# Testing Isomorphism in the Bounded-Degree Graph Model (preliminary version)

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#### Abstract

We consider two versions of the problem of testing graph isomorphism in the bounded-degree graph model: A version in which one graph is fixed, and a version in which the input consists of two graphs. We essentially determine the query complexity of these testing problems in the special case of *n*-vertex graphs with connected components of size at most poly(log n). This is done by showing that these problems are computationally equivalent (up to polylogarithmic factors) to corresponding problems regarding isomorphism between sequences (over a large alphabet). Ignoring the dependence on the proximity parameter, our main results are:

- 1. The query complexity of testing isomorphism to a fixed object (i.e., an *n*-vertex graph or an *n*-long sequence) is  $\tilde{\Theta}(n^{1/2})$ .
- 2. The query complexity of testing isomorphism between two input objects is  $\widetilde{\Theta}(n^{2/3})$ .

Testing isomorphism between two sequences is shown to be related to testing that two distributions are equivalent, and this relation yields reductions in three of the four relevant cases. Failing to reduce the problem of testing the equivalence of two distribution to the problem of testing isomorphism between two sequences, we adapt the proof of the lower bound on the complexity of the first problem to the second problem. This adaptation constitutes the main technical contribution of the current work.

Determining the complexity of testing graph isomorphism (in the bounded-degree graph model), in the general case (i.e., for arbitrary bounded-degree graphs), is left open.

Keywords: Property Testing, Graph Properties, Graph Isomorphism, Sequence Isomorphism,

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## 1 Introduction

We consider the problem of testing graph isomorphism in the bounded-degree graph model (introduced in [11] and reviewed in [9, Chap. 9])). We actually considered two versions of the problem: In one version (called the fixed graph version) the input is a single graph and the task is testing whether this graph is isomorphic to a fixed graph (which "massively parametrized" the property); in the other version, one is given two input graphs and the task is testing whether they are isomorphic.<sup>1</sup> (Both versions of the problem were considered before, but mostly in different testing models (see Section 1.3).)

The bounded-degree graph model. Recall that, in the bounded-degree graph model, graphs are represented by their incidence functions and distances between graphs are measured accordingly. Specifically, for a fixed degree bound d, a graph G = ([n], E) of maximum degree at most d is represented by a function  $g : [n] \times [d] \rightarrow [[n]]$ , where  $[[n]] = \{0, 1, ..., n\} = [n] \cup \{0\}$ , such that  $g(v, i) = u \in [n]$  if u is the *i*<sup>th</sup> neighbor of v (in G), and g(v, i) = 0 if v has less than *i* neighbors.

The graph G = ([n], E) is said to be  $\epsilon$ -far from the graph G' = ([n], E') if the symmetric difference between E and E' is larger than  $\epsilon dn/2$  (equiv., if any representations  $g : [n] \times [d] \rightarrow [[n]]$  and  $g' : [n] \times [d] \rightarrow [[n]]$  of G and G' differ on more than  $\epsilon dn$  entries (i.e.,  $|\{(v,i) : g(v,i) \neq g'(v,i)\}| > \epsilon dn)$ ). Otherwise, the graphs are  $\epsilon$ -close. The graph G = ([n], E) is said to be  $\epsilon$ -far from a graph property  $\Pi$  (i.e., a set of graphs that is closed under isomorphism) if G is  $\epsilon$ -far from any graph in  $\Pi$ .

We say that an oracle machine is an  $\epsilon$ -tester of  $\Pi$  if, when given oracle access to an incidence function of the graph, it distinguishes between the case that the graph is in  $\Pi$  and the case that the graph is  $\epsilon$ -far from  $\Pi$  (i.e., it accepts with probability at least 2/3 in the first case and rejects with probability at least 2/3 in the second case). When  $\epsilon$  is unspecified (e.g., when saying that "testing  $\Pi$ requires Q queries"), it is assumed to be some small positive constant.<sup>2</sup> Indeed, testing isomorphism to a fixed graph H is the task of testing the property that consists of the set of graphs that is isomorphic to H (i.e., H is a massive parameter that specifies the property).<sup>3</sup>

#### 1.1 Testing isomorphism between graphs with small connected components

With the foregoing preliminaries in place, we can state our first main result.

**Theorem 1.1** (testing isomorphism to a fixed graph (in the bounded-degree graph model)): Suppose that H is an n-vertex graph that consists of connected components that are each of size at most poly(log n). Then, the query complexity of  $\epsilon$ -testing isomorphism to H is at most  $\tilde{O}(n^{1/2}/\epsilon^2)$  and at least  $\tilde{\Omega}(n^{1/2})$ , where the upper bound holds for all H's and the lower bound holds for almost all 3-regular H's.

Indeed, we leave open the question of what is the query complexity of testing isomorphism to H, in the general case. The upper bound on the size of the connected components implies that the query complexity is in the same ball-park as the number of connected components that the potential tester visits. A logarithmic lower bound on the size will be used to guarantee that the connected components may be pairwise non-isomorphic (and even pairwise far from being isomorphic, see Lemma 2.2). This fact will be used when lower-bounding the query complexity of testing isomorphism.

Turning to the problem of testing isomorphism between two given graphs, we extend the testing framework to the case in which the potential tester is given a pair of input oracles. Specifically, we say that an oracle machine is an  $\epsilon$ -tester for isomorphism between two graphs if, when given oracle access to

<sup>&</sup>lt;sup>1</sup>The graphs  $G_1 = ([n], E_1)$  and  $G_2 = ([n], E_2)$  are isomorphic if there exists a bijection  $\pi : [n] \to [n]$  (called an isomorphism) such that  $\{\pi(u), \pi(v)\} \in E_2$  if and only if  $\{u, v\} \in E_1$ .

<sup>&</sup>lt;sup>2</sup>That is, saying "testing  $\Pi$  requires Q queries" means that for some  $\epsilon > 0$ , any  $\epsilon$ -tester of  $\Pi$  requires Q queries.

<sup>&</sup>lt;sup>3</sup>See [9, Sec. 12.7.2] for a brief discussion of "massively parametrized" properties.

incidence functions of the two graphs, it distinguishes between the case that the graph are isomorphic and the case that the first graph is  $\epsilon$ -far from any graph that is isomorphic to the second graph.

**Theorem 1.2** (testing isomorphism between two input graphs (in the bounded-degree graph model)): Let  $\Phi$  be the set of *n*-vertex graphs that consist of connected components that are each of size at most poly(log *n*). The query complexity of  $\epsilon$ -testing isomorphism between two graphs that are promised to be in  $\Phi$  is at most  $\widetilde{O}(n^{2/3}/\epsilon^2)$  and at least  $\widetilde{\Omega}(n^{2/3})$ . Equivalently, the query complexity of  $\epsilon$ -testing that two graphs are both in  $\Phi$  and are isomorphism to one another is at most  $\widetilde{O}(n^{2/3}/\epsilon^2)$  and at least  $\widetilde{\Omega}(n^{2/3})$ . Furthermore, the time complexity is also  $\widetilde{O}(n^{2/3}/\epsilon^2)$ .

The stated equivalence is due to the fact that membership in  $\Phi$  can be  $\epsilon$ -tested in time  $O(\epsilon^{-1} \cdot \text{poly}(\log n))$ . Again, we leave open the question of what is the query complexity of testing isomorphism between two graphs, in the general case. We comment that in the context of *one-sided error testing*, even the fixed-graph version requires linear query complexity (see Theorem 2.5).

#### 1.2 Techniques: Testing isomorphism between sequences

Theorems 1.1 and 1.2 are proved by showing the computational equivalence of these two graph-testing problems to corresponding problems of testing isomorphism between sequences (over a large alphabet). We say that two sequences,  $\sigma = (\sigma_1, ..., \sigma_n)$  and  $\tau = (\tau_1, ..., \tau_n)$ , are isomorphic if there exists a bijection  $\pi : [n] \to [n]$  such that  $\tau_{\pi(j)} = \sigma_j$  for every  $j \in [n]$ . One corresponding problem refers to testing isomorphism to a fixed sequence, and the other problem refers to testing isomorphism between two sequences (equiv., two parts of a single sequence), where we say that two *n*-long sequences are  $\epsilon$ -far if they differ on more than  $\epsilon n$  symbols (hence,  $(\sigma_1, ..., \sigma_n)$  and  $(\tau_1, ..., \tau_n)$  are  $\epsilon$ -far from being isomorphic if  $|\{j \in [n] : \sigma_j \neq \tau_{\pi(j)}\}| > \epsilon \cdot n$  for every bijection  $\pi : [n] \to ][n]$ ). Theorems 1.1 and 1.2 follow by presenting reductions between the graph-testing tasks and the corresponding sequence-testing tasks, and establishing the following results regarding the complexity of the sequence-testing tasks.

**Theorem 1.3** (testing isomorphism to a fixed sequence): Fixing any  $\Sigma$  and n, for every  $\sigma \in \Sigma^n$ and  $\epsilon > 0$ , the query complexity of  $\epsilon$ -testing isomorphism to  $\sigma$  is  $O(n^{1/2}/\epsilon^2)$ . On the other hand, if  $|\Sigma| = \Omega(n)$ , then for almost all  $\sigma \in \Sigma^n$ , the query complexity of testing isomorphism to  $\sigma$  is  $\Omega(n^{1/2})$ .

Note that the lower bound requires a large alphabet (i.e.,  $|\Sigma| = \Omega(n)$ ), whereas  $\epsilon$ -testing isomorphism to a fixed sequence over a constant-sized alphabet has query complexity  $O(1/\epsilon^2)$ . The same holds with respect to the following result.

**Theorem 1.4** (testing isomorphism between two input sequences): Fixing any  $\Sigma$  and n, the time complexity of  $\epsilon$ -testing isomorphism between n-long sequences over  $\Sigma$  is  $O(n^{2/3}/\epsilon^2)$ , provided that symbols in  $\Sigma$  can be compared in unit time. On the other hand, if  $|\Sigma| = \Omega(n)$ , then the query complexity of testing isomorphism between n-long sequences over  $\Sigma$  is  $\Omega(n^{2/3})$ .

The proof of the lower bound of Theorem 1.4 is the main technical contribution of this work, but before discussing it we briefly sketch the other three proofs.

On proving Theorem 1.3. Both directions are proved by observing that the sequence-testing problem is computationally equivalent to testing the identity of a given distribution to a fixed distribution, where the distribution-tester is given samples of the tested distribution. The fixed distribution is assigned the value v with probability  $|\{i \in [n] : \sigma_i = v\}|$ , and samples from the tested distribution correspond to the value of the tested sequence at random location. The gap between sampling with repetitions, which is available to the distribution-tester, and sampling without repetitions, which is (w.l.o.g the only thing) available to the sequence-tester, can be ignored when establishing the  $\Omega(n^{1/2})$  lower bound. (Given these reductions, Theorem 1.3 follows from the results surveyed in [9, Sec. 11.2].)<sup>4</sup>

On proving Theorem 1.4. While the upper bound follows easily by reducing the sequence-testing task to the distribution-testing task (very much as in the case of testing isomorphism to a fixed sequence), we failed to find a reduction in the opposite direction. The source of difficulty is the gap between sampling with and without repetitions, where here we cannot ignore this gap because we are considering  $\omega(n^{1/2})$  random samples taken in the probability space [n]. The main technical contribution of this work is overcoming this difficulty.

Instead of reducing the distribution-testing problem to the sequence-testing problem, we adapt Valiant's [19] proof of an  $\Omega(n^{2/3})$  lower bound for the distribution testing problem to the sequencetesting setting. The easy part is showing that it suffices to consider a "canonical" sequence-tester that rules according to the pattern of collisions among the oracle answers, while ignoring both the locations and values of the collisions. (This replaces an analogous statement regarding collisions in the samples given to the distribution-tester.) Next, we show that, for some YES and NO-instances, the pattern of collisions seen by a  $o(n^{2/3})$ -query tester for the sequence problem are statistically close. This is shown by "reducing" the *analysis* of the collision patterns seems by the sequence-tester to those analyzed (for distribution-testing) by Valiant [19].

Specifically, we transform the probability space that underlies the samples that are considered by Valiant (where random samples are drawn with repetitions) to a probability space that fits the sequence-testing setting (where random samples are drawn without repetitions). This transformation retains a tiny fraction of the original probability space and the resulting distribution of the collision patterns is different from the original distribution of the collision patterns. Still, we show that, with high probability, the difference does not occur in places that matter.

More concretely, Valiant [19] (following Batu *et. al.* [3]) considers pairs of probability distributions with heavy and light elements such that in the NO-case the distributions agree on heavy elements but disagree on light elements.<sup>5</sup> He shows that the collision pattern of the light elements (which is significantly different in the two cases (i.e., the cases of YES-instances and NO-instances)) is "masked" by the collision pattern of the heavy elements. Our transformation has an analogous effect: It *does change the collision pattern of heavy elements*, but does so *in an identically manner in the two cases* (and in a way that is oblivious of the light elements); furthermore (with high probability), the transformation *does not affect the collision pattern of light elements at all*. Hence, our transformation does not (significantly) increase the statistical difference between the collision patterns seen (by the potential tester) in the two cases (although in one case the tester has to accept and in the other case it has to reject).

#### 1.3 Related work

The two versions of the graph isomorphism testing problem were considered before [8, 15, 16], but mostly in different models. One exception that we are aware of is the work of Newman and Sohler [16], who considered testing properties of hyperfinite graphs in the bounded-degree graph model. Towards that end, they proved that testing isomorphism between two such graphs has complexity that only depends on the proximity parameter (i.e.,  $\epsilon$ ); see [16, Thm. 3.2].

A few years earlier, Fischer and Matsliah [8] studied the query complexity of both versions of the graph isomophism testing problem in the dense graph model (introduced in [10] and reviewed in [9, Chap. 8]). Interestingly, in all cases they considered, the complexity is sublinear (in the number of vertex-pairs, but polynomially related to that number). In particular, isomorphism between two *n*-vertex input graphs can be tested with one-sided error using  $\tilde{O}(n^{3/2})$  queries, and (two-sided error)

<sup>&</sup>lt;sup>4</sup>For a list of credits, which starts with [12, 4], see [9, Sec. 11.5.1].

<sup>&</sup>lt;sup>5</sup>Needless to say, the YES-instances consists of pairs of identical distributions.

testing of isomorphism to a fixed *n*-vertex graph require  $\widetilde{\Omega}(n^{1/2})$  queries.

More recently, Kusumoto and Yoshida [15] studied these testing problems in the "adjacency list model" (actually, in the general graph model (introduced in [17, 14] and reviewed in [9, Chap. 10])). They considered the case that the graph are promised to be forests (or, alternatively, are required to be forests as part of the property). They showed that in these case the query complexity is polylogarithmic in the size of the graph.

The issue of sampling with versus without repetitions (a.k.a with versus without replacement) arose also in the work of Raskhodnikova *et. al.* [18]. The context there was approximating the number of distinct elements in a sequence, and in that context they presented a reduction of  $O(\alpha)$ -factor approximation based on O(s) samples with repetitions to  $\alpha$ -factor approximation based on s samples without repetitions.

#### 1.4 Organization.

As stated in Section 1.2, our main results are proved by showing the computational equivalence of the graph isomorphism testing problems and the corresponding problems of testing isomorphism between sequences (over a large alphabet). This equivalence is shown in Section 2, and Section 3 focuses on the complexity of the sequence isomorphism testing problems. In particular, the lower bound on the complexity of testing isomorphism between two input sequences is proved in Section 3.2.

## 2 Graph Isomorphism versus Sequence Isomorphism

In this section we show the computational equivalence of the graph isomorphism testing problems to corresponding problems of testing isomorphism between sequences (over a large alphabet). Specifically, we shall focus on (n-vertex) graphs that have small connected components (i.e., each of poly(log n)-size) and on n-long sequences over the alphabet [n]. The fact that the connected components are small will be used when reducing the graph-testing problems to the corresponding sequence-testing problems. Specifically, the overhead of the reduction (presented in Section 2.1) is linearly related to the size of the connected components. When reducing in the opposite direction, we shall use a bijection between [n] and a collection of n 3-regular  $O(\log n)$ -vertex expander graphs that are pairwise far from being isomorphic to one another. (Straightforwards constructions of such a collection of gadgets are spelled out in Section 2.2.) Using this bijection we reduce the sequence isomorphism testing problems to the graph isomorphism testing problems (see Section 2.3). Lastly, in Section 2.4 we fill-up a technical gap (between equal-sized connected compenents and size-bounded connected components) and demonstrate the uselessness of *one-sided error testers* for the various isomorphism problems.

#### 2.1 Reducing the graph problems to the sequence problems

Our aim in reducing the graph problems to the sequence problems is making a step towards obtaining algorithms (i.e., testers) for the graph problems.

Reductions in the context of property testing should preserve sublinear complexities as well as distances between the objects and the corresponding properties. We refrain from presenting here an adequate notion of a reduction, while regretting that the treatment in [9, Sec. 7.4] is not general enough for the current application. Instead, we use the most generic notion possible, which merely asserts that *if one testing problem is solvable within some complexity then the other is solvable within related complexity.* 

**Proposition 2.1** (obtaining testers for the graph problems): Let  $\Phi$  be the set of n-vertex graphs that consist of connected components that are each of size s = s(n), and let d be the degree bound used in the bounded-degree graph model. Then, for every  $\epsilon > 0$ , the following holds.

- 1. If  $\epsilon$ -testing isomorphism to a fixed (n/s)-long sequence has query complexity q, then  $\epsilon$ -testing isomorphism to a fixed n-vertex graph in  $\Phi$  has query complexity  $d \cdot q$ .
- 2. If  $\epsilon$ -testing isomorphism between two (n/s)-long sequences has query complexity q, then  $\epsilon$ -testing isomorphism between two n-vertex graphs that are promised to be in  $\Phi$  has query complexity  $d \cdot q$ . Furthermore, if the time complexity of the first problem is T, then the time complexity of the second problem is  $poly(s) \cdot T$ .

The graph-testing problems refer to the bounded-degree model, and the sequence-testing problems can be restricted to the alphabet [2n/s].

Since testing  $\Phi$  is easy, we can remove the promise and test the property that consists of pairs of isomorphic graphs that are both in  $\Pi$ . On the other hand, we can easily reduce testing *m*-long sequences over [2m] to testing 2m-long sequences over [2m].

**Proof:** The fixed object case follows as a special case of the two-object case, when allowing free oracle access to one of the objects. Focusing on the two-object case, we first assume that the sequences-tester works for sequences over any alphabet.

The basic idea is viewing the connected components of the input graphs as symbols in corresponding sequences, while noting that the locations of the symbols in a sequence are immaterial (for the sequence isomorphism problem), just as the labels of vertices are immaterial for the graph-testing problem. Hence, our graph-tester invokes the guaranteed sequence-tester and answers its queries by finding and describing connected component that have not been used for that purpose before. Specifically, whenever the sequence-tester makes a new query, we select uniformly at random a vertex that was not visited so far, explore the connected component in which it resides, and answer the query with a description of the corresponding graph (either as an unlabeled graph or as a canonically labeled graph with vertex set [s]).<sup>6</sup> We stress that isomorphic copies of the same *s*-vertex graph (potentially appearing as a connected component in the tested graphs) are mapped to the same symbol.

The furthermore clause (of Item 2), referring to the running time of the resulting tester, is based on the fact that canonical labeling of bounded-degree graphs can be found in polynomial-time [2]. That is, the mapping of *s*-vertex connected components to symbols representing the set of all isomorphic copies (of the corresponding graph) can be implemented in poly(n)-time.

The analysis boils down to noting that both problems reduce to testing equality between the number of elements of each type that occur in the tested objects. In the case of graphs, these elements are the connected components of the graph and the types are the isomorphism classes, whereas in the case of sequences the elements are the different locations and the types are the symbols.

Note that the foregoing description refers to an alphabet that consists of all (unlabelled) s-vertex graphs (of maximum degree d). We now show that the sequence testing problem for m-long sequences over arbitrary alphabets reduces to the corresponding problem for the alphabet [2m]. Essentially, we invoke the tester T for the special case and answer its queries by querying our own oracle and maintaining the list of symbols viewed so far. When our oracle answers with a symbol that was not viewed so far, we answer with a random value in [2m] that was not used by us so far, and record the symbol and the selected value. (When our oracle answers with a symbol that was viewed before, we answer with the same element of [2m] that was used at that time.) Hence, when given access to the two m-long sequences over  $\Sigma$  (i.e.,  $\Sigma$  is the set of symbols that actually occur in these sequences), we *effectively* invoke T on corresponding sequences (i.e., the sequence  $(\sigma_1, ..., \sigma_m)$  is mapped to the sequence  $(\psi(\sigma_1), ..., \psi(\sigma_m))$ ).

<sup>&</sup>lt;sup>6</sup>All vertices in this explored connected component are marked as *visited*, and will not be selected when answering subsequent queries.

#### 2.2 A collection of graphs that are pairwise far from being isomorphic

As one may expect, a random collection of  $\exp(\Omega(k))$  3-regular k-vertex graphs will do for our purposes. (In fact, we get an even larger collection, whereas a collection of  $\exp(k^{\Omega(1)})$  gadgets would have sufficed too.)

**Lemma 2.2** For every sufficiently large even  $k \in \mathbb{N}$ , there exists a collection of  $\exp(\Omega(k \log k))$  3regular k-vertex graphs that are pairwise  $\Omega(1)$ -far from being isomoprohic. Furthermore, with overwhelmingly high probability, a random collection of  $\exp(\Omega(k \log k))$  3-regular k-vertex graphs satisfies the property.

Recall that, with probability 1 - o(1), a random 3-regular k-vertex graph is an expander; that is, every set of k' < k/2 vertices neighbors at least  $\Omega(k')$  vertices outside it [7].

**Proof:** Our starting point is Bollobas's estimate for the number of labeled *d*-regular *k*-vertex graphs [6], which is

$$N_d(k) \stackrel{\text{def}}{=} e^{-c-c^2} \cdot \frac{(dk)!}{(dk/2)! \cdot 2^{dk/2} \cdot (d!)^k} \tag{1}$$

where c = (d-1)/2 (and dk is even and  $d = o(\log k)^{1/2}$ ). Using a rough approximation, we have

$$N_d(k) \approx e^{-c-c^2} \cdot \frac{(dk/e)^{dk}}{(dk/2e)^{dk/2} \cdot 2^{dk/2} \cdot (d!)^k}$$
 (2)

$$= e^{-c-c^2} \cdot \frac{(dk/e)^{dk/2}}{(d!)^k}$$
(3)

Hence, for any constant  $d \ge 1$ , we have  $N_d(k) = \Omega(k)^{dk/2}$ .

We prove the existence of the claimed collection by using a greedy algorithm. For some small constant  $\epsilon > 0$ , at each step, we select an arbitrary graph that is not  $\epsilon$ -close to being isomoprophic to any of the graph already selected. The point is that the number of graphs that are  $\epsilon$ -close to being isomoprophic to a fixed k-vertex graph is at most  $M_d(k) \stackrel{\text{def}}{=} k! \cdot \binom{dk}{\epsilon dk} \cdot k^{\epsilon dk} \ll k^{k+(\epsilon+o(1))\cdot dk}$ . Hence, we can select  $N_d(k)/M_d(k)$  graphs, and the claim follows, since  $N_3(k)/M_3(k) \gg k^{k/3}$  for  $\epsilon < 1/18$  and sufficiently large k. In fact, with overwhelmingly high probability, selecting a random set of  $k^{k/4}$  (d-regular k-vertex) graphs will do.

**Remark 2.3** (large set expanders): The fact that, with very high probability, almost all the graphs in the random collection that satisfies Lemma 2.2 are expanders suffices for our main results.<sup>7</sup> We get slightly more appealing results by observing that, with very high probability, a random collection of  $\exp(\Omega(k))$  3-regular k-vertex graphs contains only large set expanders in which the expansion condition holds for every set of  $k' \in [\Omega(k), k/2]$  vertices (rather than for every  $k' \in [n/2]$ ). The point is that a random 3-regular k-vertex graph is a large set expander with probability  $1 - \exp(-\Omega(k))$ , where the hidden constant in the exception probability depends on the constant used in the definition of a large set.<sup>8</sup>

$$B(k) = \sum_{k' \in [\alpha k, 0.5k]} \binom{k}{k'} \cdot N_3(k') \cdot N_3(k-k') \cdot \sum_{k'' \in [[\gamma k']]} \binom{3k'}{3k''} \cdot \binom{3(k-k')}{3k''}$$

<sup>&</sup>lt;sup>7</sup>Specifically, it suffices for all results stated in the introduction with the exception that the lower bound of Theorem 1.1 holds only for graphs H selected from a distribution of extremely high min-entropy rather than for almost all such graphs.

<sup>&</sup>lt;sup>8</sup>Specifically, fixing  $\alpha, \gamma \in (0, 0.5)$ , consider the probability that a 3-regular k-vertex graph contains a set of  $k' \in [\alpha \cdot k, 0.5 \cdot k]$  vertices that neighbors less than  $\gamma \cdot k'$  vertices outside it. A crude upper bound for the number of such graphs is given by

#### 2.3 Reducing the sequence problems to the graph problems

Our aim in reducing the sequence problems to the graph problems is making a step towards obtaining lower bounds on the complexity of the graph-testing problems.

In this case, the basic formalism of [9, Def. 7.13] suffices for capturing the relevant reductions. Loosely speaking, a *q*-local  $(\epsilon, \epsilon')$ -reduction of  $\Pi$  to  $\Pi'$  is a mapping of objects of the first type (i.e., the type of  $\Pi$ ) to objects of the second type (i.e., the type of  $\Pi'$ ) that satisfies the following three conditions:

- 1. Locality (local reconstruction): The value of an object at the image of the reduction at any point is determined by the value of the preimage at q points; that is, if the reduction maps f to f', then the value of f' at any point is determined by the value of f at q points.
- 2. Preservation of the properties: The reduction maps objects in  $\Pi$  to objects in  $\Pi'$ .
- 3. Partial preservation of distance to the properties: An object that is  $\epsilon$ -far from  $\Pi$  is mapped to an object that is  $\epsilon'$ -far from  $\Pi'$ .

It follows that if  $\Pi'$  can be  $\epsilon'$ -tested within query complexity Q', then  $\Pi$  can be  $\epsilon$ -tested within query complexity  $q \cdot Q'$  (see [9, Thm. 7.14]). Indeed, q is the overhead of the reduction.

**Proposition 2.4** (towards deriving lower bounds on the graph-testing problems): Let  $\Phi$  be the set of  $n \cdot s$ -vertex graphs that consist of connected components that are each of size  $s = s(n) \ge \log_2 n$ , and let  $d \ge 3$  be the degree bound used in the bounded-degree graph model. Then, for every  $\epsilon > 0$ , the following holds.

1. Testing isomorphism to a fixed n-long sequence over [n] is 1-locally  $(\epsilon, \Omega(\epsilon))$ -reducible to testing isomorphism to a fixed 3-regular  $s \cdot n$ -vertex graph in  $\Phi$ .

Furthermore, for almost all 3-regular  $s \cdot n$ -vertex graph H in  $\Phi$ , testing isomorphism to the sequence (1, 2..., n) is 1-locally  $(\epsilon, \Omega(\epsilon))$ -reducible to testing isomorphism to H.

2. Testing isomorphism between two n-long sequences over [n] is 1-locally  $(\epsilon, \Omega(\epsilon))$ -reducible to testing isomorphism between two 3-regular  $s \cdot n$ -vertex graphs in  $\Phi$ .

The graph-testing problems refer to the bounded-degree model.

**Proof:** For some fixed constant  $\delta > 0$ , let C be a collection of n graphs, each being a 3-regular s-vertex expander<sup>9</sup>, that are pairwise  $\delta$ -far from being isomorphic to one another. (The existence of such a collection is guaranteed by Lemma 2.2 and [6].) Fixing a bijection  $\psi : [n] \to C$ , we spell out the claimed reduction, while focusing on the case of two input-objects, and viewing the fixed-object case as a special case (in which oracle access to the first object is for free).

$$\ll \sum_{\substack{k' \in [\alpha k, 0.5k]}} \binom{k}{k'} \cdot N_3(k) \cdot \left( (k'/k)^{3k'/2} \cdot ((k-k')/k)^{3(k-k')/2} \right) \cdot \binom{3k}{6\gamma k'}$$

$$\approx \sum_{\substack{k' \in [\alpha k, 0.5k]}} 2^{H_2(k'/k) \cdot k} \cdot N_3(k) \cdot 2^{-H_2(k'/k) \cdot 1.5k} \cdot 2^{H_2(6\gamma k'/3k) \cdot 3k}$$

$$= \sum_{\substack{k' \in [\alpha k, 0.5k]}} 2^{-0.5 \cdot H_2(k'/k) \cdot k + 3 \cdot H_2(2\gamma k'/k) \cdot k} \cdot N_3(k),$$

where  $H_2$  is the binary entropy function. Using a sufficiently small  $\gamma > 0$ , the claim holds (for any  $\alpha \in (0, 0.5]$ ).

<sup>9</sup>Recall that an s-vertex graph is an expander if any set of  $s' \leq s/2$  vertices in it neighbors at least  $\Omega(s')$  vertices outside this set.

The reduction maps each *n*-long sequence over [n], viewed as a function  $\sigma : [n] \to [n]$ , to a graph in  $\Phi$  such that its  $i^{\text{th}}$  connected component is isomorphic to  $\psi(\sigma(i))$ . The vertices of this connected component are labeled  $(i-1)\cdot s+1, ..., i\cdot s$ . Hence, a query for the neighbor of vertex  $(i-1)\cdot s+j$ , where  $i \in [n]$  and  $j \in [s]$ , is answered by querying the  $i^{\text{th}}$  symbol of  $\sigma$ , and determining the corresponding neighbor of the  $j^{\text{th}}$  vertex in  $\psi(\sigma(i))$ . That is, the sequence-tester obtains the value  $\sigma(i)$ , applies the mapping  $\psi$  to this value, considers the canonical labeling of the resulting *s*-vertex graph, and answers according to the neighborhood of the  $j^{\text{th}}$  vertex.

Note that if the input sequences are isomorphic, then the corresponding graphs are isomorphic. On the other hand, as shown next, if the input sequences, denoted  $S_1$  and  $S_2$ , are  $\epsilon$ -far from being isomorphic, then the corresponding graphs, denoted  $G_1$  and  $G_2$ , are  $\Omega(\epsilon)$ -far from being isomorphic.

Suppose that the first graph is  $\epsilon'$ -close to an isomorphic copy of the second graph, and let  $\pi'$ :  $[ns] \to [ns]$  denote the mapping that witnesses this fact (i.e.,  $\pi'(G_1)$  is  $\epsilon'$ -close to  $G_2$ ).<sup>10</sup> Let us assume first, for simplicity, that this "almost-isomorphism"  $\pi'$  maps connected components of  $G_1$  to connected components of  $G_2$ . In such a case, there exists a bijection  $\pi : [n] \to [n]$  such that for at least  $(1 - (\epsilon'/\delta)) \cdot n$  of the  $i \in [n]$  it holds that the  $i^{\text{th}}$  connected component in  $G_1$  is  $\delta$ -close to an isomorphic copy of the  $\pi(i)^{\text{th}}$  connected component in  $G_2$ . (Indeed,  $\pi$  is the mapping of components induced by  $\pi'$ .) Recalling that (1) all the connected components of  $G_1$  and  $G_2$  are in C, and that (2) different graphs in C are  $\delta$ -far from being isomorphic, it follows that the connected components that are  $\delta$ -close to being isomorphic are actually isomorphic copies of the same graph (in C). Thus, for at least  $(1 - (\epsilon'/\delta)) \cdot n$  of the  $i \in [n]$ , it holds that the  $i^{\text{th}}$  connected component in  $G_1$  is isomorphic to the  $\pi(i)^{\text{th}}$  connected component in  $G_2$ . Hence,  $S_1$  is  $(\epsilon'/\delta)$ -close to an isomorphic copy of  $S_2$  (by virtue of the sequence relocation mapping  $\pi$ ).

Recall that the foregoing analysis was based on the simplifying assumption that  $\pi'$  maps connected components of  $G_1$  to connected components of  $G_2$ . This is not necessarily the case, but the fact that the connected components are expanders can be used to show that this is essentially the case.

We say that a connected component of  $G_1$  is effectively preserved by  $\pi'$  if at least  $1-0.5\delta$  of its vertices are mapped by  $\pi'$  to the same connected component of  $G_2$ . Assume, for a moment, that at least a  $1-\gamma\epsilon'$ fraction of the connected components of  $G_1$  are effectively preserved by  $\pi'$ , where the constant  $\gamma$  will be determined later. Then, there exists a bijection  $\pi : [n] \to [n]$  such that for at least  $(1-\gamma\epsilon'-(\epsilon'/0.5\delta))\cdot n$ of the  $i \in [n]$  it holds that the  $i^{\text{th}}$  connected component in  $G_1$  is effectively preserved and is  $\delta$ -close to an isomorphic copy of the  $\pi(i)^{\text{th}}$  connected component in  $G_2$ . (This is the case because otherwise the distance between  $G_1$  and its image under  $\pi'$  is greater than  $(\epsilon'/0.5\delta)) \cdot (\delta - 0.5\delta)$ , where the first factor accounts for the fraction of effectively preserved components that are  $\delta$ -far from the corresponding mapped components, and the second factor accounts for the fraction of mapped vertex-incidences in which these components differ.)<sup>11</sup> So it follows that for at least  $(1 - (\gamma + (2/\delta)) \cdot \epsilon')) \cdot n$  of the  $i \in [n]$ it holds that the  $i^{\text{th}}$  connected component in  $G_1$  is isomorphic to the  $\pi(i)^{\text{th}}$  connected component in  $G_1$ . Hence,  $S_1$  is  $(\gamma + (2/\delta)) \cdot \epsilon'$ -close to an isomorphic copy of  $S_2$  (again, by virtue of the sequence relocation mapping  $\pi$ ).

The foregoing analysis relied on the assumption that at least a  $1 - \gamma \epsilon'$  fraction of the connected

<sup>&</sup>lt;sup>10</sup>Indeed,  $\pi'(G_1)$  denotes the graph obtained from  $G_1$  by relabeling the vertices according to  $\pi'$ ; that is,  $\{\pi'(u), \pi'(v)\}$  is an edge of  $\pi'(G_1)$  if and only if  $\{u, v\}$  is an edge of  $G_1$ .

<sup>&</sup>lt;sup>11</sup>Indeed,  $\pi$  is any bijection that fits the preservation condition of  $\pi'$  (i.e., the connected components of  $G_1$  that are effectively preserved by  $\pi'$  are mapped by  $\pi$  to the corresponding connected components of  $G_2$ ). Let I denote the set of *i*'s such that the *i*<sup>th</sup> connected component in  $G_1$  is effectively preserved and is  $\delta$ -far from an isomorphic copy of the  $\pi(i)^{\text{th}}$  connected component in  $G_2$ . Assuming towards the contradiction that  $|I| > (\epsilon'/0.5\delta) \cdot n$ , recall that for each  $i \in I$  it holds that  $\pi'$  maps at least  $1 - 0.5\delta$  fraction of the vertices of the *i*<sup>th</sup> component of  $G_1$  to the  $\pi(i)^{\text{th}}$  component of  $G_2$ . Hence, for each  $i \in I$ , the incidences of vertices in the *i*<sup>th</sup> component differ from the incidences of their image under  $\pi'$  in at least  $\delta \cdot 3s - 3 \cdot 0.5\delta \cdot s$  entries (i.e., the number of incidence differences between the components minus the number of incidences that belong to vertices of the *i*<sup>th</sup> component that were not mapped to the  $\pi(i)^{\text{th}}$  component). It follows that, under  $\pi'$  witnesses a distance of at most  $\epsilon'$ .

components of  $G_1$  are effectively preserved by the witness mapping  $\pi'$ . However, this assumption is actually a fact (i.e., it must hold), because otherwise expansion (w.r.t the non-preserved components) implies that  $\pi'(G_1)$  is  $\gamma \epsilon' \cdot \Omega(\delta)$ -far from  $G_2$ , in contradiction to the hypothesis regarding  $\pi'$  (provided the constant  $\gamma$  is chosen to be sufficiently large). This establishes the main claims of the proposition and leaves us with the furthermore claim of Part 1, which refers to the case that the fixed graph is uniformly selected among the 3-regular graphs in  $\Phi$  (i.e., that the fixed graph is uniformly selected among all 3-regular graphs that have *n* connected components that is each of size *s*).

Recall that the furthermore claim refers to reducing from the problem of testing isomorphism to the fixed sequence (1, 2, ..., n), whereas the instance produced by reduction refers to a fixed  $n \cdot s$ -vertex graph H that consists of the connected components  $\psi(1), ..., \psi(n)$ . Recalling that  $\psi : [n] \to C$  is a bijection, it follows that a random H corresponds to a random collection C. Recall that such a random collection contains n - o(n) expanders, whereas we have assumed that all graphs in C are expanders.<sup>12</sup> However, a closer look at the foregoing argument reveals that it holds even if all graphs in C are only large set extractors (in the sense defined in Remark 2.3). Since the latter condition holds with very high probability, the furthermore claim follows.

#### 2.4 Implications for testing graph isomorphism

Using the first (resp., second) part of Propositions 2.1 and 2.4, the claims of Theorem 1.1 (resp., Theorem 1.2) almost follow from Theorem 1.3 (resp., Theorem 1.4). The remaining gap is in the upper bound, since Proposition 2.1 only handled graphs in which all connected components are of the *same size* (whereas the upper bounds in Theorems 1.1 and 1.2 refer to *n*-vertex graphs in which all connected components are of *size at most* poly( $\log n$ )).

The gap can be bridged by modifying the reduction used in the proof of Proposition 2.1 so that *n*-vertex graphs (with connected components of size at most *s*) are associated with *n*-long sequences (initially over  $[\exp(\tilde{O}(s))])^{13}$  such that a connected component of size  $s' \leq s$  is associated with s'locations in the sequence. The reduction itself is modified too: Whenever the sequence-tester makes a new query, we answer it by selecting at random a vertex that was not *selected* before (rather than not *visited* before), exploring the connected component in which it resides, and returning the corresponding graph (either as an unlabeled graph or as a canonically labeled graph with vertex set [s]).

**One-sided error testing.** Recall that a one-sided error tester is required to *always* accept any object that has the property (i.e., accept with probability 1 any such object), while rejecting with high probability (i.e., with probability at least 2/3) any object that is far from the property, as a usual tester. It is quite easy to see that all testing problems considered in this work have no ose-sided error tester of sublinear query complexity. We prove this assertion for the problem of testing isomorphism to a fixed graph (with very small connected components).

**Theorem 2.5** (a lower bound on one-sided testers in the fixed-graph model): There exists a n-graph H of degree bound two and connected components of size three such that, in the bounded-degree graph model, any one-sided error tester of isomorphism to H requires  $\Omega(n)$  queries.

**Proof:** Let H be composed of n/6 isolated triangles and n/6 isolated 2-paths, and assume (w.l.o.g.) that the tester for isomorphism decides based on the number of isolated triangles and 2-paths that it

<sup>&</sup>lt;sup>12</sup>Note that, with probability 1/poly(s), a random 3-regular s-vertex graph is not even connected, whereas we have  $s = \text{poly}(\log n)$ .

<sup>&</sup>lt;sup>13</sup>Recall that we latter reduce the problems regarding m-long sequences over arbitrary alphabet to the corresponding problems regarding 2m-long sequences over [2m].

sees.<sup>14</sup> Then, a one-sided error tester for isomorphism to H that makes at most n/6 queries must always accept when seeing any proportion of isolated triangles and 2-paths, since such a proportion may occur when querying a random isomorphic copy of H. It follows that such a tester (always) accepts when inspecting a random graph composed on n/9 isolated triangles and 2n/9 isolated 2-paths, whereas such a graph is  $\Omega(1)$ -far from the property.

## 3 On the Complexity of Testing Isomorphism between Sequences

In this section we focus on the two versions of the sequence-testing problem, and establish Theorems 1.3 and 1.4. We shall often view *n*-long sequences over  $\Sigma$  as functions from [n] to  $\Sigma$ .

#### 3.1 Testing isomorphism to a fixed sequence

We start with the problem of testing isomorphism to a fixed sequence, which serves as a good warm-up towards our study of the complexity of testing isomorphism between two input sequences.

**Theorem 3.1** (the query complexity of testing isomorphism to a fixed sequence (Theorem 1.3, restated)):

- 1. For every  $\epsilon > 0$ , the query complexity of  $\epsilon$ -testing isomorphism to any fixed n-long sequence is  $O(n^{1/2}/\epsilon^2)$ . Furthermore, the time complexity is the same if the tester can determine the number of occurrences of a symbol in the fixed sequence in constant time.
- 2. Testing isomorphism of an n-long sequence over [n] to the sequence (1, 2, ..., n) requires  $\Omega(n^{1/2})$  queries. The same holds with respect to almost all n-long sequences over [n].

Of course, testing isomorphism to a fixed sequence may be easier in some cases (e.g., the all-1 sequence). In fact, the following reduction that establishes the upper bound yields such results via the results of [20] (see also [5]).

**Proof:** The upper bound follows by a reduction to testing identity to a corresponding fixed (*n*-grained) distribution (where a distribution is called *n*-grained if all elements in its support appear with probability that is a multiple of 1/n). Specifically, given a fixed sequence  $\sigma : [n] \to \Sigma$  and access to an input sequence  $\tau : [n] \to \Sigma$ , we define the fixed distribution D and the input distribution X such that  $\Pr[D=v] = |\{i \in [n] : \sigma(i)=v\}|/n$  and  $\Pr[X=v] = |\{i \in [n] : \tau(i)=v\}|/n$  for every  $v \in \Sigma$ . We then invoke the guaranteed distribution-tester (for the distribution D which is determined by  $\sigma$ ), and provide it with samples of X in the obvious matter; that is, if we need to provide s samples, we select uniformly and independently  $i_1, ..., i_s \in [n]$ , and use the values  $\tau(i_1), ..., \tau(i_s)$ . (The straightforward analysis of this reduction is given in the first paragraph of the proof of Theorem 3.2.)

Using the known distribution-testers (see [9, Thm. 11.11]), the claimed upper bound follows. The time bound (asserted in the furthermore clause) follows by observing that the reduction of *testing identity to a fixed n-grained distribution* to *testing that a distribution is uniform on* [n] only requires an evaluation oracle to the fixed distribution, and makes one query to this oracle per each example obtained from the input distribution (see [9, Sec. 11.2.2.1]).

The lower bound regarding testing isomorphism to the fixed sequence (1, 2, ..., n) is proved by using a reduction in the opposite direction; that is, by reducing testing that a distribution is uniform over [n] to testing isomorphism to the sequence (1, 2, ..., n). Here we capitalized on the fact that we are proving a lower bound of the form  $\Omega(n^{1/2})$ , since in that regime the difference between sampling indices

 $<sup>^{14}</sup>$ This is analogous to a claim in [13]. Specifically, if the tester is guaranteed that the input is composed of connected components that are each of size 3, then we may assume that it merely samples random vertices and inspects their respective connected components.

in [n] without repetitions and sampling them with repetitions can be ignored. This is relevant since a q-query sequence-tester can be assumed to query the input sequence at q random locations, but the set of q locations is distributed uniformly among all q-subsets of [n] (i.e., it samples without repetitions). In contrast, the distribution-tester obtains samples of some n-grained distribution X, which may be thought of as being generated by selecting uniformly  $i \in [n]$  and outputting a value G(i) for a suitable  $G : [n] \to [n]$ , but these samples are generated independently of one another (i.e., they are generated with repetitions). Details follow.

We reduce testing whether an n-grained distribution X over [n] is uniform to testing isomorphism to the sequence  $\sigma = (1, 2, ..., n)$ . The point is that the input distribution X can be viewed as generated by selecting at random  $i \in [n]$  and outputting the value G(i) for an adequate  $G : [n] \to [n]$ ; that is,  $\Pr[X = e] = |\{i \in [n] : G(i) = e\}|/n$ . So testing whether X is uniform over [n] corresponds to testing whether G viewed as a sequence is isomorphic to  $\sigma$ . (Recall that this distribution-testing problem has complexity  $q = \Omega(\sqrt{n})$ , even when restricted to n-grained distributions.) In the reduction, given a tester T for isomorphism to the fixed sequence  $\sigma$ , we construct a distribution-tester that invokes T and answers its (distinct, w.l.o.g.) queries by using the samples provided to it. That is, for every  $j \in [q]$ , the  $j^{\text{th}}$  query is answered by the  $j^{\text{th}}$  sample.

In the analysis, we think of the q samples given to our tester as being generated by selecting  $i_1, ..., i_q \in [n]$  uniformly and independently (with repetitions), and being presented with  $G(i_1), ..., G(i_q)$ . In contrast, without loss of generality, we can think of T as selecting uniformly a sequence of q distinct elements in [n] and querying its oracle for their value.<sup>15</sup> That is, whereas T should be given the answers  $G(i_1), ..., G(i_q)$  such that  $i_1, ..., i_q$  are selected uniformly in [n] without repetitions, we gave it corresponding answers with respect to  $i_1, ..., i_q$  that are selected uniformly in [n] with repetitions. However, the statistical difference between these two n-long samples is at most  $\binom{q}{2}/n$ , and we can ignore this difference when establishing a lower bound of the form  $q = \Omega(\sqrt{n})$ .

#### 3.2 Testing isomorphism between two input sequences

We now turn to the problem of testing isomorphism between two input sequences, while adopting slightly different notation than the one used so far. The first three paragraph recap ideas that were already presented in Section 3.1.

Given two sequences  $S_1, S_2 \in \Sigma^n$ , presented as functions  $S_1, S_2 : [n] \to \Sigma$ , the sequence isomorphism testing problem is to determine whether there exists a permutation  $\pi : [n] \to [n]$  such that  $S_1(j) = S_2(\pi(j))$  for every  $j \in [n]$  (i.e.,  $S_1 = S_2 \circ \pi$ ) or  $S_1$  is far from  $S_2 \circ \pi$  for every permutation  $\pi$ . This can be captured as a property testing problem by considering  $S : \{1,2\} \times [n] \to \Sigma$  such that  $S(i,j) = S_i(j)$ .

Clearly, the sequence isomorphism testing problem is reducible the testing equality between distributions, by considering random variables  $X_1$  and  $X_2$  such that  $\Pr[X_i = \sigma] = |\{j \in [n] : S_i(j) = \sigma\}|/n$  for every  $\sigma \in \Sigma$ . Hence, the complexity of testing sequence isomorphism is upper-bounded by the complexity of testing equality between distributions, which is  $O(1/\epsilon^2) \cdot n^{2/3}$  (see [9, Sec. 11.3], presenting the best result known, which in turn improves over earlier work of Batu *et. al.* [3]).

It is tempting to hope that a reduction in the opposite direction holds if we restrict the distributions to be *n*-grained [9, Def. 11.7], where a distribution is *n*-grained if each element in its support is assigned a probability mass that is a multiple of 1/n. Indeed, one can show that, without loss of generality, a tester of sequence isomorphism queries the sequences at a random set of location, but the distribution-tester obtains a sample that corresponds to a random multi-set of locations. That is, we face a gap between sampling with and without repetitions, and this gap matters because we are interested in the case that the number of samples is larger than the square root of the support size.

<sup>&</sup>lt;sup>15</sup>Given an arbitrary tester T that tests  $\tau : [n] \to [n]$ , suppose that we answer its  $j^{\text{th}}$  query with  $\tau(i_j)$ , where  $(i_1, ..., i_q)$  is uniformly distributed among all q-long sequences of *distinct* elements in [n]. Then, we actually emulate an execution of T in which it is given access to a random isomorphic copy of  $\tau$  (i.e., to the function  $\tau \circ \pi$ , where  $\pi : [n] \to [n]$  is a uniformly distributed bijection).

We were unable to show a reduction of the distribution-testing problem to the sequence-testing problem, which would have allowed to infer a lower bound on the sequence-testing problem from a lower bound on the distribution-testing problem; but we were able to adapt the proof of the known  $\Omega(n^{2/3})$  lower bound (of Valiant [19]) for the distribution-testing problem to thesequence-testing problem.

**Theorem 3.2** (on the complexity of sequence isomorphism (Theorem 1.4, restated)):

- 1. For every  $\epsilon > 0$ , the time complexity of  $\epsilon$ -testing sequence isomorphism (for n-long sequences) is  $O(n^{2/3}/\epsilon^2)$ , provided that symbols can be compared in unit time.
- 2. Testing sequence isomorphism for n-long sequences over [n] requires  $\Omega(n^{2/3})$  queries.

**Proof:** The upper bound follows by the reduction outlined above; that is, given sequences  $S_1, S_2$ :  $[n] \to \Sigma$ , consider the random variables  $X_1$  and  $X_2$  such that  $\Pr[X_i = \sigma] = |\{j \in [n] : S_i(j) = \sigma\}|/n$ . If  $S_1$  is isomorphic to  $S_2$ , then  $X_1 \equiv X_2$ . On the other hand, if  $S_1$  is  $\epsilon$ -far from being isomorphic to  $S_2$ (i.e.,  $\min_{\pi \in \text{Sym}_n}\{|\{j \in [n] : S_1(j) \neq S_2(\pi(j))\}|\} > \epsilon \cdot n$ ), then  $\sum_{\sigma} |\#_{\sigma}(S_1) - \#_{\sigma}(S_2)| > 2 \cdot \epsilon n$ , where  $\#_{\sigma}(S)$  denotes the number occurrences of  $\sigma$  in the sequence S (i.e.,  $\#_{\sigma}(S) = |\{j \in [n] : S(j) = \sigma\}|$ ).<sup>16</sup> Hence,  $\epsilon$ -testing of sequence isomorphism reduces to  $\epsilon$ -testing of identity of distributions, by virtue of invoking the distribution-tester and providing it with samples of  $X_1$  (resp.,  $X_2$ ) by querying  $S_1$ (resp.,  $S_2$ ) at random locations. Using the distribution-tester that works in time  $O(1/\epsilon^2) \cdot n^{2/3}$  (see [9, Sec. 11.3]), the theorem's upper bound follows.

Turning to the lower bound, we adapt Valiant's proof [19] of the lower bound for the distributiontesting problem into a lower bound for the sequence-testing problem. We first mimic the relatively easy argument showing that all that matters is the distribution of "histograms" seen by the tester. Then, we show that the relevant histograms (of some YES and NO-instances) seen by a  $o(n^{2/3})$ -query tester for the sequence problem are statistically close. This is done by "reducing" the *analysis* to the case analyzed by Valiant [19]. The histograms that we define next refer to a pair of sequences; these sequences are not the input sequences (or distributions) but rather samples that the tester obtains from these sequences (or samples). (Indeed, this generalizes the more basic notion of a histogram of a single sequence.)<sup>17</sup>

**Definition 3.2.1** (the relevant histograms): For a pair of m-long sequences  $((s_1, ..., s_m), (s'_1, ..., s'_m)) \in \Sigma^{m+m}$ , the corresponding histogram is an (m+1)-by-(m+1) matrix  $H = (h_{t,t'})_{t,t' \in [[m]]}$  such that  $h_{t,t'}$  is the number of values that occur exactly t times in the first sequence and t' times in the second sequence; that is,

$$h_{t,t'} = \left| \left\{ \sigma \in \Sigma : \ \#_{\sigma}(s_1, ..., s_m) = t \& \ \#_{\sigma}(s'_1, ..., s'_m) = t' \right\} \right|, \tag{4}$$

where 
$$\#_{\sigma}(\sigma_1, ..., \sigma_m) = |\{j \in [m] : \sigma_j = \sigma\}|.$$
 (5)

(Indeed, we use the notation  $[[m]] \stackrel{\text{def}}{=} \{0, 1, ..., m\}.$ )

Note that  $\sum_{t,t'} h_{t,t'} = |\Sigma|$ , since each  $\sigma \in \Sigma$  contributes to exactly one  $h_{t,t'}$ , and  $\sum_{t,t'} h_{t,t'} \cdot (t+t') = 2m$ , since each location  $j \in [m+m]$  (resp., occurrence of a symbol) is counted once in the sum.

Claim 3.2.2 (histograms are all that matters): If isomorphism of n-long sequences over  $\Sigma$  can be tested within query complexity  $q = q(n, \Sigma, \epsilon)$ , then it can be tested by a canonical tester that obtains the values of each of the two sequences in q random positions and rules according to the corresponding histogram. In other words, when testing the sequences  $S_1, S_2 : [n] \to \Sigma$ , the tester rules according to the

<sup>&</sup>lt;sup>16</sup>It may be easier to see that the distance of  $S_1$  from being isomorphic to  $S_2$  equals  $\sum_{\sigma} \max(0, \#_{\sigma}(S_1) - \#_{\sigma}(S_2))$ .

<sup>&</sup>lt;sup>17</sup>A histrogram of a sequence  $\overline{s} = (s_1, ..., s_m)$  is a sequence  $(h_0, h_1, ..., h_m)$  such that  $h_t$  equals the number of elements that occur exactly t times in  $\overline{s}$  (i.e.,  $h_t = |\{\sigma \in \Sigma : \#_{\sigma}(\overline{s}) = t\}|$ , where  $\#_{\sigma}(s_1, ..., s_m) = |\{j \in [m] : s_j = \sigma\}|$ ).

histogram of  $((S_1(j_1), ..., S_1(j_q)), (S_2(k_1), ..., S_2(k_q)))$ , where  $(j_1, ..., j_q)$  and  $(k_1, ..., k_q)$  are distributed uniformly and independently in the set of q-long sequences of distinct elements in [n].<sup>18</sup>

Proof: For sake of clarity, we proceed in two steps (where the first step details an argument already used before (see Footnote 15)). Given an arbitrary tester T as in the hypothesis, and fixing  $(n, \Sigma, \epsilon)$ and  $q = q(n, \Sigma, \epsilon)$ , we first construct an algorithm T' that obtains, as input, a pair of q-long sequences, denoted  $((s_1, ..., s_q), (s'_1, ..., s'_q))$ , invokes T, while answering its  $j^{\text{th}}$  query to the first (resp., second) sequence with  $s_j$  (resp.,  $s'_j$ ), and outputs the verdict of T. When analysing T', we consider, for any two sequences  $S_1, S_2 : [n] \to \Sigma$ , what happens when selecting  $(j_1, ..., j_q)$  and  $(k_1, ..., k_q)$  uniformly and independently among all possible q-long sequences of distinct elements in [n], and feeding T' with  $((S_1(j_1), ..., S_1(j_q)), (S_2(k_1), ..., S_2(k_q)))$ . In this case the output of T' is distributed identically to the output of T when given oracle access to random isomorphic copies of  $S_1$  and  $S_2$  (i.e., to the oracles  $S_1 \circ \pi_1$  and  $S_2 \circ \pi_2$ , where  $\pi_1$  and  $\pi_2$  are uniformly and independently distributed permutations of [n]). Hence, T' distinguishes between the case that  $S_1$  is isomorphic to  $S_2$  and the case that  $S_1$  is  $\epsilon$ -far from being isomorphic to  $S_2$ .

Next, we present the desired canonical tester, denoted T''. On input a (valid) historam, denoted  $H = (h_{t,t'})_{t,t'\in[[q]]}$ , algorithm T'' selects at random a pair of q-long sequences that fits the histogram H, feeds it to T', and outputs its verdict. That is, T'' selects uniformly at random a pair of q-long sequences  $((s_1, ..., s_q), (s'_1, ..., s'_q)) \in \Sigma^{q+q}$  such that

$$|\{\sigma \in \Sigma : \#_{\sigma}(s_1, ..., s_q) = t \& \#_{\sigma}(s'_1, ..., s'_q) = t'\}| = h_{t,t'}$$

holds for every  $t, t' \in [[q]]$ , and outputs  $T'((s_1, ..., s_q), (s'_1, ..., s'_q))$ . Then, for any two sequences  $S_1, S_2 : [n] \to \Sigma$ , the output of T'' when given a *histogram* of a sample of q distinct random values in  $S_1$  and q distinct random values in  $S_2$  equals the output of T' when given the corresponding samples (*themselves*!) from  $\psi(S_1)$  and  $\psi(S_2)$ , where  $\psi$  is a random permutation of  $\Sigma$ . Hence, T'' distinguishes between the case that  $S_1$  is isomorphic to  $S_2$  and the case that  $S_1$  is  $\epsilon$ -far from being isomorphic to  $S_2$ .

(Indeed, our argument only relies on the fact that T distinguishes between the case that  $\psi \circ S_1 \circ \pi_1$  is isomorphic to  $\psi \circ S_2 \circ \pi_1$  and the case that  $\psi \circ S_1 \circ \pi_1$  is  $\epsilon$ -far from being isomorphic to  $\psi \circ S_2 \circ \pi_1$ , where  $\pi_1, \pi_2 : [n] \to [n]$  and  $\psi : \Sigma \to \Sigma$  are random permutations (as above).)

The core of the proof. We wish to show that a canoncal sequence-tester must make  $\Omega(n^{2/3})$  queries in order to distinguish between the case that it is provided with a histograph that corresponds to a YES-instance and the case that it is provided with a histograph that corresponds to a NO-instance. The YES-instance will consists two copies of the sequence  $S_1$ , whereas the NO-instance will consist of the sequences  $S_1$  and  $S_2$ , where  $S_1$  and  $S_2$  will correspond to the *n*-grained distributions analyzed by Valiant [19], which we will denote  $X_1$  and  $X_2$ .

Each of the distributions used in Valiant's proof [19] has n/4 elements of individual probability weight 2/n (called light), and  $n^{2/3}$  elements each of weight  $n^{-2/3}/2$  (called heavy). Furthermore, the proof refers to NO-instances that are pairs of distributions that agree on the heavy elements, but are disjoint on their light elements. We deconstruct these *n*-grained distributions by detailing the underlying probability space [n], and the way this space is mapped to values, which are either heavy or light. The mapping goes through sets, denoted H and L, which correspond to the heavy and light elements.

Fixing two disjoint sets H and L such that |H| = n<sup>2/3</sup> and |L| = n/4, we consider a mapping of the probability space [n] to these sets such that each element in H is assigned n<sup>1/3</sup>/2 elements and each element in L is assigned two elements. We denote this mapping by G : [n] → H ∪ L; that is, for each h ∈ H (resp., l ∈ L) it holds that Pr<sub>j∈[n]</sub>[G(j)=h] = n<sup>-2/3</sup>/2 (resp., Pr<sub>j∈[n]</sub>[G(j)=l] = 2/n).

<sup>&</sup>lt;sup>18</sup>That is, the set  $\{(i_1, ..., i_q) \in [n]^q : |\{i_1, ..., i_q\}| = q\}\}.$ 

• Fixing a set L' of size |L| that is disjoint of  $H \cup L$ , we consider a bijection  $\phi : H \cup L \to H \cup L'$ that is invariant on H (i.e.,  $\phi(H) = H$  and  $\phi(L) = L'$ ). We shall consider the random variables (or distributions)  $X_1$  and  $X_2$  such that  $X_1(j) = G(j)$  and  $X_2(j) = \phi(G(j))$ .<sup>19</sup>

(We shall later view  $X_1$  and  $X_2$  as values sampled from corresponding *n*-long sequences  $S_1$  and  $S_2$ , where  $S_1(j) = G(j)$  and  $S_2(j) = \phi(G(j))$ . Note that  $X_i$  represents the value of a uniformly distributed location in  $S_i$ , and that the statistical distance between  $X_1$  and  $X_2$  equals the relative Hamming distance between  $S_1$  and  $S_2$ , which in turn equals 1/2 (since  $S_1(j) \neq S_2(j)$  if and only if  $G(j) \in L$ , and  $|\{j \in [n] : G(j) \in L\}| = n/2$ ).

• For a fixed  $m = \Theta(n^{2/3})$ , we denote by Y a pair of m-long sequences, each consisting of m independent samples of  $X_1$ ; that is, using  $[n]^{2m}$  as an undelying probability space, we have

$$Y(j_1, ..., j_m, k_1, ..., k_m) = ((X_1(j_1), ..., X_1(j_m)), (X_1(k_1), ..., X_1(k_m))).$$
(6)

(Indeed, here  $j_1, ..., j_m$  are distributed independently in [m], and ditto for  $k_1, ..., k_m$ . This is the setting that is suitable for distribution-testing, and it was analyzed in [19]. Our challenge would be to move from this setting to the one in which  $j_1, ..., j_m$  are distict and ditto for  $k_1, ..., k_m$  (as is suitable for sequence-testing).)

Likewise, we denote by Z a pair of m-long sequences such that the first sequence consists of m independent samples of  $X_1$  and the second sequence consists of m independent samples of  $X_2$ ; that is,

$$Z(j_1, ..., j_m, k_1, ..., k_m) = ((X_1(j_1), ..., X_1(j_m)), (X_2(k_1), ..., X_2(k_m))).$$
(7)

Hence, Y and Z differ only in the values assigned to light samples that occur in the second m-long sequence; that is, the  $m + \ell^{\text{th}}$  element of  $Y(j_1, ..., j_m, k_1, ..., k_m)$  differs from the  $m + \ell^{\text{th}}$  element of  $Z(j_1, ..., j_m, k_1, ..., k_m)$  if and only if  $G(k_\ell) \in L$ .

• The histograms of  $Y = Y(j_1, ..., j_m, k_1, ..., k_m)$  and  $Z = Z(j_1, ..., j_m, k_1, ..., k_m)$  are denoted h(Y)and h(Z), respectively. Recall that the  $(t, t')^{\text{th}}$  entry in h(Y) (resp., h(Z)) is the number of values  $\sigma$  that occur t times in  $((X_1(j_1), ..., X_1(j_m))$  and t' times in  $(X_1(k_1), ..., X_1(k_m))$  (resp., in  $(X_2(k_1), ..., X_2(k_m)))$ .

The crucial fact, proved in [19], is that, for a sufficiently small  $m = \Omega(n^{2/3})$ , the (random variables representing the) histograms h(Y) and h(Z) are statistically close (say, are at total variation distance at most 0.1).

Our aim is to show that the foregoing fact continues to hold when the probability space (underlying these random variables) is restricted to pairs of *m*-long sequences of *distict elements*. Note that this restriction retains only a tiny portion of the original probability space, since  $m = \Omega(n^{2/3}) \gg O(n^{1/2})$ ; nevertheless, we shall show that the corresponding histographs remain statistically close. In other words, we modify the distributions Y and Z so that they fit the samples viewed by the sequence-tester. Recall that the underlying probability space for Y (and likewise for Z) is  $[n]^{2m}$ , whereas here we wish the underlying space to include only pairs of *m*-long sequences of distinct elements in [n]. We shall extensively use the assumption that  $m = c \cdot n^{2/3}$ , for a sufficiently small constant c > 0.

Starting from two uniformly and identically distributed sequences  $(j_1, ..., j_m), (k_1, ..., k_m) \in [n]^m$ , we first observe that, with probability at least 0.99, there are no three-way collisions in  $(j_1, ..., j_m)$ . Furthermore, with probability at least 0.99, there are at most  $2 \cdot {m \choose 2} \cdot (1/n) < m^2/n \ll n^{1/3}$  pairwise collisions in  $(j_1, ..., j_m)$ . Ditto for  $(k_1, ..., k_m)$ . Conditioned on the foregoing case, for each collision

<sup>&</sup>lt;sup>19</sup>Following Batu *et. al.* [3], Valiant [19] uses  $H = [n^{2/3}]$ , L = [0.5n + 1, ..., 0.75n], and L' = [0.75n + 1, ..., n]. This choice is, of course, immaterial. Also note that, for simplicity, we use the elements of the intermediate sets H and L also as final value (as reflected in the fact that  $\phi$  is only applied in the generation of  $X_2$  and that  $\phi(H) = H$ ).

 $\{j_p, j_q\}$  (such that  $j_p = j_q$ ), we re-select at random (i.e., re-randomize) one of the colliding indices (where the index to be re-randomized is selected obliviously of the sequence  $(k_1, ..., k_m)$ ). Note that the number of *potential* new collisions (between the re-randomized indices and all other indices) is upper-bounded by  $\frac{m^2}{n} \cdot m \ll n$ . Hence, with probability 0.99, the re-randomization yields an *m*long sequence of distinct elements (which is uniformly distributed among all such sequences). We do the same for the sequence  $(k_1, ..., k_m)$ . Let us denote the resulting pair of *m*-long sequences by  $((j'_1, ..., j'_m), (k'_1, ..., k'_m))$ .

Recall that in the likely case in which there are no three-way collisions in  $(j_1, ..., j_m)$  (resp.,  $(k_1, ..., k_m)$ ), with high probability, the resulting  $(j'_1, ..., j'_m)$  (resp.,  $(k'_1, ..., k'_m)$ ) is uniformly distributed among all *m*-long sequence of distinct elements in [n], and in any case  $(j'_1, ..., j'_m)$  and  $(k'_1, ..., k'_m)$  are distributed independently of one another. Hence, with high probability, the indices  $((j'_1, ..., j'_m), (k'_1, ..., k'_m))$  are distributed as expected by the sequence-tester. The crucial fact is that the re-randomization of indices does not change the statistical difference between the histograms of the modified Y and Z by much, because such a change occurs only due to collisions of indices in  $G^{-1}(L)$  whereas such collisions are rare.<sup>20</sup> The foregoing fact will be proved next, when using  $\Delta(.,.)$  to denote the statistical difference (a.k.a. total variation difference) between distributions.

**Claim 3.2.3** (the effect of re-randomization): For uniformly distributed  $(j_1, ..., j_m, k_1, ..., k_m) \in [n]^{2m}$ and  $(j'_1, ..., j'_m, k'_1, ..., k'_m)$  as generated above, consider the random variables

- $Y = Y(j_1, ..., j_m, k_1, ..., k_m),$
- $Z = Z(j_1, ..., j_m, k_1, ..., k_m),$
- $Y' = Y(j'_1, ..., j'_m, k'_1, ..., k'_m)$ , and
- $Z' = Z(j'_1, ..., j'_m, k'_1, ..., k'_m).$

Suppose that  $m = c \cdot n^{2/3}$ , for a sufficiently small constant c > 0 (c = 1/20 will do). Then,  $\Delta(\mathbf{h}(Y'), \mathbf{h}(Z')) \leq \Delta(\mathbf{h}(Y), \mathbf{h}(Z)) + 0.05$ .

Recalling that, for a sufficiently small constant c > 0, it holds that  $\Delta(h(Y), h(Z)) < 0.1$  (cf. [19]), we get  $\Delta(h(Y'), h(Z')) < 0.15$ .

Proof: The key observation is that h(Y) and h(Z) (and likewise h(Y') and h(Z')) reflect the pattern of collisions among  $X_i$ -values, where Y (resp., Y') contains only  $X_1$ -values and Z (resp., Z) contains also  $X_2$ -values in its second part. Hence, only collisions between the  $X_1$ -values of indices of the first part (i.e.,  $j_1, ..., j_m$ ) and the  $X_i$ -values of indices of the second part (i.e.,  $k_1, ..., k_m$ ) contribute to  $\Delta(h(Y), h(Z))$ .<sup>21</sup> Furthermore, such a contribution (which arises from the difference between  $X_2$  and  $X_1$ ) occurs only for indices in  $G^{-1}(L)$ , since indices in  $G^{-1}(H)$  are always assign the value of  $X_1$ . The same consideration applies to  $j'_1, ..., j'_m$  and  $k'_1, ..., k'_m$  regarding their contribution to  $\Delta(h(Y'), h(Z'))$ . Hence, the difference between  $\Delta(h(Y), h(Z))$  and  $\Delta(h(Y'), h(Z'))$  is due to re-randomization of indices  $j_p$  (resp.,  $k_p$ ) that reside in  $G^{-1}(L)$  either initially or after re-randomization, and to the collision of their  $X_i$ -value with the  $X_i$ -value of some  $k_q$  or  $k'_q$  (resp.,  $j_q$  or  $j'_q$ ). Details follow.

Recalling that, with probability at least 0.98, there are no three-way collisions in  $(j_1, ..., j_m)$  and at most  $m^2/n$  pairwise collisions in it, we consider the re-randomization applied to one index in each pair  $(j_p, j_q) \in [m^2]$  such that  $j_p = j_q$  (and  $p \neq q$ ). The same analysis is applied to  $(k_1, ..., k_m)$ . Suppose that we re-randomized  $j_p$ , replacing it by a uniformly distributed  $j'_p$ . We consider four cases.

<sup>&</sup>lt;sup>20</sup>Since less than  $m^2/n$  indices get re-randomized, and collisions may occur only with the other O(m) indices, we get a total of  $O(m^3/n)$  collisions each occurring with probability 2/n (because these are collisions of indices in  $G^{-1}(L)$ ). This yields a total difference of  $O(m^3/n^2) = O(c^3)$ , which can be made small enough by a suitable choice of c > 0.

<sup>&</sup>lt;sup>21</sup>We stress that collisions inside each part do not contribute to  $\Delta(h(Y), h(Z))$ , regardless if they are in the first part (i.e., between  $j_1, ..., j_m$ ) or in the second part (i.e., between  $k_1, ..., k_m$ ).

1. If  $G(j_p) \in H$  and  $G(j'_p) \in H$ , then the replacing of  $j_p$  by  $j'_p$  does not affect the difference between  $\Delta(\mathfrak{h}(Y), \mathfrak{h}(Z))$  and  $\Delta(\mathfrak{h}(Y'), \mathfrak{h}(Z'))$ , because these statistical differences are only due to light indices.

(Recall that  $X_2(k) = X_1(k)$  if  $G(k) \in H$ , whereas  $Y = (X_1(j_1), ..., X_1(j_m), X_1(k_1), ..., X_1(k_m))$ and  $Z = (X_1(j_1), ..., X_1(j_m), X_2(k_1), ..., X_2(k_m))$ . Therefore replacing  $j_p \in G^{-1}(H)$  by  $j'_p \in G^{-1}(H)$  may change the pattern of  $X_1$ -value collisions within the Y sequence, but the same change will occur in the Z sequence (since  $G(j_p)$  and  $G(j'_p)$  have values in H). In other words, h(Y') may differ from h(Y) due to the replacement of  $j_p$  by  $j'_p$ , but exactly the same effect occurs between h(Z') and h(Z), because in all cases we refer to collisions of  $X_1$ -values.)

- 2. If  $G(j_p) \in H$  and  $G(j'_p) \in L$ , then the probability that  $G(j'_p)$  hits  $\{G(k_q) : q \in [m]\}$  (equivalently, hits  $\{G(k_q) : q \in [m] \& G(k_q) \in L\}$ )<sup>22</sup> is at most  $m \cdot \frac{2}{n} = 2m/n$ , and otherwise this re-randomization does not affect the difference between  $\Delta(h(Y), h(Z))$  and  $\Delta(h(Y'), h(Z'))$ . (That is, if  $G(j_p) \in H$  and  $G(j'_p) \in L$  do not hit  $\{G(k_q) : q \in [m] \& G(k_q) \in L\}$ , then the replacement of  $j_p$  by  $j'_p$  does not affect the foregoing statistical difference.)<sup>23</sup> Hence, recalling that there are at most  $m^2/n$  collisions among the  $j_p$ -indices (i.e., pairs  $(j_p, j_q)$  such that  $j_p = j_q$  and  $p \neq q$ ), it follows that the total contribution of this case is at most  $\frac{m^2}{n} \cdot \frac{2m}{n} = \frac{2m^3}{n^2} < 0.001$ , by an appropriate choice of the constant c > 0 (in  $m = c \cdot n^{2/3}$ ).
- 3. If  $G(j_p) \in L$  and  $G(j'_p) \in H$ , then the probability that  $G(j_p)$  hits  $\{G(k_q) : q \in [m]\}$  is at most 2m/n, and otherwise this re-randomization does not affect the difference between  $\Delta(\mathfrak{h}(Y), \mathfrak{h}(Z))$  and  $\Delta(\mathfrak{h}(Y'), \mathfrak{h}(Z'))$ . So, again, the total contribution of this case is at most  $2m^3/n^2 < 0.001$ .
- 4. If  $G(j_p) \in L$  and  $G(j'_p) \in L$ , then the probability that either  $G(j_p)$  or  $G(j'_p)$  hits  $\{G(k_q) : q \in [m]\}$  is at most  $2 \cdot 2m/n$ , and otherwise this re-randomization does not affect the difference between  $\Delta(\mathfrak{h}(Y), \mathfrak{h}(Z))$  and  $\Delta(\mathfrak{h}(Y'), \mathfrak{h}(Z'))$ . So the total contribution of this case is at most  $4m^3/n^2 < 0.002$ .

To summarize: The only contribution of the re-randomization of the indices that form collisions in  $(j_1, ..., j_m)$  to the difference between  $\Delta(h(Y), h(Z))$  and  $\Delta(h(Y'), h(Z'))$  arises from indices  $j_p$  that are re-randomized to  $j'_p$  such that either  $j_p$  or  $j'_p$  hits  $\{G(k_q) : q \in [m] \& G(k_q) \in L\}$ . But the probability of this event is small (i.e., smaller than  $O(m^3/n^2) < 0.005$ ), since the number of re-randomized indices is relatively small (i.e., smaller than  $m^2/n$ ) and the probability of each hit is small (i.e., at most 2m/n). Applying the an analogous analysis to  $(k_1, ..., k_m)$ , where here we consider collisions with either  $\{G(j_q) : q \in [m] \& G(j_q) \in L\}$  or  $\{G(j'_q) : q \in [m] \& G(j'_q) \in L\}$ , the claim follows (because we were considering an event that occurs with probability at least 0.96 and showed that in that case the difference is smaller than  $2 \cdot 0.005$ ).

Conclusion. The theorem follows by combining Claims 3.2.2 and 3.2.3, while recalling that (with high probability) the sequences  $(j'_1, ..., j'_m)$  and  $(k'_1, ..., k'_m)$  are uniformly and independently distributed among the *m*-long sequences of distinct elements in [*n*]. Specifically, with high probability, the canonical tester is presented with histograms (of either Y' or Z') that are statistically close (i.e.,  $\Delta(h(Y'), h(Z')) < 0.15$ ), and so it cannot distinguish them. On the other hand, Y' represents answers to *m* distinct random queries made to each of the two copies of the sequence  $S_1$ , whereas Z' represents answers to *m* distinct random queries made to the sequences  $S_1$  and  $S_2$ , which are at distance 1/2 of one another.<sup>24</sup> That is,

<sup>&</sup>lt;sup>22</sup>Indeed, for  $G(j'_p) \in L$  to hit  $\{G(k_q) : q \in [m]\}$ , it must hit  $\{G(k_q) : q \in [m] \& G(k_q) \in L\}$ , which has cardinality at most m. This fact implies the probability bound of  $m \cdot \frac{2}{n}$ .

<sup>&</sup>lt;sup>23</sup>In particular, a possible collision of  $G(j'_p)$  with a value in  $\{G(j'_q) : q \in [m]\}$  does not contribute to  $\Delta(h(Y'), h(Z'))$ . The same holds, of course, for the original collision of  $G(j_p)$  with  $G(j_q)$ .

<sup>&</sup>lt;sup>24</sup>Recall that  $S_1$  and  $S_2$  are *n*-long sequences (viewed as functions defined over [n]) such that  $S_1(j) = G(j)$  and  $S_2(j) = \phi(G(j))$  for every  $j \in [n]$ , whereas  $X_1$  and  $X_2$  are defined in the same manner but viewed as random variables

Y' (resp., Z') represents answers from a pair of sequences that should be accepted (resp., rejected) by a tester with probability at least 2/3. It follows that for a sufficiently small constant c > 0, isomorphism between sequences cannot be tested using  $m = c \cdot n^{2/3}$  queries.

# 4 Concluding Comments

The results in this paper determine the query complexity of both version of the problem of testing graph isomorphism, in the bounded-degree graph model, up to a factor that is linear in the size of the largest connected components. In fact, Theorems 1.1 and 1.2 are special cases of the following general results (which are stated only for constant  $\epsilon > 0$  and  $s > \log_2 n$ ):<sup>25</sup>

- 1. For every sufficiently small constant  $\epsilon > 0$ , the query complexity of  $\epsilon$ -testing isomorphism to a fixed *n*-vertex graph that consists of connected components of size at most *s* is between  $O(s \cdot n)^{1/2}$  and  $\Omega(n/s)^{1/2}$ .
- 2. For every sufficiently small constant  $\epsilon > 0$ , the query complexity of  $\epsilon$ -testing isomorphism between a pair of *n*-vertex graphs that consist of connected components of size at most *s* is between  $O(s^{1/3} \cdot n^{2/3})$  and  $\Omega(n/s)^{2/3}$ .

The proofs of these results employ graph theoretic arguments (i.e., the notion of an expander) only in order to reduce the analysis to problems that ignore the graph theoretic origin. We believe that determining the query complexity of testing graph isomorphism in the general case (e.g., for connected graphs, let alone for expanders) may require some graph theoretic insights. Two concrete challenges follow.

**Open Problem 4.1** (sublinear complexity for testing isomorphism to a fixed graph): Is it possible to 0.01-test isomorphism to a fixed 3-regular graph in query complexity that is sublinear in the number of vertices?

Note that such a tester must have two-sided error probability (cf. Theorem 2.5).

**Open Problem 4.2** (higher lower bounds for testing isomorphism between two input graphs): Does 0.01-testing isomorphism between two n-vertex graphs (of bounded degree) require  $\omega(n^{2/3})$  queries?

As hinted above, the focus should be on graphs that cannot be particitoned into small parts with few edges among them. Recall that  $\epsilon$ -testing isomorphism between hyperfinite graphs (i.e., graphs that can be partitioned to  $P(\epsilon)$ -sized parts with at most  $\epsilon dn/6$  edges among them) is doable using  $Q(\epsilon)$  queries [16, Thm. 3.2], where  $P, Q: (0, 1] \to \mathbb{N}$ .

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 $\Pr[X_1 \neq X_2] = \Pr_{j \in [n]}[S_1(j) \neq S_2(j)] = \Pr_{j \in [n]}[G(j) \in L] = 1/2.$ 

<sup>25</sup>Actually, the following bounds hold also for  $s > \frac{3 \log_2 n}{\log_2 \log_2 n}$ , but for smaller s the query complexity is exponential in  $\Theta(s \log s)$ .

over [n]. Also recall that

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