent manner.

The foregoing discussion suggests the following implementation of Algorithm 4.6. For Step 1 use a randomness-efficient hitter that hits any set of density $\Omega(\epsilon)$ with constant probability. More importantly, for Step 2 use a randomness-efficient construction of K four-wise independent random strings, each specifying a random walk of length L (i.e., each being a string of length $L \log_2 d$). By the foregoing discussion, this implementation preserves the performance guarantees of Algorithm 4.6; that is, this implementation is also an ϵ -tester for bipartiteness. The crucial point, however, is that Step 2 is now implemented using $4 \cdot L \log_2 d = \text{poly}((\log N)/\epsilon)$ random coins (rather than $K \cdot L \log_2 d = \Omega(\sqrt{N})$ random coins). Thus, we obtain:

Theorem 4.7 There exists a bipartite tester (in the incidence function model) of time-complexity $\operatorname{poly}(\epsilon^{-1} \cdot \log N) \cdot \sqrt{N}$ and randomness-complexity $\operatorname{poly}(\epsilon^{-1} \cdot \log N)$. Furthermore, as Algorithm 4.6, this tester always accepts a bipartite graph, and in case of rejection it provides a witness of length $\operatorname{poly}(\epsilon^{-1} \cdot \log N)$ (that the graph is not bipartite).

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References

- N. Alon, L. Babai and A. Itai. A fast and Simple Randomized Algorithm for the Maximal Independent Set Problem. J. of Algorithms, Vol. 7, pages 567–583, 1986.
- [2] N. Alon, S. Dar, M. Parnas, and D. Ron. Testing of Clustering. SIAM Journal on Discrete Mathematics, Vol. 16 (3), pages 393-417, 2003.
- [3] N. Alon, E. Fischer, M. Krivelevich, and M. Szegedy. Efficient testing of large graphs. In 40th IEEE Symposium on Foundations of Computer Science, pages 645–655, 1999.
- [4] N. Alon, E. Fischer, I. Newman, and A. Shapira. A Combinatorial Characterization of the Testable Graph Properties: It's All About Regularity. In 38th ACM Symposium on the Theory of Computing, pages 251-260, 2006.
- [5] N. Alon and M. Krivelevich. Testing k-Colorability. SIAM Journal on Discrete Mathematics, Vol. 15 (2), pages 211-227, 2002.
- [6] N. Alon and J.H. Spencer. The Probabilistic Method. John Wiley & Sons, Inc., 1992. Second edition, 2000.
- [7] E. Ben-Sasson, M. Sudan, S. Vadhan, and A. Wigderson. Randomness-efficient low degree tests and short PCPs via epsilon-biased sets. In Proc. 35th ACM Symposium on the Theory of Computing, June 2003, pp. 612–621.
- [8] M. Blum, M. Luby, and R. Rubinfeld. Self-testing/correcting with applications to numerical problems. Journal of Computer and System Science, Vol. 47, pages 549-595, 1993.
- [9] R. Canetti, G. Even, and O. Goldreich. Lower Bounds for Sampling Algorithms for Estimating the Average. Information Processing Letters, Vol. 53, pages 17–25, 1995.
- [10] E. Fischer. The art of uninformed decisions: A primer to property testing. Bulletin of the European Association for Theoretical Computer Science, Vol. 75, pages 97–126, 2001.
- [11] O. Goldreich. A Sample of Samplers A Computational Perspective on Sampling. ECCC, TR97-020, May 1997.
- [12] O. Goldreich. Another motivation for reducing the randomness complexity of algorithms. Position paper, *ECCC*, 2006.
- [13] O. Goldreich, S. Goldwasser, E. Lehman, D. Ron, and A. Samorodnitsky. Testing Monotonicity. Combinatorica, Vol. 20 (3), pages 301–337, 2000.
- [14] O. Goldreich, S. Goldwasser, and D. Ron. Property testing and its connection to learning and approximation. Journal of the ACM, pages 653-750, July 1998.
- [15] O. Goldreich and D. Ron. Property testing in bounded degree graphs. Algorithmica, pages 302–343, 2002.
- [16] O. Goldreich and D. Ron. A sublinear bipartite tester for bounded degree graphs. Combinatorica, Vol. 19 (3), pages 335-373, 1999.
- [17] O. Goldreich and M. Sudan. Locally testable codes and PCPs of almost linear length. Journal of the ACM, Vol. 53 (4), pages 558–655, 2006.

- [18] O. Goldreich and L. Trevisan. Three theorems regarding testing graph properties. Random Structures and Algorithms, Vol. 23 (1), pages 23-57, 2003.
- [19] T. Kaufman and M. Sudan. Algebraic Property Testing: The Role of Invariances. In 40th ACM Symposium on the Theory of Computing, 2008, pages 403–412.
- [20] L. Lovász and N. Young. Lecture Notes on Evasiveness of Graph Properties. Technical Report TR-317-91, Princeton University, Computer Science Department, 1991. Available from http://arxiv.org/abs/cs.CC/0205031
- [21] I. Newman. Private vs. Common Random Bits in Communication Complexity. Information Processing Letters, Vol. 39 (2), pages 67-71, 1991.
- [22] J. Radhakrishnan and A. Ta-Shma. Bounds for Dispersers, Extractors, and Depth-Two Superconcentrators. SIAM Journal on Discrete Mathematics, Vol. 13 (1), pages 2-24, 2000.
- [23] D. Ron. Property testing. In Handbook on Randomization, Volume II, pages 597-649, 2001.
 (Editors: S. Rajasekaran, P.M. Pardalos, J.H. Reif and J.D.P. Rolim.)
- [24] R. Rubinfeld and M. Sudan. Robust characterization of polynomials with applications to program testing. SIAM Journal on Computing, Vol. 25 (2), pages 252-271, 1996.
- [25] R. Shaltiel. Recent Developments in Explicit Constructions of Extractors. Bulletin of the European Association for Theoretical Computer Science, Vol. 77, pages 67–95, 2002.
- [26] R. Shaltiel and C. Umans. Simple Extractors for All Min-Entropies and a New Pseudo-Random Generator. In 32nd IEEE Symposium on Foundations of Computer Science, pages 648–657, 2001.
- [27] O. Sheffet. Reducing the Randomness Complexity of Property Testing, with an Emphasis on Testing Bipartiteness. M.Sc. Thesis, Weizmann Institute of Science, December 2006. Available from http://www.wisdom.weizmann.ac.il/~oded/msc-os.html
- [28] A. Shpilka and A. Wigderson. Derandomizing Homomorphism Testing in General Groups. SIAM Journal on Computing, Vol. 36-4, pages 1215–1230, 2006.
- [29] A. Ta-Shma, D. Zuckerman, and S. Safra. Extractors from Reed-Muller Codes. In 32nd IEEE Symposium on Foundations of Computer Science, pages 638–647, 2001.
- [30] D. Zuckerman. Randomness-Optimal Oblivious Sampling. Journal of Random Structures and Algorithms, Vol. 11, Nr. 4, December 1997, pages 345–367.