

Mathematical Logic and Probability

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Preface

In the late 19th and early 20th centuries, logic and probability were frequently treated as closely related disciplines. Each has, in an important sense, gone its own way, so that neither, in its modern form, is in any proper sense a systematization of the “Laws of Thought,” as Boole called them.

However, the last four decades have seen a remarkable rapprochement. On the most obvious level, the various probability logics have developed as formal systems of reasoning in the modern sense of logic.

At a deeper level, though, attempts have been made to formulate logics in which model theory of random variables, stochastic processes, and randomized structures can be explored from the perspective of model theory. Continuous first-order logic as a context for stability theory on metric structures is perhaps the most conspicuous example, but others exist.

At the same time, algorithmic randomness in its various forms has come to play a core role in computability theory, while probabilistic computation of various kinds (randomized computation, interactive proofs, and others) has come to dominate major parts of computational complexity. The older recursion-theoretic program of machine learning, initiated by Gold in the 1960s, has become much more important thanks to Valiant’s reformulation in probabilistic terms to allow for reasonable errors.

The model theory of random objects, Fraïssé limits, and pseudofinite structures, each of which embodies some important aspect of 0-1 laws, has been important for longer, but advances in stability, simplicity, and the transition from finite to infinite model theory have enriched this subject.

In set theory, too, the study of dynamics that respect probability measures has played a central role in the study of equivalence relations. Probability is frequently at the center of modern descriptive set theory.

Nor have these developments been independent. The PAC learning theory of Valiant is inextricably linked to the model theory of NIP theories. The dynamics of computable Polish spaces have become an important emerging area in computability. Randomized computation is the natural computation on metric structures. Notions of random structures have become intertwined with algorithmic randomness, and are naturally described in continuous first order logic.

Many of these developments have been adequately treated in isolation by various books. Probability logic has been discussed at length from various perspectives in [10, 239, 242, 394, 419]. Bayesian networks are well-covered, for instance, in [219, 397, 398], and a monograph on adapted distributions also exists [180]. Randomized computation has a detailed treatment in [30]. Algorithmic randomness is the subject of three relatively recent books, [158, 387, 194]. Zero-one laws are treated at length in [165, 230], and other places, and [266] includes an

extended treatment of Fraïssé limits. Random graphs are extensively covered in [74, 125, 338]. The definitive reference on PAC learning is [297]. In the field of set-theoretic dynamics, there have been several treatments at several levels of detail, among which [55, 260, 298, 303] merit special mention. There is no shortage of book-length treatments of subjects within the range of this book.

However, a reader in a well-stocked library might well pass all these separate books without knowing that they had anything substantive in common. Indeed, one could read most of them in detail — in addition to the long papers that give strong expositions on many related subjects (the seminal paper [61] on continuous first-order logic comes to mind) — without finding a commonality.

It is true that [238] describes connections between probability logics and Bayesian networks. However, it is silent on the rest of these issues.

The present book, then, attempts to take a unified — or, at least, unifying — approach to this subject. The expanding literature in each of these fields has seen more interaction between them, so that a model theorist might well want to know more about the frontier of probabilistic work in set theory, or a computability theorist more about the relevant work in model theory.

We focus here on *mathematical* logic and probability. Probability logic and its relatives seem frequently to arise as works of *philosophical* logic, and this has implications for the questions that are asked about it. Frequently it is seen in connection with the theory of rational decision, as in [242]. Mathematical logic, by contrast, asks about computability and undecidability; about theories and their models; about reducibilities and regularity of sets. Alternate logics are of interest to mathematical logic inasmuch as they provide the necessary infrastructure for carrying out this program in interesting settings. Applications of logic to artificial intelligence and other modeling contexts are important, but they arise as applications of the theory, not as its defining elements.

Chapter 1 begins to lay out the central thesis of the book: that all the other chapters have something to say to one another. This is done by identifying several important cross-cutting themes that come up in several of the other chapters.

In the next chapter, we begin the technical section of the book by describing the various logics useful for probability. Continuous first-order logic has a central role, not least because it generalizes many others. Probability logic is extensively studied, and is explored here as well, as are some other approaches.

In a third chapter, we will consider the theory of algorithmic randomness, with special attention to normal numbers, Martin-Löf randomness, and their relation to computation. This treatment will not be complete, of course — the subject is well-covered elsewhere. Rather, the focus will be on those aspects of algorithmic randomness that interact with other areas of advance in the logic and probability community.

The chapter on randomized computation involves the leap of reasoning that computability and complexity still have something to say to one another. Recent work on generic and coarse computability, as well as that on derandomization, descriptive complexity, and continuous first-order logic support this hypothesis.

The following two chapters will take up the various approaches to random structures. The investigation of random structures seems to have arisen historically from the study of random graphs, which invited generalization to 0-1 laws, and which

connected with the earlier beginnings of Fraïssé limits. More recent approaches consider the “random” structure as a single structure that somehow embodies the possible variation — graphons, Keisler randomizations, invariant random subgroups, and the like. Others use algorithmic randomness to define the structure.

In taking up the problem of learning theory, there is a fair viewpoint from which learning after the tradition of Gold, probably approximately correct (PAC) learning after the tradition of Valiant, and the model theory of NIP structures are wildly different fields. The chapter devoted to these topics takes the opposite view. Valiant’s definition is a natural extension of Gold’s framework, and the theory of Vapnik-Chervonenkis dimension governs both PAC learning and NIP theories.

The final chapter surveys the general area of dynamics. An introduction to orbit equivalence relations and Borel cardinality is given, and several topics on the relation of measure to equivalence relations are considered, including the implications of ergodicity and Hjorth’s notion of turbulence. Recent model-theoretic approaches to Szemerédi Regularity and Furstenberg Correspondence belong here, too, as does the characterization of 1-randomness by the Ergodic Theorem and the emerging theory of computable Polish spaces.

Of course, some limits must be set on the content of such a book. For instance, a new line of thought has arisen in recent years over categorical treatments of probability [199, 200, 395]. In view of traditional [350] and recent [119, 248, 247] work on connections between category theory and logic, this work is certainly interesting and relevant, but it is hard, at this stage of the theory, to explain its relationship to the other work.

The book is to be formally self-contained, but realistically anticipates a reader who has completed a first course in logic at the graduate or upper undergraduate level. Such a reader will, after reading the book, be prepared to understand the frontier of the research literature in probability-related areas of computability, model theory, set theory, and logical aspects of artificial intelligence. There is an important place in the world for a reader equipped in this way: A major part of logic in the coming years will involve connections between these fields, and those who understand something of all of them will be well-poised to contribute.

CHAPTER 4

Nondeterminism and Randomized Computation

4.1. Nondeterminism

4.1.1. Nondeterministic Machines. One of the greatest insights of complexity theory is that of designing an algorithm that proceeds according to a distribution of behavior. This insight is, at first face, obviously impractical. Only on further reflection does it become so practical as to be the industry standard in many applications.

The most naive approach to this paradigm, perhaps, is to allow a machine to do almost anything, as long as we have an efficient algorithm to verify success. To be concrete, we give the following definition.

DEFINITION 4.1.1. Let $S \subseteq \{0, 1\}^*$, and $f : \mathbb{R} \rightarrow \mathbb{R}$. We say that S is of class **NTIME**(f) if there is some Turing machine T such that on input $x \in \{0, 1\}^*$, the machine T halts in at most $f(|x|)$ steps, returning output from $\{0, 1\}$, with the additional provision that $x \in S$ if and only if there is some $\sigma \in 2^{<\omega}$ such that $T(x, \sigma) = 1$.

We call T the *verifier* for S , and we call σ the *proof* that $x \in S$. The idea is that $x \in S$ if and only if there is a proof of this fact, and the verifier T checks whether the proof is, in fact, a “correct” proof. Note, of course, that because of the bounded running time of the verifier, we have a natural bound of $f(|x|)$ on the length of σ , without needing to explicitly bound the length of the proof. Since the verifier can inspect at most one bit of σ at each time step, if there is no σ shorter than that on which $T(x, \sigma) = 1$, then there is none at all.

This definition is not transparently one of randomized computation, but of deterministic verification. Consequently, we also give the following definition.

DEFINITION 4.1.2. Let $S \subseteq \{0, 1\}^*$ and $f : \mathbb{R} \rightarrow \mathbb{R}$. We say that S is of class **NTIME**^{comp}(f) if there is some oracle Turing machine T such that on input $x \in \{0, 1\}^*$, the machine T halts in at most $f(|x|)$ steps, returning output from $\{0, 1\}$, with the additional provision that $x \in S$ if and only if there is some $\sigma \in \{0, 1\}^{<\omega}$ such that $T^\sigma(x) = 1$.

We will show that these two classes are equal. However, our provisional definition of **NTIME**^{comp}(f) displays the “random” character more clearly: The oracle is viewed as a set of coin flips, and the machine “accepts” or “rejects” x , possibly in light of the values of those coin flips. The condition for membership in **NTIME**^{comp}(f) is exactly that there is some sequence of coin flips on which the machine accepts. The randomness in this oracle is exactly the “nondeterminism” of nondeterministic computation.

PROPOSITION 4.1.3. *The class **NTIME**^{comp}(f) is equal to the class **NTIME**(f).*

PROOF. Let S be of class $\mathbf{NTIME}(f)$, with verifier T . Then we define an oracle machine \bar{T} so that $\bar{T}^\sigma(x) = T(x, \sigma)$ for all $x \in \{0, 1\}^*$, with equal running time. Now \bar{T} witnesses that S is of class $\mathbf{NTIME}^{\text{comp}}(f)$. The converse is exactly symmetric. \square

Interesting as the questions of nondeterminism may be at the level of complexity (frequently runtime, occasionally space or some other resource), much of the distinction disappears entirely at the level of computability. Since the proof σ has bounded length, a Turing machine with adequately large time bounds could scan all possible proofs, and report whether it accepts any of them. As we will see in a later section, something of the distinction reemerges in more generalized models of computation.

4.1.2. NP and the Polynomial Hierarchy. The nondeterministic model of computation is fundamentally not one of probability, except in the most naive sense. Membership in the class is determined by the existence or nonexistence of a single element of the space of random coins — in the limit, a measure zero set.

On the other hand, this model does give rise to one of the most storied problems of complexity theory, and this problem is, as we shall see, closely entangled with problems on other models of computation which are more genuinely probabilistic. We state this problem now.

DEFINITION 4.1.4. Let $S \subseteq \{0, 1\}^*$.

- (1) We say that S is of class \mathbf{P} if there is some Turing machine T and some polynomial p such that on input $x \in \{0, 1\}^*$, the machine T halts in at most $p(|x|)$ steps, returning output from $\{0, 1\}$, where $T(x) = 1$ if and only if $x \in S$.
- (2) We say that S is of class \mathbf{NP} if S is of class $\mathbf{NTIME}(p)$ for some polynomial p .

PROBLEM 4.1.5. Is $\mathbf{P} = \mathbf{NP}$?

The literature on this problem is extensive, but the problem is still so open that Gasarch [210] has polled computer scientists to find their opinions on not only the answer, but also whether and when a solution will be found (in fairness, a majority feel that a solution is possible and negative). Since the subject of nondeterministic computation is, in its own right outside the proper scope of the present book, we will not even attempt a cursory review of the literature, although the interested reader will do well to consult [30].

Just as the Continuum Hypothesis ($2^{\aleph_0} = \aleph_1$), when it resisted solution, gave rise to the much stronger Generalized Continuum Hypothesis ($2^{\aleph_\alpha} = \aleph_{\alpha+1}$), which at first seemed no more obviously true or false, the \mathbf{P} versus \mathbf{NP} problem has a more general analogue, the collapse of the Polynomial Hierarchy.

DEFINITION 4.1.6. We define the Polynomial Hierarchy, Σ_i^p , Π_i^p , and Δ_i^p , as follows:

- (1) $\Sigma_1^p = \mathbf{NP}$
- (2) A set S is of class Π_n^p if and only if its complement is of class Σ_n^p
- (3) A set $S \subseteq \{0, 1\}^*$ is of class Σ_{n+1}^p if and only if there is a Π_n^p set $Q(\tau, y)$ and for any $x \in \{0, 1\}^*$ there is a $\sigma \in \{0, 1\}^{q(|x|)}$ such that $x \in S$ if and only if $Q(\sigma, x)$.

$$(4) \Delta_n^p = \Sigma_1^p \cap \Pi_1^p.$$

By contrast with the arithmetical hierarchy, in which we know that every possible inclusion is strict, relatively little is known about the strictness of this hierarchy. It is not even obvious that $\Delta_1^p = \mathbf{P}$. There are, for each i , problems that are Σ_i^p complete. Moreover, we can give some criteria for collapse.

PROPOSITION 4.1.7. *If $\mathbf{P} = \mathbf{NP}$, then $\Sigma_i^p = \Sigma_{i+1}^p$ for all i .*

PROOF. We proceed by induction on i . For $i = 1$, since $\Sigma_1^p = \mathbf{NP}$, then under the stated hypothesis we have $\Sigma_1^p = \mathbf{P} = \Pi_1^p$. Consequently, $\Sigma_2^p = \mathbf{NP} = \Sigma_1^p$. Similarly, by induction, $\Sigma_{i+1}^p = \mathbf{NP} = \Sigma_i^p$. \square

This is, *a priori*, not the only way in which the hierarchy could collapse. It is, for instance, possible that $\Sigma_i^p \subsetneq \Sigma_{i+1}^p$ exactly for $i < k$ for some k . In that case, we say that the Polynomial Hierarchy collapses at level k . The uncertainty concerning whether the hierarchy collapses at some point, and, if so, where, enhances the contrast of this structure with the arithmetical hierarchy — for instance, the arithmetical hierarchy can have no complete set precisely because the hierarchy is strict: if an “arithmetical hierarchy complete” set occurred in Σ_i^0 , it could not be complete, as it would not admit reduction from a properly Σ_{i+1}^0 set. In the Polynomial Hierarchy, though, that could just be the level at which the hierarchy collapses.

Really, the better analogy is with the analytic hierarchy — although the same differences remain: the analytic hierarchy is known to be strict in a very strong sense, and little is known unconditionally about the structure of the polynomial hierarchy. In essence, the step to pass from Π_n^p to Σ_{n+1}^p is very much like a function quantifier, an issue to be explored more fully in the next section.

4.1.3. Descriptive Complexity. One of the most powerful insights of computability theory is the connection between degrees of unsolvability on one hand and logics on the other. In an appropriate context, the sets defined by Σ_1^0 formulas are precisely the computably enumerable ones, the sets defined by computable infinitary formulas are exactly those computable from (perhaps transfinitely) iterated Turing jumps, and those defined by Π_1^1 formulas exactly those computable from Kleene’s \mathcal{O} .

This connection continues at the level of complexity, as well. We consider structures in the *language of strings*, which consists of one binary predicate that is axiomatically determined to be a linear ordering, and one unary predicate X , which will be interpreted as the set of bits with a 1. A finite structure in this language is thus interpreted as a binary string, of length equal to its cardinality.

The standard reference in this area is Immerman’s survey [275], and the treatment in this section is indebted to his. The following theorem, first proved by Fagin in his 1973 Ph.D. thesis [178], is fundamental in the area.

THEOREM 4.1.8. *Let $S \subseteq 2^{<\omega}$. Then S is of class \mathbf{NP} if and only if S is axiomatizable in the language of strings by a finitary Σ_1^1 formula.*

PROOF. Let T be an oracle Turing machine running in time n^k witnessing that $S \in \mathbf{NP}$. We can arrange (if necessary, by slowing T) that T uses exactly one random bit at each time step. Let $\Gamma = (\gamma_i : i \leq g)$ be a list of all pairs (q, σ) , where q is either a state of T or a new symbol $-$; and σ is from the tape alphabet of T .

We let $(\mathfrak{C}_{s,t} : s, t \leq n^k)$ be an array in which $\mathfrak{C}_{s,t}$ is a pair $(q, \sigma) \in \Gamma$ so that σ is the contents of cell s at time t , and such that if T is on cell s at time t , then q is the state of the machine at that time, and otherwise $q = -$. For each $i \leq g$, we define a predicate $C_i(s, t)$ to denote the pairs s, t such that $\mathfrak{C}_{s,t} = \gamma_i$.

Let Δ be the string of random bits used by T in the first n^k steps. We then construct a formula as follows: We let $\alpha(\mathfrak{C}, \tau)$ say that $\mathfrak{C}_{s,0}$ codes the input string τ , and let $\beta(\mathfrak{C})$ say that $C_i(s, t) \rightarrow \neg C_j(s, t)$ for $i \neq j$ (that is, that at each time, a particular cell may have only one value). We let $\xi(\mathfrak{C})$ state that $(\mathfrak{C}_{s,n^k} : s \leq n^k)$ includes the accept state. Finally, we write a sentence $\eta(\mathfrak{C}, \Delta)$ stating that for all t , the stage $\mathfrak{C}_{s,t+1}$ is the next step prescribed by T after $\mathfrak{C}_{s,t}$ on random bit $\Delta(t)$. Now let

$$\varphi(\mathfrak{C}, \Delta, \tau) = \alpha(\mathfrak{C}, \tau) \wedge \beta(\mathfrak{C}) \wedge \xi(\mathfrak{C}) \wedge \eta(\mathfrak{C}, \Delta).$$

Now $\tau \in S$ if and only if $\exists(\mathfrak{C}, \Delta) \varphi(\mathfrak{C}, \Delta, \tau)$.

On the other hand, any finitary Σ_1^1 formula axiomatizing S can be expressed as $\exists f \psi(f, \tau)$, where ψ has no second-order quantifiers. Now a Turing machine can verify in polynomial time that a particular sequence f and input τ satisfy $\psi(f, \tau)$, so that S is of class **NP**. \square

This result generalizes throughout the polynomial hierarchy, as seems to have been first recorded by Stockmeyer [467].

THEOREM 4.1.9. *Let $S \subseteq 2^{<\omega}$. Then S is of class Σ_n^p (respectively, Π_n^p) if and only if S is axiomatizable in the language of strings by a finitary Σ_n^1 (respectively, Π_n^1) formula.*

PROOF. Notice that Π_n^p sets are exactly the complements of Σ_n^p sets, just as Π_n^1 sets are the complements of Σ_n^1 sets. Consequently, it suffices to prove the result for Σ_n^p . We have already established the result for Σ_1^p .

Suppose that the Σ_n^p (respectively, Π_n^p) sets are exactly the Σ_n^1 (respectively, Π_n^1) sets. Then the result $\Sigma_n^1 \subseteq \Sigma_n^p$ follows exactly as in the original case. For the opposite inclusion, the inductive definition of the polynomial hierarchy admits a straightforward translation from a Σ_n^p definition to a Σ_n^1 definition. \square

4.2. Randomized Turing Machines

4.2.1. Complexity Classes Defined by Randomization. We now turn to a more properly probabilistic notion of computation. It is natural enough to ask that a computation give the correct answer with some high probability, but it is important to ask in what domain the probability lives. One can (and we do, in Section 4.5) carry out a satisfactory theory in which we consider a probability measure on the space of inputs to the computation, and consider the probability that a randomly selected input is one on which the computation is correct. The larger body of work, though, has been in a different direction.

Consider the Miller-Rabin primality test, first put forward in [374, 410]. To determine whether n is prime, we perform several tests. In each test, we choose a random positive integer $a < n$, and use it as a test for whether n is composite. To do this, we compute $x = a^{n-1} \bmod n$. If $x \neq 1$, then n cannot be prime by Fermat's Little Theorem. In the course of the calculation, we might also discover a non-trivial square root of 1 modulo n , in which case we again know that n is composite (a corollary to Fermat's Little Theorem). We perform these tests (with new random

choices of a) as often as we like (in practice, there is a formula that will give the number of tests necessary to give the desired level of certainty of correctness). If these checks reveal that n is composite, we report as much. Otherwise, we report it to be prime.

To those familiar with computably enumerable (or perhaps Π_1^0) sets, the issue here is clear: we are giving output of both positive (prime) and negative (composite) kinds, based only on negative information. Of course, if we report that n is composite, it is based on good evidence, and what we reported was correct. If we report that n is prime, it is possible that we just got unlucky with the choice of the a 's in all of the tests. Under appropriate (and very strong) number-theoretic assumptions (a strengthening of the Riemann Hypothesis), such unluckiness is impossible, and the algorithm is simply correct for determining whether n is prime. Without venturing for the wild fancies of number theory quite this far, if n truly is composite, there are still quite a few of the possible values for a that must be witnesses, and the probability of not finding any of them in several attempts is quite small (indeed, the probability is 2^{-t} , where t is the number of tests performed; the interested reader may find a full exposition and analysis of the algorithm in [138]). While a polynomial-time algorithm for primality testing is now known, methods like the Miller-Rabin test are still the industrial standard for most applications, because their correctness is high enough for industrial needs, and they run faster.

The one part of the previous algorithm that is not obviously “algorithmic” is the clause, “choose a random positive integer $a < n$.” Indeed, depending on what, exactly, we mean by it, this operation is not likely to be strictly computable. As we saw in Chapter 3, an important foundation of algorithmic randomness is that if we have an algorithm to compute something, it cannot be very random. Consequently, a broader definition is needed. The following definition, introduced in [146], formalizes the notion of a Turing machine that can make some random choices, and then produce an output that is correct with high probability in the domain of the random choices.

DEFINITION 4.2.1. A *randomized Turing machine* (sometimes called a *probabilistic Turing machine*) is a Turing machine equipped with an oracle for an element of 2^ω , called the *random bits*.

We can then talk reasonably about a randomized Turing machine M having output k on input n with probability p if the Lebesgue probability measure of the set of random bit strings x such that $M^x(n) = k$ is equal to p .

The addition of the random bits presents an abundance of new questions. For instance, we classically ask about whether a Turing machine halts. We can now ask, for instance about the probability of halting. We consider a Turing functional $T : 2^{<\omega} \rightarrow 2^{<\omega}$, in the sense that T is a Turing machine which, on oracle σ will output the sequence $T(\sigma)$. We call such a functional *prefix-free* if, whenever $T(\sigma_1) \downarrow$ and $T(\sigma_2) \downarrow$, then σ_1 is not an initial segment of σ_2 . A *universal prefix-free machine* is a Turing functional U such that for any prefix-free machine T there is a string $\nu_T \in 2^{<\omega}$ such that $T(\sigma) = U(\nu_T \sigma)$ for all $\sigma \in 2^{<\omega}$.

Following Chaitin [112], we define Ω_U , for a fixed universal prefix-free machine U , to be probability (in the sense of the standard probability measure on $2^{<\omega}$) of the set of σ such that $U(\sigma) \downarrow$. This is often described as the probability that a random (prefix-free) Turing machine halts on a random input. To see this interpretation, interpret U as a randomized Turing machine whose output, for a fixed oracle, is

either undefined or constant (over all inputs). We then interpret the random bits as giving both the U -code for an algorithm (in the first several bits) and the input (whose arity may be an arbitrary finite number). In this interpretation, all U -codes of the same length are given equal probability, and, given a U -code, all inputs (of the correct length) have equal probability.

THEOREM 4.2.2 (Chaitin [112]). *The probability Ω that a random prefix-free Turing machine will halt on a random input is a left-c.e. 1-random real.*

PROOF. To show that Ω is left-c.e. we take the computably enumerable set $H = \{\sigma \mid U(\sigma) \downarrow\}$. Since each element of U has finite length, we can, uniformly in n , add the probabilities $2^{|\sigma|}$ for all σ enumerated into H by time n . Since this sequence is monotonically increasing and approximates Ω from the left, the result follows.

To show that Ω is 1-random, we follow the proof of [158]. We first describe a particular prefix-free machine M in the following way. At stage s , we consider whether $U(\tau)$ threatens to be a name for Ω witnessing that $K(\Omega \upharpoonright s) < s - c$, where c is the coding constant for M in U . That is, we find n such that $\omega_n \upharpoonright s = \Omega \upharpoonright s$ (this is possible by the proof that Ω is left-c.e.), and search for a τ of length at most $s - c$ such that $U(\tau)_n = \omega_n \upharpoonright s$. If such a τ exists, we find λ outside the range of U_n , and set $M(\tau) = \mu$.

We now verify that this procedure prevents τ as a name for $\Omega \upharpoonright s$. We must have ν with $|\nu| \leq |\tau| + c < s$ such that $U(\nu) = M(\tau)$. However, $U_n(\sigma) \neq \mu$ for any σ , so that $U(\nu)$ does not halt within n steps, and so $\Omega \upharpoonright s \neq \omega_n \upharpoonright s$. Thus, $U(\tau) \neq \Omega \upharpoonright s$ for any τ shorter than $s - c$, as required. \square

Several similar results have been achieved recently, in work of Barmpalias, Cenzer, and Porter [44, 45]. Of course, the exact numerical values in all of these results, including Chaitin's, depend dramatically on the choice of a universal machine. The following result from [44] is typical.

THEOREM 4.2.3 (Barmpalias–Cenzer–Porter). *The following conditions on a real number x are equivalent:*

- (1) *There is a universal oracle Turing machine U such that the probability that U produces a computable output when reading from a random oracle is x .*
- (2) *x satisfies both of the following conditions:*
 - (a) $x \in (0, 1)$,
 - (b) $x = \alpha - \beta$, where α and β are \emptyset' -left-c.e. real numbers.

PROOF. Assuming 1, condition 2a is trivial, so we establish 2b. We begin by showing that x is the difference of two \emptyset' -right-c.e. numbers. Let C be a Π_2^0 class. Then \emptyset' can produce a decreasing sequence of rationals $(c_n : n \in \omega)$ converging to the measure of C , since, as non-members of C are enumerated, measure can be subtracted. Now given a Turing functional T , the class of all reals X such that $T(X)$ is total is a Π_2^0 class. If we can subtract from its measure the measure of the X such that $T(X)$ is total and not computable, we will have the intermediate result. To this end, consider the set I of all X such that $T(X)$ is total and $X \in \bigcup_{i \in \omega} V_i^{T(X)}$, where $(V_i : i \in \omega)$ is a universal Martin-Löf test. Let X be a 2-random real. We will show that $X \in I$ if and only if $T(X)$ is total and $T(X)$ is not computable.

Indeed, let $T(X)$ be total and non-computable. Then if $X \notin \bigcap_{i \in \omega} V_i^{T(X)}$, then X would be random relative to $T(X)$, but $T(X) \leq_T X$, and so $T(X)$ must be Δ_2^0 . However, no 2-random X computes a non-computable Δ_2^0 set, so that $X \in I$. On the other hand, if $X \in I$ and $T(X)$ is 2-random, it must follow that $T(X)$ is not computable.

Now this set I is a Π_2^0 class (and thus has \emptyset' -right-c.e. measure), and has the same measures as the set of X such that $T(X)$ is total and not computable. Since being a difference of \emptyset' -left-c.e. reals is equivalent to being a difference of \emptyset' -right-c.e. reals, condition 2b follows.

The proof from 2 to 1 is more difficult. It makes use of the following well-known fact (see Section 5.1 of [158]), which will be useful in other results later.

LEMMA 4.2.4. *The following are equivalent:*

- (1) *The real number α is X -left-c.e.*
- (2) *There is a prefix-free machine M such that α is the measure of the domain of M^X .*
- (3) *There is a $\Sigma_1^0(X)$ -class of measure α .*

Moreover, these equivalences hold uniformly.

PROOF. If α is X -left-c.e., we can express α as an infinite sum $\sum_{i \in \omega} 2^{-n_i}$ where $(n_i : i \in \omega)$ is a computable sequence. By the Kraft-Chaitin theorem, suitably relativized, we have a machine M whose domain has measure α . The domain of this machine will then be a $\Sigma_1^0(X)$ -class of appropriate measure. Finally, X can enumerate the lower cut of the measure of a $\Sigma_1^0(X)$ -class. \square

Let α, β be \emptyset' -left-c.e. reals and let γ be a 2-random \emptyset' -left-c.e. real. By a result of Demuth [150], the numbers $\alpha_0 = \alpha + \gamma$ and $\beta_0 = \beta + \gamma$ are 2-random \emptyset' -left-c.e. reals, which, of course, have the same difference as α and β . So we can assume that α, β are 2-random.

If $\alpha - \beta$ is also 2-random, then by a result of Rettinger and Zheng [506], we know that $\alpha - \beta$ must be either \emptyset' -left-c.e. or \emptyset' -right-c.e. In the case that it is \emptyset' -left-c.e., we take a universal machine V_0 such that $P_X(V_0(X) \equiv_T \emptyset) = \alpha_0 - \beta_0$, where α_0, β_0 are \emptyset' -left-c.e. reals. Now by a result of [324] there must be some i such that $(\alpha - \beta) - 2^{-i}\alpha_0$ is \emptyset' -left-c.e. and $(\alpha - \beta) - 2^{-i}(\alpha_0 - \beta_0) \in (0, 1 - 2^{-i})$. From these two points, it follows that $(\alpha - \beta) - 2^{-i}(\alpha_0 - \beta_0)$ is \emptyset' -left-c.e. We can then find a machine M_0 such that $M_0(\sigma)$ is undefined for σ comparable with 0^i and where $P_X(M(X) \equiv_T \emptyset) = (\alpha - \beta) - 2^{-i}(\alpha_0 - \beta_0)$. We then define a machine V so that

$$V(\rho) = \begin{cases} V_0(\sigma) & \text{if } \rho = 0^i \sigma \\ M_0(\rho) & \text{otherwise} \end{cases}$$

Now $P_X(V(X) \equiv_T \emptyset) = \alpha - \beta$.

If $\alpha - \beta$ is 2-random and \emptyset' -right-c.e., then we let δ, ϵ be such that

$$(\alpha - \beta) - 2^{-\epsilon}\delta \in (2^{-\epsilon}, 1)$$

and such that δ is a \emptyset' -left-c.e. real. Then, as before, we can find a machine V such that $P_X(V(X) \equiv_T \emptyset) = \delta$.

Now we can find a Σ_2^0 prefix-free set S of strings incomparable with 0^ϵ and such that $\mu[S] = 1 - ((\alpha - \beta) - 2^{-\epsilon}\delta)$. Consequently, we can build a machine N

such that $P_X(N(X) \equiv_T \emptyset) = (\alpha - \beta) - 2^{-e}\delta$. We define a machine M so that

$$M(\rho) = \begin{cases} N(\rho) & \text{if } \rho \perp 0^e \\ V(\sigma) & \text{if } \rho = 0^e\sigma \end{cases}$$

Now $P_X(M(X) \equiv_T \emptyset) = P_X(N(X) \equiv_T \emptyset) + 2^{-e}P_X(V(X) \equiv_T \emptyset) = \alpha - \beta$.

In the case that $\alpha - \beta$ is not 2-random, we pick a 2-random \emptyset' -left-c.e. δ . By a result of [46], it follows (for any e that $(\alpha - \beta) - 2^{-e}\delta$ must be \emptyset' -right-c.e. We then pick e such that $(\alpha - \beta) - 2^{-e}\delta \in (2^{-e}, 1)$, and pick N, V , and M as in the previous case. \square

The question already arises in the original paper [146] whether randomized Turing machines represent any genuine gain in power over deterministic Turing machines. Of course, something of that question depends on what we expect the machine to do. The authors of that paper prove that there is a set which is not computably enumerable, but which is enumerated with positive probability by an object much like a randomized Turing machine, although their formalism is not quite equivalent and this result fails under the definitions given above. Modern approaches to the question have asked either more or less. On the one hand, we might ask that the algorithm be correct with probability bounded above $\frac{1}{2}$, something like the Miller-Rabin algorithm. In that case, we could simulate the algorithm on a (much slower) deterministic machine by the following algorithm: To compute $f(n)$, list sequences in $2^{<\omega}$, and run the randomized machine with those sequences as random bits. When the machine gives the same value on a set of such strings that collectively have measure more than $\frac{1}{2}$, that must be the correct answer, so our deterministic machine reports it.

Of course, in the contexts where the Miller-Rabin algorithm is favored over the AKS class of deterministic primality tests on the basis of being slightly faster but not quite as certain, this method of massive simulation is not practical. Consequently, a large literature now centers on this second path of setting some correctness standard for a randomized Turing machine, and asking whether the same performance could be matched by a deterministic machine, *under given time constraints*. This gives rise to the following definitions.

DEFINITION 4.2.5 (Gill [216]). Let $S \subseteq \{0, 1\}^*$.

- (1) We say that S is of class **BPP** if and only if there is some randomized Turing machine T and some polynomial p such that on input $x \in \{0, 1\}^*$, the machine T halts in at most $p(|x|)$ steps, returning output from $\{0, 1\}$, where $T(x) = \chi_S(x)$ with probability at least $\frac{2}{3}$.
- (2) We say that S is of class **RP** if and only if there is some randomized Turing machine T and some polynomial p such that on input $x \in \{0, 1\}^*$, the machine T halts in at most $p(|x|)$ steps, returning output from $\{0, 1\}$, where if $x \in S$, we have $T(x) = 1$ with probability at least $\frac{2}{3}$, and if $x \notin S$, then $T(x) = 1$ with probability 0.
- (3) We say that S is of class **coRP** if and only if the complement of S is of class **RP**.
- (4) We say that S is of class **ZPP** if and only if there is some randomized Turing machine T and some polynomial p such that on input $x \in \{0, 1\}^*$, the machine T halts in $p(|x|)$ steps *in expectation*, returning output from $\{0, 1\}$, where $T(x) = \chi_S(x)$.

As in many parts of complexity theory, much is unknown about how these classes relate to one another, to \mathbf{P} , to \mathbf{NP} , and to the enormous collection of other classes that have been defined. The following standard result, though, exemplifies the connectedness of these classes and their tractability (at least, relative to the rest of complexity theory).

PROPOSITION 4.2.6. $\mathbf{ZPP} = \mathbf{RP} \cap \mathbf{coRP}$.

PROOF. Suppose T_1 witnesses that $S \in \mathbf{RP}$, and that T_2 witnesses that $S \in \mathbf{RP}$. Then we can simultaneously run $T_1(x)$ and $T_2(x)$, in total time deterministically equal to the sum of two polynomials in $|x|$. If T_1 and T_2 agree, then they must be correct. If not, we repeat the procedure. Since they agree with probability at least $\frac{2}{3}$ on each repetition, we expect agreement on the first trial, so that the full procedure runs in expected polynomial time.

On the other hand, if T witnesses that $S \in \mathbf{ZPP}$, with polynomial p bounding the expected run time of T , we run T for $3p(|x|)$ steps, and then return the output of T , if it halts in that time, and 1 otherwise.

LEMMA 4.2.7 (Markov-Chebyshev Inequality; see p. 47 of [448]). *Let X be a non-negative random variable with expected value $E(X)$. Then $P(X \geq \epsilon) \leq \frac{E(X)}{\epsilon}$.*

Since the run-time is non-negative, we apply the Markov-Chebyshev inequality to see that the probability that the actual run time exceeds three times its expectation is at most $\frac{1}{3}$. Consequently, this algorithm witnesses that S is in \mathbf{RP} . Similarly, if we output 0 when T does not halt in the prescribed time, we see that S is in \mathbf{coRP} . \square

The second half of the preceding proof illustrates an important method in probability: the use of so-called “concentration inequalities.” These results give (often quite loose) bounds on the probability that a random variable takes values in some range. Frequently, the conditions for the inequality to hold are broad enough that, by appropriate choice of parameters, these loose bounds can be made good enough to prove something useful. We will see more examples, especially in Chapter 6, but the technique is worth noting in this proof. The Markov-Chebyshev inequality is an important and simple example of a concentration inequality. It has as consequences several other concentration inequalities of interest. Proofs of these statements can be found in [448].

COROLLARY 4.2.8. *Let X be a random variable and $\epsilon > 0$. Then the following hold:*

- (1) $P(|X| \geq \epsilon) \leq \frac{E(X)}{\epsilon}$
- (2) $P(|X| \geq \epsilon) \leq \frac{E(X^2)}{\epsilon^2}$
- (3) $P(|X - E(X)| \geq \epsilon) \leq \frac{\text{var}(X)}{\epsilon^2}$

The centrality of the concentration inequality method in the proof of Proposition 4.2.6 may explain the often-cited contrast between the case of randomized computation (where we know that $\mathbf{ZPP} = \mathbf{RP} \cap \mathbf{coRP}$) and the case of nondeterministic computation (where we emphatically do not know whether $\mathbf{P} = \mathbf{NP} \cap \mathbf{coNP}$). While we know at least something about the spread of even an arbitrary random variable, no such structure exists on mere path existence, as is called for in nondeterministic computation.

While work on the descriptive complexity of **BPP** is ongoing, it is worthwhile to note that progress has been made. For instance, Eickmeyer and Grohe [168] have reported a logic that captures **BPP**.

4.2.2. Effective Completeness for Continuous First Order Logic. The classical effective completeness theorem is foundational for computable model theory. Let T be a complete first-order theory such that there is a Turing machine to decide, for each first-order sentence of the signature, whether that sentence is or is not in T (we call such a theory *decidable*). Then there is a model $\mathcal{M} \models T$ such that the atomic diagram of \mathcal{M} is classically computable as a set of Gödel codes. This is, in effect, due to the constructive character of Henkin's proof of classical completeness.

Of course, this is no guarantee that an arbitrary model of T will be computable. Indeed, unless T is automorphically trivial any computable model is isomorphic to a non-computable model. However, even when we weaken the question to which models have isomorphic copies which are computable, there are typically many which do not, and a large literature exists on the question of which models of a given decidable theory have, or do not have, computable copies in this sense.

The converse to effective completeness is false. The standard model of arithmetic, $(\mathbb{N}, +, \cdot, 0, 1)$, for instance, is computable, but is a model of true arithmetic, a highly undecidable theory. Moreover, effective completeness is far from the only way to guarantee the existence of a computable structure. Construction of a model by Barwise compactness, or even by direct construction are widespread in the literature.

Nevertheless, the effective completeness theorem provides an important connection between effectiveness of the theory and effectiveness of at least some models. In the present section, we will show that a similar result relates continuous first order logic and randomized computation. The results of this section were originally proved in [93].

The first consideration in a randomized effective completeness theorem is to describe the kind of object that will be said to be effective. On the side of the theory, this is straightforward. As in the first order case, we define a complete theory to be one that is the theory of some model — in the continuous case, if $T = Th(\mathcal{M})$ is the set of all continuous \mathcal{L} -sentences φ such that $\mathcal{M}(\varphi) = 0$. We now define what it means for a continuous theory to be decidable.

DEFINITION 4.2.9 ([63]). Let L be a continuous signature and Γ a set of formulas of L .

- (1) We define

$$\varphi_\Gamma^\circ := \sup \{ \mathcal{M}(\varphi) : \mathcal{M} \models \Gamma \}.$$

- (2) If T is a complete continuous first-order theory, we say that T is *decidable* if and only if there is a (classically) computable function f , called a *decision procedure for T* such that $f(\varphi)$ is an index for a computable real number equal to φ_T° .

In effect, this means that we can, uniformly in φ , compute the truth value of φ required by T of all its models. It is worth remembering that if φ is a sentence, then $1 \div \varphi$ is also a sentence, so that f will also compute the truth value of $1 \div \varphi$ required by T . Since a particular structure will have a single truth value for φ , a complete theory T will include information on both, so that the value of φ is

uniquely determined. In a decidable theory, it is uniquely determined to be a real number which is computable, uniformly in φ .

We now turn our attention to the models. As in the classical case, a model is identified with its atomic diagram, but since constants are not continuous in the most interesting continuous structures, some modification is required. We replace each true constants of classical first-order logic with a unary predicate for the distance from the constant's interpretation. In this way, we arrive at the following definition.

DEFINITION 4.2.10. Let L be a computable continuous signature (i.e. one where the sets of predicate and function symbols, the set of moduli, and the arity function are all computable). Let \mathcal{M} be a continuous L -structure. Let $L(\mathcal{M})$ be the expansion of \mathcal{L} by a constant c_m for each m in the universe of \mathcal{M} (i.e. a unary predicate $c_m \in \mathcal{R}$ where $c_m^{\mathcal{M}}(x) := d(x, m)$). Then

- (1) The *continuous atomic diagram* of \mathcal{M} , written $D(\mathcal{M})$ is the set of all pairs (φ, p) , where φ is a quantifier-free (i.e. sup- and inf-free) sentence in $L(\mathcal{M})$ and $\mathcal{M}(\varphi) = p$.
- (2) The *continuous diagram* of \mathcal{M} , denoted $D^*(\mathcal{M})$, is the set of all pairs (φ, p) , where φ is a sentence in $L(\mathcal{M})$ and $\mathcal{M}(\varphi) = p$.

Equipped with this definition, we define what it means for a structure to be effective in the sense of randomized computation.

DEFINITION 4.2.11. Let \mathcal{M} be a continuous structure.

- (1) We say that \mathcal{M} is *probabilistically computable* if and only if there is some randomized Turing machine T such that, for every pair $(\varphi, p) \in D(\mathcal{M})$, the machine T accepts φ with probability p .
- (2) We say that \mathcal{M} is *probabilistically decidable* if and only if there is some randomized Turing machine T such that for every pair $(\varphi, p) \in D^*(\mathcal{M})$, the machine T accepts φ with probability p .

We now set about to prove the effective completeness theorem for randomized computation and continuous first order logic. Much of the work of constructing the model is carried out in [63]; only some effectivization was necessary to complete the following theorem in [93].

THEOREM 4.2.12. *Let T be a complete decidable continuous first-order theory. Then there is a probabilistically decidable continuous structure $\mathcal{M} \models T$.*

PROOF. We first extend T to a consistent set Γ of formulas which is Henkin complete; that is, a formula in which for every formula φ , every variable x , and every pair $p < q$ of diadic rationals, there is a constant c such that

$$\left(\sup_x \varphi \div q \right) \wedge (p \div \varphi[c/x]) \in \Gamma.$$

This notion of Henkin completeness was used in [63]. We let $L_0 = L$ and $\Gamma_0 = T$. At stage n , we add, for each formula $\varphi \in L_n$ a new constant $c_{(\varphi, x, p, q)}$ to form L_{n+1} . We form Γ_{n+1} by adding the formula

$$\left(\sup_x \varphi \div q \right) \wedge (p \div \varphi[c_{(\varphi, x, p, q)}/x]).$$

We observe that this construction of $\Gamma = \bigcup_i \Gamma_i$ is effective uniformly in L and T .

We can also — still uniformly in L, T , and s , compute a sequence Δ_s of sets such that $\Delta^0 := \bigcup_{s \in \omega} \Delta_s$ such that $\Gamma \subseteq \Delta^0$ and such that for any $\varphi, \psi \in L^* = \bigcup_{n \in \omega} L_n$, we have either $\varphi \dot{\vdash} \psi \in \Delta^0$ or $\psi \dot{\vdash} \varphi \in \Delta^0$. Indeed, we start with $\Delta_0 = \Gamma_0$, and at stage s , we take the first φ, ψ requiring attention, and check (effectively, since the theory is decidable) whether Δ_s proves that $\psi \dot{\vdash} \varphi$ has value less than $\frac{1}{2^n}$. Accordingly, we add either $(\psi \dot{\vdash} \varphi) \dot{\vdash} \frac{1}{2^n}$ or $(\varphi \dot{\vdash} \psi) \dot{\vdash} \frac{1}{2^n}$ to form Δ_{s+1} .

As an intermediate step, we construct two machines: an accepting machine M_A and a rejecting machine M_R . At stage s , if $\Delta_s \vdash \varphi \dot{\vdash} \frac{k}{2^n}$, we will enumerate $1 - \frac{k}{2^n}$ into the left cut of $\alpha_{\varphi, A}$. Similarly, if $\Delta_s \vdash \frac{k}{n} \dot{\vdash} \varphi$, we enumerate $\frac{k}{2^n}$ into the left cut of $\alpha_{\varphi, R}$.

Now, using the uniformity of Lemma 4.2.4, we produce a single machine M_A such that $M_A(\varphi)$ halts with probability $\alpha_{\varphi, A}$ and a single machine M_R such that $M_R(\varphi)$ halts with probability $\alpha_{\varphi, R}$, so that the domains of $M_A(\varphi)$ and $M_R(\varphi)$ are disjoint. We now define our final machine M by running both M_A and M_R on φ , accepting if M_A halts and rejecting if M_R halts. Now by construction of Δ_s , the machine M accepts φ with probability exactly $\mathcal{M}(\varphi)$. \square

Complexity is a notoriously fraught area for computable model theory, because of sensitivity to representation of the elements of an infinite structure — indeed, a choice of whether a natural number is represented by tallies or by binary representation can make a difference of exponential time. Still, it is worthwhile to note the following connection to the complexity-theoretic aspects of randomized computation, proved in [93].

PROPOSITION 4.2.13. *We say that a continuous structure is polynomial time if and only if there is some polynomial-time randomized Turing machine that, for any $(\varphi, p) \in D(\mathcal{M})$, accepts φ with probability p .*

- (1) *Let $A \in \mathbf{BPP}$. Then there is a polynomial-time probabilistically computable structure \mathcal{M} , with a polynomial-time function $f : \{0, 1\}^* \rightarrow \mathcal{M}$ such that there is a quantifier-free formula $\varphi(x)$ such that $\mathcal{M}(\varphi(x)) \leq \frac{1}{3}$ for $x \in f(A)$, and $\mathcal{M}(\varphi(x)) \geq \frac{2}{3}$ for $x \notin f(A)$.*
- (2) *Let \mathcal{M} be a polynomial time probabilistically computable structure, and let A, B be quantifier-free definable subsets of \mathcal{M}^n , whose distance is bounded away from zero, and where $A \cup B$ is polynomial-time computable. Then each of A and B is of class \mathbf{BPP} .*

PROOF. For the first point, we make a structure on the universe of which A is naturally a subset, with the discrete metric, and a unary predicate $\mathcal{A}(x)$, interpreted as the probability that x is accepted by the machine that witnesses $A \in \mathbf{BPP}$. For the second point, for any $\bar{a} \in A \cup B$, we can decide whether $\bar{a} \in A$ by evaluating $\mathcal{M}(\varphi(\bar{a}) \dot{\vdash} \psi(\bar{a}))$. \square

A more recent approach to effective completeness — and, indeed, to computable metric structures generally, has been followed by Camrud, McNichol, and others. In this approach, more closely mirroring the constructivizations of the Russian school [227], the key concept is the *presentation*. Given a continuous L -structure \mathcal{M} , a presentation of \mathcal{M} is a pair (\mathcal{M}, g) , where $g : \mathbb{N} \rightarrow |\mathcal{M}|$, where the algebra generated by $\text{ran}(g)$ in $|\mathcal{M}|$ under the functions of L is dense in \mathcal{M} . We say that the presentation is *computable* if and only if the predicates of \mathcal{M} are uniformly computable on the elements of this dense algebra. This treatment of effectivity for

continuous structures can be found in [89, 192], and the corresponding effective completeness theorem can be found in [104].

4.2.3. Pseudorandom Generators and Derandomization. Consider a set S of class **BPP**. The obvious quick way to check whether $n \in S$ is to provide some random bits and see what the randomized algorithm does on them. Since the algorithm is required to be right a great majority of the time, we could take a majority vote of the results.

Of course, this doesn't work quite so easily. In particular, the entirety of Chapter 3 explores the many ways in which it is impossible for us to simply "provide some random bits." An interesting hypothesis, which is the subject of intense current interest in complexity theory, is that it may not be necessary to be random. It may only be necessary to be so much like random that the algorithm can't tell that it isn't random.

Obviously the literature in this area is large and fast-moving. Knuth [316] described many particular algorithms for generation of numbers that were "random enough" for certain applications, and the use of a random variable with one distribution to "simulate" (to generate a random variable with) another distribution is well-established in probability (see [421]). However, the literature on general-purpose pseudorandom generators of the type we describe here seems to have started with a paper of Yao [504]. The interested reader will find a much more extensive survey in [224] and [30], including most of the material of the present section.

Many of the results in this area are — perhaps necessarily (see Theorem 4.2.16) expressed in the language of Boolean circuits. In this model a k -ary Boolean circuit of size m takes k boolean inputs, and applies binary AND and OR, and unary NOT operations to compute a function, with m total operations performed.

DEFINITION 4.2.14. Let $\ell : \mathbb{N} \rightarrow \mathbb{N}$ be monotonically increasing. A *pseudorandom generator of stretch ℓ* is a deterministic Turing machine G such that for any $\sigma \in \{0, 1\}^k$,

- (1) The machine G runs in time polynomial in $2^k \ell(k)$ steps and outputs a string of length $\ell(k)$, and
- (2) If U_j is a uniform random binary string of length j , then for every Boolean circuit D_k of size $\ell(k)^2$, the circuit accepts $G(U_k)$ and $U_{\ell(k)}$ with probabilities that differ by less than $\frac{1}{6}$.

Something should be said about the formulation of this definition. In particular, the use of Boolean circuits in item 2 is striking. Generally, Boolean circuits are a non-uniform model of computation — the circuits on different numbers of inputs are not, *a priori*, required to have anything to do with one another, which allows such counterintuitive results as the fact that every set of natural numbers in unary representation is computable by a system of relatively small circuits. The point in the present application, though, is that nothing that a randomized Turing machine might want to do with its random bits should depend so sensitively on the difference between $G(U_k)$ and $U_{\ell(k)}$ that the outcome of a majority vote could be changed.

It also needs to be recognized that the *existence* of pseudorandom generators is already deeply problematic. As with much of complexity theory, so much is unknown that one can find entire books developing the consequences of and conditions for the existence of objects whose existence itself is unknown. However, this custom is at odds with the usual practice of mathematics, and so it needs to be noted. The

payoff is that many complexity classes based on randomization collapse under the assumption that certain pseudorandom generators exist.

Of course, even if a pseudorandom generator exists, it only reduces the number of random bits needed. If the randomized algorithm needs $\ell(k)$ random bits, we must still come up with k of them. Another obstacle still to be overcome is that the definition only calls for the pseudorandom generator to run in time exponential in its input. It is not obvious that an exponential time algorithm should require anything built from it to run in polynomial time. Nevertheless, the following theorem shows that the definition given is enough.

THEOREM 4.2.15. *If for some $\epsilon > 0$ there is a pseudorandom generator of stretch $k \mapsto 2^{\epsilon k}$, then $\mathbf{BPP} = \mathbf{P}$.*

PROOF. Let G be a pseudorandom generator of stretch $k \mapsto 2^{\epsilon k}$. Let T be a randomized Turing machine witnessing that $S \in \mathbf{BPP}$. We define a deterministic Turing machine U as follows. On input n , suppose that T runs in s steps. Let k be the least natural number such that $2^{\epsilon k} \geq s$. We let U search over all binary strings σ of length k , computing, for each one, $T^{G(\sigma)}(n)$, and output the majority result. Note that the computation $\sigma \mapsto T^{G(\sigma)}(n)$ can be represented by a circuit of size $2^{2\epsilon k}$, for each n , this transformation must accept with the same probability (plus or minus error of less than $\frac{1}{6}$) as the transformation $\tau \mapsto T^\tau(n)$, where τ ranges over all strings of length $2^{\epsilon k}$. Since T was correct on set of $\{0, 1\}^{2^{\epsilon k}}$ with probability greater than $\frac{2}{3}$, and since the error introduced by G was less than $\frac{1}{6}$, the majority vote still gives the correct solution. Since U searches over strings of length k , it must compute G a total of 2^k times, each of which takes time polynomial in $2^{k+\epsilon k}$. However, k is approximately $\log s$, where s is polynomial in the size of n , so that U runs in polynomial time. \square

The same strategy applies much more broadly, allowing many complexity classes to be collapsed if a pseudorandom generator of the right stretch exists.

As has been pointed out, the existence of pseudorandom generators is problematic. There are several well-known results giving other difficult conditions necessary or sufficient to the existence of pseudorandom generators with particular stretch. An important example is a result of Nisan and Wigderson [388].

THEOREM 4.2.16. *Let $\ell : \mathbb{N} \rightarrow \mathbb{N}$ be monotonically increasing, and $\ell(n) \leq 2^n$ for all n . Then the following are equivalent:*

- (1) *There is some $c > 0$ such that $\mathbf{EXPTIME}$ cannot be approximated by Boolean circuits of size $\ell(m^c)$.*
- (2) *There is some $c > 0$ such that there is a pseudorandom generator of stretch ℓ .*

While the existence of pseudorandom generators sufficient to collapse \mathbf{BPP} to \mathbf{P} is currently unknown, there are objects called expander graphs, unconditionally known to exist, that give some reduction in the length of the string of random bits needed by certain randomized Turing machines. These graphs arise naturally in the study of random graphs, and will be seen in Section 5.1.1.

4.3. Interactive Proofs

4.3.1. Interactive Proofs as Games. The complexity classes arising from randomized computation can be viewed as a modification of those arising from

nondeterminism: Instead of requiring that there exist one path, we require that there exist a lot of paths. Another modification in a similar spirit views **NP** as a proof environment. We have already hinted at this view in the description of the “proof” and the “verifier” in comments after the definition of **NTIME**.

In this view, an agent (the “prover”) using a lot of computing power finds a proof, if one exists, and then communicates it to the verifier, who is required to check it in polynomial time. We can imagine an environment in which more complex communication takes place between the prover and the verifier. This is the idea behind interactive proofs, introduced in slightly different forms independently by [226, 39].

DEFINITION 4.3.1. Let $S \subseteq \{0, 1\}^*$. We say that S is of class **IP** if and only if there is a randomized polynomial-time Turing machine V and a natural number constant k with the following properties:

- (1) If $\sigma \in S$, there is a function $P : \{0, 1\}^* \rightarrow \{0, 1\}^*$ so that the sequence $V(\sigma), P(\sigma, V(\sigma)), V(\sigma, V(\sigma)), P(\sigma, V(\sigma)), \dots$ achieves a 1 in the k th term with probability at least $\frac{2}{3}$.
- (2) If $\sigma \notin S$, then for any function $P : \{0, 1\}^* \rightarrow \{0, 1\}^*$, the sequence described in part 1 achieves a 1 in the k th term with probability at most $\frac{1}{3}$.

The idea is that if $\sigma \in S$, there should be some prover that can prove this to the verifier’s general satisfaction, but if $\sigma \notin S$, no prover should reliably convince the verifier otherwise.

Some examples will illustrate the significance of the definition. Recall that the isomorphism on finite graphs is known to be of class **NP**, since it is easy to check whether a given function is an isomorphism.

PROPOSITION 4.3.2 (Goldreich–Micali–Wigderson 1991 [225]). *The isomorphism problem for finite graphs is of class **IP**.*

PROOF. A straightforward proof arises from the fact that the isomorphism problem for finite graphs is **NP**. To achieve this, the prover need only find an isomorphism and send it to the verifier. The verifier then checks (in polynomial time) that the function given is, in fact, an isomorphism.

Another proof system is possible, though, which has the additional interesting property that the prover does not have to reveal any information to the verifier beyond the fact that the two graphs are isomorphic (a so-called *zero knowledge proof*). On input (G_0, G_1) , the prover selects uniformly at random a permutation π of the finite set of vertices of G_0 , and interprets this as a graph $H \cong G_0$, and sends H to the verifier. The verifier chooses $\alpha \in \{0, 1\}$ uniformly at random, and sends α to the prover. If $\alpha = 0$, the prover sends π , and if $\alpha = 1$, the prover sends $\pi \circ f$, where f is an isomorphism from G_0 to G_1 , if any such isomorphism exists. The verifier checks whether the function sent is an isomorphism. If so, the procedure is repeated a second time, accepting if the second function is also an isomorphism. If either function fails to be an isomorphism, it rejects.

If $G_0 \cong G_1$, then the prover will always be able to send the verifier an isomorphism. Otherwise, the verifier will reject with probability at least $\frac{3}{4}$, since with that probability it will have asked about isomorphism of H with G_1 at least once. \square

Graph non-isomorphism is not known to be in **NP**. In that sense, the following result was initially viewed as surprising.

PROPOSITION 4.3.3 (Goldreich–Micali–Wigderson 1991 [225]). *The non-isomorphism problem for finite graphs is of class **IP**.*

PROOF. On input (G_0, G_1) , the verifier chooses α_0, α_1 uniformly at random, and selects uniformly at random a permutation π_i of G_{α_i} , interpreting it as a graph $H_i \cong G_{\alpha_i}$, and sends (H_0, H_1) to the prover. The prover computes, for each i , whether $H_i \cong G_0$ or $H_i \cong G_1$, and sends the verifier (β_0, β_1) , where $H_i \cong G_{\beta_i}$. The verifier accepts if $\beta_i = \alpha_i$ for each i , and rejects otherwise.

If G_0 and G_1 are not isomorphic, then each H_i is isomorphic to exactly one of G_0, G_1 , and the prover can distinguish the two cases, so that $\vec{\beta} = \vec{\alpha}$ and the verifier accepts. Otherwise, the probability that both $\alpha_0 = \beta_0$ and $\alpha_1 = \beta_1$ is at most $\frac{1}{4}$. \square

Although much is unknown about the complexity of graph isomorphism and non-isomorphism, these two results suggest, at least, that **IP** is a very large class. In 1990, it was shown that **IP** contains the polynomial hierarchy, and shortly afterward that it is equal to **PSPACE**, the class of problems decidable in arbitrary time using a polynomially bounded number of cells on the tape of the Turing machine.

PROPOSITION 4.3.4 ([343]). ***IP** contains the polynomial hierarchy.*

PROOF. The permanent $\text{per}(A)$ of an $r \times r$ matrix A is the sum

$$\sum_{\sigma \in S_r} \left(\prod_{i=1}^r a_{i\sigma(i)} \right).$$

Equivalently,

$$\text{per}(A) = \sum_{i=1}^r \text{per}(A_{\{1i\}}),$$

where the matrix minor $A_{\{ij\}}$ is the result of deleting the i th row and j th column of A . There is a nondeterministic machine M such that for any 0-1 matrix A , the computation $M(A)$ has $\text{per}(A)$ accepting paths (this follows from the recursive definition just given); that is, the permanent function is of class **#P**.

Moreover, the permanent function is known to be complete in this class [489]. Consequently, the set

$$L = \{(A, s) : A \text{ is a 0-1 matrix and } \text{per}(A) = s\}$$

is complete among problems solvable in polynomial time with an oracle for solving problems in **#P**. Since this class contains **PH**, by a result of Toda [485], it suffices to show that L has an interactive proof system.

To prove that $(A, s) \in L$, where A is an $r \times r$ matrix with entries from $\{0, 1\}$, we use the following procedure. The prover picks a prime p such that $r! < p < 2^r$, and sends both p and a proof that p is prime to the verifier. All later arithmetic in the procedure is done modulo p . Set $\mathcal{L}_0 = \{(A, s)\}$.

In stage t , the verifier checks whether \mathcal{L}_t is a singleton (B, q) where B is a 1×1 matrix. If so, the verifier will accept if $q = \text{per}(B)$, and reject otherwise.

If \mathcal{L}_t is a singleton with a larger matrix, the verifier will construct the minors $B_{\{1,i\}}$ for each i , and send them to the prover. The prover returns the permanent

$q_i = \text{per}(B_{\{1,i\}})$. The verifier then computes

$$\sum_{i=1}^r b_{1i} q_i.$$

If this quantity is not equal to q , the verifier rejects. Otherwise,

$$\mathcal{L}_{t+1} = \{(B_{\{l,i\}}, q_i) : 1 \leq i \leq \dim(B)\}.$$

If \mathcal{L}_t is not a singleton, then the verifier chooses the first two pairs in \mathcal{L}_t , say (B_1, q_1) and (B_2, q_2) and sends them to the prover. Note that at any time t , all matrices in \mathcal{L}_t have the same dimension. The prover computes

$$f(x) = \text{per}(C + x(D - C)),$$

a polynomial of degree at most equal to the dimension of C and D , and sends f_0, \dots, f_k to the verifier, where

$$f(x) = \sum_{i=0}^k f_i x^i.$$

The verifier then evaluates this polynomial on 0 and 1, and checks whether $f(0) = c$ and $f(1) = d$. If either is not equal, the verifier rejects. Otherwise, the verifier sets

$$\mathcal{L}_{t+1} = (\mathcal{L}_t - \{(C, c), (D, d)\}) \cup \{((C + a(D - C)), g(a))\}$$

for some a chosen uniformly at random from \mathbf{Z}_p .

The verifier will certainly either accept or reject by the time t at which \mathcal{L}_t consists of a singleton whose matrix is 1×1 . This will happen in at most $2(r - 1) + (r - 2) + \dots + 1$ steps. If $\text{per}(A) = s$, then the verifier will accept (A, s) .

We now consider the case in which $\text{per}(A) \neq s$, and show that the probability of accepting is low. Notice that the verifier will eventually accept if and only if at some stage t every pair $(B, q) \in \mathcal{L}_t$ has the property that $\text{per}(B) = q$. This cannot attain in either of the cases where \mathcal{L}_{t-1} is a singleton. We assume, then, that we have a stage t in which there is some $(B, q) \in \mathcal{L}_t$ such that $\text{per}(B) \neq q$, but there is no such $(B, q) \in \mathcal{L}_{t+1}$. What must be shown is that the step which replaces two entries of \mathcal{L}_t by one will, with high probability, preserve the property that $\text{per}(B) \neq q$ for some element of \mathcal{L} .

Let $(C, c), (D, d) \in \mathcal{L}_t$, with either $\text{per}(C) \neq c$ or $\text{per}(D) \neq d$. Note that if C, D are of dimension $r \times r$, the polynomial $f(x) = \text{per}(C + x(D - C))$ over \mathbb{Z}_p is of degree at most r . At $x = 0$ this polynomial should compute the permanent of C , and at 1 it should compute the permanent of D . Suppose that the prover has sent a degree r polynomial g different from f . If the verifier does not reject, $g(0) = c$ and $g(1) = d$, so one of them does not match the permanent of the appropriate matrix, so that f and g are not identical. Consequently, they can agree on at most r elements of \mathbb{Z}_p , so that the probability that a , chosen uniformly at random from \mathbb{Z}_p is a point of agreement is at most $\frac{r}{p}$. Since $p > r!$ and the total probability of a spurious acceptance is at most

$$\frac{(2(r - 1) + (r - 2) + \dots + 1)r}{p},$$

the proof is complete. \square

The following result is originally due to Shamir [440], but we follow the simpler proof by Shen [446] as described in [30].

THEOREM 4.3.5. **IP = PSPACE.**

PROOF. Consider the set B of propositional formulas $\varphi(X_1, \dots, X_n)$ with propositional atoms X_1, \dots, X_n . A *quantified Boolean formula* is an expression of the kind

$$Q_1 X_1 Q_2 X_2 \cdots Q_n X_n \varphi(X_1, \dots, X_n),$$

where $\varphi \in B$ and where for each i , the quantifier Q_i is either \forall or \exists . If ψ is a quantified Boolean formula, we say that it is true if it is a true predicate sentence where the quantifiers range over the Boolean constants \top and \perp . The problem of determining which quantified Boolean formulas are true is known to be **PSPACE** complete (see, for instance, [30]). Consequently, to show that **PSPACE** \subseteq **IP** it suffices to show that this problem has an interactive proof system.

We can read any propositional formula as a polynomial: we interpret $\neg X_i$ as $(1 - X_i)$, we interpret conjunction as a product; disjunction may be interpreted by deMorgan's law. If the variables take values from $\{0, 1\}$, the polynomial P_φ interpreting φ will also take values from $\{0, 1\}$, and this value will correspond to the truth of the formula. We can interpret $\forall X_i$ as a product of the cases $X_i = 0$ and $X_i = 1$, and $\exists X_i$ as a sum, where the formula is true if the polynomial evaluates to some value greater than zero. A first intuition to give interactive proof for true Boolean formulas is simply to give interactive proofs for an equation asserting the value of the appropriate polynomial. However, since the products tend to increase the degree of the polynomial, they may give us a polynomial that the time-bounded verifier cannot evaluate.

To avoid this, we introduce another “quantifier”-like operator. We define

$$L_i : \mathbf{Z}_p[X_1, \dots, X_n] \rightarrow \mathbf{Z}_p[X_1, \dots, X_n]$$

by

$$\begin{aligned} L_i(p(\overline{X})) &:= X_i p(X_1, \dots, X_{i-1}, 1, X_{i+1}, \dots, X_n) + \\ &\quad + (1 - X_i) p(X_1, \dots, X_{i-1}, 0, X_{i+1}, \dots, X_n). \end{aligned}$$

This “linearization” operator has the property that for all $\overline{X} \in 2^n$, we have $L_i(p(\overline{X})) = p(\overline{X})$. Note also that $L_i(p)$ always has degree 1 in X_i . Consequently, we have

$$\begin{aligned} &Q_1 X_1 Q_2 X_2 \cdots Q_n X_n \varphi(X_1, \dots, X_n) = \\ &= Q_1 X_1 L_1 Q_2 X_2 L_1 L_2 \cdots Q_n X_n L_1 L_2 \cdots L_n \varphi(X_1, \dots, X_n). \end{aligned}$$

Moreover, the expression on the right has size $O(n^2)$.

Now let $f \in \mathbf{Z}_p[X_1, \dots, X_n]$, and suppose, for induction, that for any $\bar{a} \in \mathbf{Z}_p^n$, there is an interactive proof system for $f(\bar{a}) = C_0$ in which the verifier accepts with probability 1 if the equation is true and rejects with probability $1 - \epsilon$ if it is false. To verify $\exists X_1 f(X_1, a_2, \dots, a_n) = C_1$, the prover sends a polynomial $g \in \mathbf{Z}_p[X_1, \dots, X_n]$ of appropriate degree r , and the verifier checks it for equality to $f(X_1, a_2, \dots, a_n)$ in the following way: If $g(0) + g(1) \neq C_1$, the verifier rejects. Otherwise, it sends the prover a , uniformly randomly chosen from \mathbf{Z}_p , and asks for a proof that $g(a, a_2, \dots, a_n) = f(X_1, a_2, \dots, a_n)$. As in the proof of Theorem 4.3.4, the probability that distinct g, f of degree r agree on a is at most $\frac{r}{p}$. Similarly, to verify $\forall X_1 f(X_1, a_2, \dots, a_n)$, we follow the same procedure, except that the verifier checks $g(0)g(1) = C_1$. Finally, to verify $LX_1 f(X_1, a_2, \dots, a_n)$, the prover sends a polynomial $g(X_1, \dots, X_n)$ of appropriate degree, and the verifier checks if $a_1 g(0) +$

$(1 - a_1)g(0) = C_1$. If not, it rejects. Otherwise, it picks a random $a \in \mathbb{Z}_p$, and asks for a proof that $g(a) = f(a, a_2, \dots, a_k)$.

As in the proof of the previous theorem, with high probability the new statement to be verified is still false if the original was false. \square

Note that the statement that $S \in \mathbf{IP}$ is a statement about winning strategies for certain games. The games in question are similar to, but not precisely like, several types of games that are well-studied. At least one reasonable construction is this: If $\sigma \notin S$, then there is no winning strategy for player P in a game where P wins if V accepts and loses otherwise. If $\sigma \in S$, then P does have a winning strategy for this game.

The random coins are at least one level of preventing a straightforward interpretation of this, and they are essential to the definition of \mathbf{IP} . This randomness can be incorporated by viewing the game as a *stochastic game* in the sense of Shapley [442]. In these games, when the players choose their alternatives at a particular stage, the game moves to a new position determined by a random variable that depends on the position and the alternatives chosen.

The usual treatment of these games, however, centers on their expectation, rather than the concentration required in the definition of the class \mathbf{IP} . For instance, these games have equilibria [369, 368, 494]. Condon [130] showed that the complexity of deciding which player has the greatest chance of winning is of class $\mathbf{NP} \cap \mathbf{coNP}$, at least for a large class of stochastic games. What is wanted for \mathbf{IP} is not the expected value of the game, but the probability that each player will win.

An approach to this has been made in recent research of Kiefer and others [311]. In this paper, it is shown that these games sometimes exhibit a property called “strong determinacy.” This is exactly the condition that one player has a strategy that will enforce that a winning condition holds with probability at least (or at most) c . If $c = 1$, all such games are strongly determined. However, if $c \in (0, 1)$, which is what we care about for interactive proofs, this condition may not hold.

There is still another obstacle. As Condon [130] points out, much of the literature on stochastic games does not impose resource bounds on the players, and the asymmetric resource bound (polynomial-time verifier, unbounded prover) in interactive proofs is a significant barrier between the game research and the \mathbf{IP} research.

While the quantum features of \mathbf{MIP}^* are beyond the scope of the present work, it is germane to point out to the reader that the interactive proofs described here are a natural entry point to understanding the recent result that $\mathbf{MIP}^* = \mathbf{IP}$, proved in [280], and the related work on the Connes Embedding Problem [221].

4.3.2. Interactive Proofs as Proofs. While complexity theorists are perfectly comfortable calling the objects of the previous section “proofs,” this will likely seem strange to logicians. We are accustomed to seeing a proof as a sequence, to be sure, but it is a sequence whose every step consists of syntactically meaningful “calculations” to determine the truth value of a proposition. More exact definitions could be given, and have been. Is there, then, some meaningful sense in which an interactive proof (in the sense of the previous section) is a proof?

There is actually something like interactive proof in logic. Hintikka and others have described a “Game-theoretic semantics” for various logics, including classical

first-order logic (see, for instance, the survey [255]). This does not transparently encompass the full power of **IP**, but it suggests a direction for further work in this area.

DEFINITION 4.3.6. Given a first-order L -structure \mathcal{M} and an L -sentence φ , we define the game $G(\varphi, \mathcal{M})$ between a *prover* and a *verifier* by induction on the form of φ as follows:

- (1) If φ is an atomic formula, the prover wins if φ is true in \mathcal{M} , and loses otherwise.
- (2) If $\varphi = \neg\psi$, then the prover and the verifier trade positions and continue the game as $G(\psi, \mathcal{M})$.
- (3) If $\varphi = \psi_1 \vee \psi_2$, then the prover chooses $i \in \{1, 2\}$, and the game continues as $G(\psi_i, \mathcal{M})$.
- (4) If $\varphi = \psi_1 \wedge \psi_2$, then the verifier chooses $i \in \{1, 2\}$, and the game continues as $G(\psi_i, \mathcal{M})$.
- (5) If $\varphi = \exists x\psi(x)$, then the prover chooses some c in the domain of \mathcal{M} , and the game continues as $G(\psi(c), \mathcal{M})$.
- (6) If $\varphi = \forall x\psi(x)$, then the verifier chooses some c in the domain of \mathcal{M} , and the game continues as $G(\psi(c), \mathcal{M})$.

Now if φ is true, then the prover always has a winning strategy — a way of making all necessary choices in such a way as to win. If φ is false, then the verifier always has a winning strategy. Indeed, in keeping with the results of Section 4.1.3, if \mathcal{M} is finite, then both the verifier and the prover, when they have a winning strategy, have one that they can compute in time polynomial in the size of the problem instance (φ, \mathcal{M}) .

In that sense, the game-theoretic semantics of Hintikka represent another, perhaps more concrete proof, that for any finite structure \mathcal{M} , the set of sentences true in \mathcal{M} is of class **IP**. At the same time, the moves of the game look very much like an actual proof of φ (from, say, the complete diagram of \mathcal{M}).

Of course, this game uses very little of the power available in **IP**. Not only is the prover using a time-bounded strategy, but the randomization is completely absent, and with it the opportunity for the wrong conclusion to be reached with some positive probability.

It is not immediately clear what strengthening of the proof system of Definition 4.3.6 would be logically sensible and have a chance of matching the power of **IP**.

Hintikka and Sandu point out that game-theoretic semantics have been used for quite a long time for extremely diverse logics. It may be reasonable, then, to ask what a game-theoretic semantics for continuous first-order logic would look like. This seems like a difficult question for several reasons. At the level of quantifiers, there is, in general, no *a priori* reason that suprema and infima must be realized, meaning that those clauses must look very different from the ones in Definition 4.3.6.

Moreover, even handling $\psi_1 \div \psi_2$ seems difficult. However, if the players are allowed randomization, it seems germane to point out that if r is picked uniformly at random from $[0, 1]$, the truth value of $\psi_1 \div \psi_2$ corresponds exactly to the probability that $\mathcal{M}(\psi_2) \leq r \leq \mathcal{M}(\psi_1)$. Making a game for that is still not straightforward, but it gives some cause for hope that there may be something like a proof system corresponding to **IP**.

4.4. Word Problems

In Section 4.2, we suggested two kinds of randomization that could be considered in computation. The first kind, which we have considered up to this point, involves an extrinsic randomization: the algorithm has access to some random bits that it can use, for instance, to simulate, or pick some “place” to check. A second possibility is randomization in the input. This approach has a long history in the theory of algorithms (see, for instance, [138]), where it is used to consider how an algorithm would perform under conditions other than the best or worst case. Examples arising in group theory have led to a collection of new work in computability.

4.4.1. Groups with Solvable Word Problem. Up to this point, our description of randomized computation has focused on a randomization within the algorithm. There is a very different dimension of randomization possible, and it occurs commonly in practice: randomization in the input.

Consider, for example, the simplex algorithm in linear programming. It is well-known that the worst case complexity of the algorithm is exponential [314]. However, the algorithm empirically runs in such short time on almost all practically useful problem instances to make it the industry standard [83]. Intuitively, the “hard” instances of the problem that establish the worst-case complexity are relatively rare, and “most” instances allow a short running time.

In complexity, one way to handle this is with “average-case” complexity analysis, which can be sensitive to a choice of distribution on inputs. Moreover, a computation that does not halt at all, if it arises with any nonzero probability, must cause significant problems with a complexity measure based on the expected value. A more robust method is used for computability, and it first arose in the context of group theory.

Group theory is perhaps a natural field for the consideration of such problems. The word problem for groups is not solvable, but the standard examples proving that it is unsolvable are not only special groups; they are groups in which particular — apparently rare — words witness the unsolvability.

We first consider the classical problem, and in the following section we will take up the model of computability used to formalize the informally rare nature of the obstructions to computability.

It is well-known that a group can be presented by a list of generators and a list of relations. In particular, we write

$$G = \langle \{g_i : i \in I\} | \{r_i : i \in J\} \rangle$$

where each r_i is a word in the letters g_i and their inverses. In this presentation system, G is the quotient of the free group on generators $\{g_i : i \in I\}$ by the smallest normal subgroup containing all the relators $\{r_i : i \in J\}$. Any group can be presented in this way, and the presentations of groups are highly non-unique. Indeed, a significant open problem in group theory, the Andrews-Curtis Conjecture, involves recognizing presentations of the trivial group [22]. Additional background on group presentations can be found in standard algebra texts, such as [274], or in the group theory text [422], which gives one of the standard expositions on the unsolvability of the word problem.

In 1911, Dehn [147] identified three “Fundamental Problems Of Infinite Discontinuous Groups.” He envisioned each group being given by finitely many generators and relations. The three problems were as follows:

The Word Problem: (Dehn called this the “Identity Problem.”) In a particular presentation of a particular group, give an algorithm which will, given an element of the group (in the form of a word on the generators and their inverses, decide whether that element is equal to the identity or not.

The Conjugacy Problem: (Dehn called it the “Transformation Problem.”) In a particular presentation of a particular group, give an algorithm which will, given two elements of the group, decide whether those elements are conjugate or not.

The Isomorphism Problem: Given two group presentations decide whether the two groups are isomorphic or not.

Of course, for many well-known classes of groups, these problems are known to be solvable. Dehn himself showed that the word problem for fundamental groups of certain manifolds must be solvable [148]. Of course, positive solutions tend to require less logical sophistication than negative solutions. However, after the definition of computability in the 1930s, it was clear, at least, what a negative solution could look like.

Indeed, not long after Dehn, Thue had posed the word problem for semigroups [483], and Post [408] proved it unsolvable in 1947 by a method that, with appropriate modifications, would also settle the word problem for groups.

THEOREM 4.4.1 (Post). *There is a finitely presented semigroup P such that the word problem for P is not computable.*

PROOF. Let T be a universal Turing machine with states $Q = \{q_0, \dots, q_{n_s}\}$, where q_0 is the halting state, and tape alphabet $S = \{\sigma_1, \dots, \sigma_{n_A}\}$. Consider the free semigroup F generated by $Q \cup S$. Within this semigroup computation states are represented by the words with exactly one letter from Q . We form the semigroup P with relations prescribing that two computation states are equal if T prescribes a transition from one to another, and that q_0 is equal to the identity.

The result follows from the fact that it is not possible to decide, from input n , whether a universal Turing machine will halt with an empty tape on input n . \square

To produce a *group* with unsolvable word problem, we must define the computation state words more carefully: a word is said to be *special* if it is of the form $\Sigma_1 q_i \Sigma_2$, where $\Sigma_1, \Sigma_2 \in S^{<\omega}$. The difficulty of groups is that with inverses, words may be equivalent in more subtle ways.

Novikov [389], and independently Boone [76, 77] demonstrated that there is indeed a finitely generated group with unsolvable word problem. The parts of the proof that concern us here are not the most difficult parts of the proof. The full algebraic details, including the proof of the critical lemma, can be found in Rotman’s book [422]. Since most of those details are not germane to the present work, the reader is referred to that treatment, while we give only a brief outline of the proof below.

THEOREM 4.4.2. *There is a group with unsolvable word problem.*

PROOF. Let S and Q be as in Post’s construction, and let $R = \{B_i = \Gamma_i\}$ be the set of relators in Post’s construction. Let G be generated by

$$S \cup Q \cup \{k, t, x, y\} \cup \{\ell_i : i \in R\} \cup \{r_i : i \in R\}$$

and defined by the following relations (for all values in the range of b and i):

$$\begin{aligned}
s_b y &= y y s_b \\
x s_b &= s_b x x \\
s_b \ell_i &= y \ell_i s_b \\
r_i s_b &= s_b x r_i x \\
B_i &= \ell_i \\
\Gamma_i &= r_i \\
t \ell_i &= \ell_i t \\
t y &= y t \\
r_i k &= k r_i \\
x k &= k x \\
(q^{-1} t q) k &= k (q^{-1} t q).
\end{aligned}$$

We note that all of the words of Post's semigroup are still words of G , and the definition of “special” words above is still meaningful. Again, it is the special words that will allow us to encode computation.

Let Δ be a special word in G which is equal in P to q_0 . Then we can show from the relators that in G , we have $\Delta = L q_0 R$, where L is a word in $\{y, \ell_i : i \in R\}$ and R is a word in $\{x, r_i : i \in R\}$. Then, by the commutativity relations of G , it follows that $(\Delta^{-1} t \Delta) k = k (\Delta^{-1} t \Delta)$. Through some involved group-theoretical arguments, it is possible to show that the converse also holds. At this point, we know that there can be no algorithm to decide, for any special word, whether $((\Delta^{-1} t \Delta) k) (k (\Delta^{-1} t \Delta))^{-1}$ is the identity. \square

4.4.2. Groups with Generically Solvable Word Problem. In some sense the proof of Theorem 4.4.2 is unsatisfying: While the result is, in fact, established, there is nothing to prevent the existence of an algorithm that would solve the word problem for the vast majority of words in the group. The words representing the obstruction are called “special” for a reason (considerably predating any of the results or considerations of the present section). They are, in fact, relatively rare.

In 2003, Kapovich, Myasnikov, Schupp, and Shpilrain formalized this dissatisfaction [288]. In particular, they showed that for large classes of groups — including the group constructed in Theorem 4.4.2, the word problem is almost solvable, in the sense that there is an algorithm (indeed, a linear time algorithm) that will solve almost all instances of the word problem.

To this point, we have not yet defined what “almost all” should mean in this context, and a reader who has kept in mind that all of the structures involved are countable is doubtless ready to see a definition.

DEFINITION 4.4.3. Let Σ be a finite alphabet, and let $(X^*)^k$ be, as usual, the set of k -tuples of words on X . Let B_n be the set of elements of $(X^*)^k$ where the sum of the lengths of the words involved is at most n , and $S \subseteq (X^*)^k$. Then the *asymptotic density* of S , denoted $\rho_a(S)$ is defined as

$$\limsup_{n \rightarrow \infty} \frac{|S \cap B_n|}{|B_n|}.$$

In this sense, “almost all” means on a set of density 1, or (equivalently) except for a set of density 0. Of course, there is still some flexibility in the definition, and we will see in the following section how this flexibility leads to several notions of effectiveness. For the present, however, we follow the original work.

DEFINITION 4.4.4. Let $P \subseteq (X^*)^k$. We say that P is *generically computable* if there is a Turing machine T with domain S such that $\rho_a(S) = 1$, and such that if $\sigma \in S$, we have $T(\sigma) = \chi_P(\sigma)$.

THEOREM 4.4.5. *Let G be a finitely generated group, such that G has a finite index subgroup with infinite quotient \bar{G} for which the word problem is solvable. Then the word problem for G is generically solvable.*

Before considering the proof of this theorem, we point out the following corollary, which clarifies the situation of the Boone group.

COROLLARY 4.4.6. *The group G constructed in Theorem 4.4.2 has generically solvable word problem.*

PROOF. Consider the subgroup \bar{G} of G generated by the set $\{r_i : i \in R\}$. Now this \bar{G} is a quotient of G , and is a non-Abelian free group on its generators. Since such a group has solvable word problem, the Corollary follows from the Theorem. \square

Actually, Theorem 4.4.5 follows from another technical result of the same paper. Given a group presentation and a subgroup H , the *membership problem* for H is that of determining whether a word is an element of H . Note that the word problem for a group is equivalent to the membership problem for its trivial subgroup.

PROPOSITION 4.4.7. *Let G be a finitely generated group and $H \leq G$ a finitely generated subgroup of infinite index. Let $G_1 \geq H$ be a finite-index subgroup of G , and $\phi : G_1 \rightarrow \bar{G}$ an epimorphism to some group \bar{G} . If $\bar{H} := \phi(H)$ is contained in an infinite index $\bar{K} \leq \bar{G}$ and the membership problem for \bar{K} is solvable, then the membership problem for H is generically solvable.*

Before entering the interesting details of Proposition 4.4.7, let us see how it implies Theorem 4.4.5.

PROOF OF THEOREM 4.4.5. Let G_1 be a finite index subgroup of G with infinite quotient \bar{G} , where \bar{G} has solvable word problem, with $\phi : G_1 \rightarrow \bar{G}$ the standard projection. Let H be the trivial subgroup of G and \bar{K} the trivial subgroup of \bar{G} . Then $\phi(H) \leq \bar{K}$. Note that \bar{K} has infinite index in \bar{G} , and that the word problem of \bar{G} is exactly the membership problem of \bar{K} , so that the membership problem of \bar{K} is, by assumption, solvable. By Proposition 4.4.7, the membership problem for H is generically solvable. But this problem is exactly the word problem for G . \square

We now undertake the proof of Proposition 4.4.7. As usual, the interested reader will want to follow up with the original paper, which has both stronger conclusions and more details of the proof.

PROOF OF PROPOSITION 4.4.7. Let A be a generating set of length k for G and let B be a finite generating set for G_1 . Let $\pi : F(A) \rightarrow G$ be the standard projection of the free group on A , denoted $F(A)$ to G . Let $K_1 := \phi^{-1}(\bar{K})$, and

$K_2 := \pi^{-1}(K_1)$. Note that \overline{K} has infinite index in \overline{G} by assumption, and K_1 has infinite index in G_1 . Consequently, K_2 has infinite index in $F(A)$.

We let z_n denote the number of words σ in $A \cup A^{-1}$ of length n , with $\pi(\sigma) \in K_2$, and note that there are

$$\frac{(2k)^{n+1} - 1}{2k - 1}$$

distinct words in $(A \cup A^{-1})$ of length n . Consequently, the asymptotic density of K_2 is given by

$$\lim_{n \rightarrow \infty} \frac{z_n}{\frac{(2k)^{n+1} - 1}{2k - 1}}.$$

By assumption, $H \leq K_1$, so that if σ is a word in $A \cup A^{-1}$, but not in K_2 , it follows that $\pi(w) \in G - H$. Consequently, if we can prove that the asymptotic density of K_2 is 0, the proof is done.

We define the *Cayley graph of $F(A)$* , as the graph whose vertices are the elements of $F(A)$, and where two vertices σ, τ are adjacent if and only if $\sigma = \tau x$ for some $x \in A$. In the standard graph metric, where adjacent vertices have distance 1, the ball of radius n about the origin contains exactly the words of length at most n .

LEMMA 4.4.8. *Let Γ be the Cayley graph of a group with d -generators, where $d \geq 3$. Let ζ_n denote the number of paths of length at most n from the identity to the identity. Then*

$$\lim_{n \rightarrow \infty} \frac{\zeta_n}{d^n} = 0.$$

PROOF. if b_n denotes the number of paths without backtracks of length at most n from the identity to the identity, and $\beta := \limsup_{n \rightarrow \infty} \sqrt[n]{b_n}$. It is known [48] that we can assume $\beta < d$, so that there is some $N_0 \geq 1$ and $0 < b < d$, so that

$$\lim_{n \rightarrow \infty} \frac{b_n}{d^n} \leq \frac{b^n}{d^n} \rightarrow 0.$$

The lemma follows, by some well-established limit comparison theorems. \square

It follows from the definition of the Cayley graph that $z_n = \zeta_n$. Consequently, by the lemma, K_2 has asymptotic density 0, as required. \square

The group theory is important in this theorem, and to generalize this notion of computability it is necessary to replace it. In particular, the notion of asymptotic density depended on properties of random words in a group, or equivalently, random walks in a graph. It will be the business of the next section to pose a definition of a notion of computation that carries the sense of a generically solvable word problem, but in a more general setting.

While we turn from the group theoretic origins of generic computability for the present, we will return to them. There is important work in this area on dynamics, to which we will return in Chapter 8, and random walks in a graph will be important when we take up the subject of expander graphs in Chapter 5.

4.5. Generic and Coarse Computability

4.5.1. Generic Computability. In 2012, Jockusch and Schupp [281] extended the theory of generic solvability to subsets of \mathbb{N} , the native realm of computability theory. By identifying $n \in \mathbb{N}$ with its unary representation, the definitions of the previous section reduce to something reasonable on the natural numbers.

DEFINITION 4.5.1. Let $S \subseteq \mathbb{N}$.

- (1) The *density of S up to n* , denoted $\rho_n(S)$ is given by

$$\frac{|S \cap \{0, 1, 2, \dots, n\}|}{n + 1}$$

- (2) The *density of S* , denoted $\rho(S)$ is given by

$$\lim_{n \rightarrow \infty} \rho_n(S).$$

Under the interpretation of natural numbers by their unary representations, this definition coincides exactly with the one in Section 4.4.2. We have the corresponding definition.

DEFINITION 4.5.2. *generically computable*—textbf Let $P \subseteq \mathbb{N}$. We say that P is generically computable if and only if there is a Turing machine T such that $T(n) = \chi_P(n)$ wherever T is defined, and T is defined on a set of density 1.

Jockusch and Schupp made several initial observations about these sets.

PROPOSITION 4.5.3. *Every Turing degree contains a set which is generically computable.*

PROOF. Let \mathbf{d} be a Turing degree, with $A \in \mathbf{d}$. We set $B = \{2^m : m \in A\}$. Note that $B \in \mathbf{d}$, since the set of natural numbers which are not powers of 2 is computable. We define a Turing machine T so that

$$T(n) = \begin{cases} 1 & \text{if } n = 2^k \text{ for some } k \\ \uparrow & \text{otherwise} \end{cases}$$

Now $T(n) = \chi_B(n)$ for all $n \in \text{dom}(T)$, and $\text{dom}(T)$ has density 1. \square

Even after the structural evidence of the generically solvable word problem, it is still natural, after Proposition 4.5.3 to wonder how nearly trivial the concept of generic computability could be.

PROPOSITION 4.5.4. *Every nonzero Turing degree contains a set which is not generically computable.*

PROOF. Let $\mathbf{d} >_T \emptyset$, and let $A \in \mathbf{d}$. Let

$$R_k = \{m : 2^k | m \text{ and } 2^{k+1} \nmid m\}.$$

We write

$$\mathcal{R}(A) = \bigcup_{k \in A} R_k.$$

Now if T were a Turing machine witnessing that $\mathcal{R}(A)$ is generically computable, we could decide whether $n \in \mathcal{R}(A)$ by running T on all multiples of 2^n until it answers on one. We know that T must answer on some element of R_n , since this set has density $2^{-k} > 0$. Since $\mathcal{R}(A) \equiv A$, the result follows. \square

Recall that a set is said to be *bi-immune* if both it and its complement are infinite, with no infinite computably enumerable subsets.

PROPOSITION 4.5.5. *No bi-immune set is generically computable.*

PROOF. Suppose that Turing machine T witnesses that A is generically computable. Let C_0 be the set of n such that $T(n) = 0$ and C_1 the set of n such that $T(n) = 1$. Now $C_0 \subseteq \bar{A}$ and $C_1 \subseteq A$, and $C_0 \cup C_1$ has density 1. Consequently, C_0 and C_1 cannot both be finite, so that A cannot be bi-immune. \square

Of course, probabilities other than 1 are still interesting, although they do depend more sensitively on the distribution of possible inputs. Since we, by definition, have fixed a distribution (the uniform distribution), it is possible to talk about these sets. Downey, Jockusch, and Schupp gave the following definition, which extends generic computability in this natural way.

DEFINITION 4.5.6. Let $S \subseteq \mathbb{N}$. We define the lower density of S , denoted $\rho_\ell(S)$ as

$$\liminf_{n \rightarrow \infty} \frac{|S \cap \{0, 1, \dots, n\}|}{n + 1}.$$

We then say, for any $r \in \mathbb{R}$, that a set $A \subseteq \mathbb{N}$ is partially computable at density r if there is a Turing machine T such that $T(n) = A(n)$ for all n in the domain of T , and such that the domain of T has lower density greater than or equal to r .

In the same paper, several observations were made about this definition. To begin with, this definition differs significantly from the definition of generic computability.

PROPOSITION 4.5.7. *Every nonzero Turing degree contains a set which is partially computable exactly at the densities strictly less than 1.*

PROOF. Let the sets R_k and the operator \mathcal{R} be as in the proof of Proposition 4.5.4. Let \mathbf{d} be a nonzero Turing degree, with $A \in \mathbf{d}$. Let $\epsilon > 0$, and note that $\mathcal{R}(A)$ is partially computable at density $1 - \epsilon$, for a Turing machine witnessing its partial computability need not have in its domain any element of R_k for sufficiently large k . As we observed, however, $\mathcal{R}(A)$ cannot be partially computable at density 1. \square

In this sense, it is easier to believe in a set which is partially computable at all densities less than 1 than it is to believe in one which is partially computable at some proper subset of those densities. The following result shows that such sets do exist.

THEOREM 4.5.8. *Let $r \in [0, 1]$. Then there is a set A which is partially computable at density r , but not at any higher density, if and only if r is left- Σ_3^0 .*

PROOF. Let $r \in [0, 1]$ be left- Σ_3^0 . We first produce a computably enumerable set C with lower density r and a simple set S of density 0.

Toward the first goal, we first produce a Δ_2^0 set \tilde{C} with lower density r . We construct a Δ_2^0 -computable sequence $(q_s : s \in \mathbb{N})$ where $\liminf_{s \rightarrow \infty} q_s = r$. Let A be the set of rationals less than r , a Σ_3^0 set. We can then construct a Δ_2^0 sequence $(A_s : s \in \mathbb{N})$ of finite sets such that for all $x \in A$, we have $x \in A_s$ for all sufficiently large s , such that there are infinitely many s such that $A_s \subseteq A$, and such that

$0 \in A_s$ for all s . We set $q_s := \max A_s$. Now we will inductively define $\tilde{C} = \bigcup_{i \in \mathbb{N}}$ and a sequence $(s_i : i \in \mathbb{N})$. At stage 0, we set $s_0 = 1$ and $\Gamma_0 = \{0\}$. At stage t , we check whether

$$\frac{|\Gamma_{t-1}|}{s_{t-1} + 1} < q_t.$$

If so, we find the least k such that

$$\frac{|\Gamma_{t-1}| + k}{s_{t-1} + 1 + k} \geq q_t$$

and set $s_{t+1} = s_t + k$ and $\Gamma_t = \Gamma_{t-1} \cup \{s_t + 1, \dots, s_t + k\}$. Otherwise, we find the least k such that

$$\frac{|\Gamma_{t-1}|}{s_{t-1} + 1 + k} < q_t$$

and set $s_{t+1} = s_t + k$ and $\Gamma_{t+1} = \Gamma_t$. All of these operations can be completed with a Δ_2^0 oracle, so that \tilde{C} is Δ_2^0 , as required.

Now let \tilde{C}_s be a computable approximation to \tilde{C} . At stage 0, set $C_0 = \emptyset$ and $t(n, 0) = n + 1$, and $t(-1, n) = -1$ for all n . At stage $s + 1$, we let n_s be the least $n \leq s$, if any, such that $\rho_{t(n,s)}(\tilde{C}_s) \neq \rho_n(C_s)$. We then find a finite set $C_{s+1} \supseteq C_s$ and a number \hat{t} greater than any number in C_s and greater than $t(n, s)$ such that $\rho_c(C_{s+1}) = \rho_n(C_s)$, with the additional requirements that C_{s+1} agrees with C_s between n_s and s , and that C_{s+1} includes an initial segment of $[t(n, s), \infty)$. It can be shown that for each $k > 0$ there are only finitely many s where $n_s = k$, and that $\rho_{\lim_{s \rightarrow \infty} t(k,s)}(C_s) = \rho_k(\tilde{C})$. We set $C = \bigcup_{s \in \mathbb{N}} C_s$, so that C is a computably enumerable set of lower density r .

We also produce a simple set S of density 0. Indeed, for each e , we look for the first $n > e^2$ with $n \in W_e$. If such an e exists, we include it in S . Now any computably enumerable set disjoint from S must be finite, but S contains at most e elements less than e^2 for each e , so that

$$\rho(S) \leq \lim_{n \rightarrow \infty} \frac{n}{n^2} = 0.$$

Let $A = C \cup S$. Then A is computable at density r , because $C \subseteq A$. However, if $r' > r$ and T witnesses that A is partially computable at density r' , then the set $\{n : T(n) = 0\}$ must be disjoint from S (since $S \subseteq A$), infinite (since density $r' - r > 0$ of the domain of T must be outside A), and computably enumerable. Since S is simple, this is impossible.

On the other hand, let r fail to be left- Σ_3^0 , and let T witness that A is partially computable at density r . Suppose T has domain D . From the definitions, it is clear that $r' := \rho_\ell(D) = \liminf_{n \rightarrow \infty} q_n$, where $(q_n : n \in \mathbb{N})$ is a Δ_2^0 sequence of rationals. If r' is irrational, then for any rational q , and for all but finitely many of the q_n , we have $q < q_n$, so that r' is left- Σ_3^0 . Since $r' \geq r$, we know that $r' > r$, with A partially computable at density r' . \square

It is worth noting the similarities between the construction of \tilde{C} in the last proof and the proof of Theorem 4.2.12. In both cases, the construction proceeds by recruiting witnesses to the necessary case to establish a needed probability.

DEFINITION 4.5.9. Let $A \subseteq \mathbb{N}$. Then the *partial computability bound* of A , denoted $\alpha(A)$, is the supremum of all r such that A is computable at density r .

PROPOSITION 4.5.10. *Let $r \in (0, 1)$. Then there is a set with partial computability bound r .*

PROOF. Let $r = \sum_{i=1}^{\infty} b_i 2^{-i}$ where $b_i \in \{0, 1\}$ and infinitely many of the b_i are

1. Let S be the simple set of density 0 constructed above, and let

$$R_k = \{m : 2^k | m \text{ and } 2^{k+1} \nmid m\}$$

as before. We set $D = \bigcup_{b_i=1} R_i$, and set $A = D \cup S$. As before, A is partially computable at any density strictly less than r , but not at any density greater than r . \square

4.5.2. Coarse Computability. Jockusch and Schupp [281] identified and studied a second notion of approximate computability. As we suggested earlier, the key difference is in what it means — even after agreeing on the meaning of “almost all” — for an algorithm to solve almost all instances of the problem. Of course, all correct solutions are alike. The real question concerns what the algorithm does on the places where it does not give a correct solution. In generic computability, the algorithm must be correct everywhere it halts: it can either be correct or not halt at all.

In this second notion, however, the algorithm is required to halt on all inputs, and be correct almost everywhere.

DEFINITION 4.5.11. Let $P \subseteq \mathbb{N}$. We say that P is coarsely computable if there is a Turing machine T which is defined on all natural numbers, and such that for some set $S \subseteq \mathbb{N}$ of density one, we have $T(n) = \chi_P(n)$ for all $n \in S$.

Jockusch and Schupp point out that this notion says something important about the initial issue of word problems for groups. Indeed, the following result, in light of the proofs of the previous section, suggests that the “coarse” in the definition is very coarse indeed. Importantly, we need not know *where* the machine goes wrong, only that it do so very seldom.

PROPOSITION 4.5.12. *Every finitely generated group has coarsely computable word problem.*

PROOF. If G is a finitely generated infinite group, then the set of words not equal to the identity has density 1, so that a Turing machine rejecting every input coarsely computes the word problem. If G is finite, then its word problem is computable. \square

In view of the near-triviality of the preceding proof, it is tempting to think that coarse computability would be a notion permissive to the point of uselessness. However, a recent paper of Greenberg, J. Miller, Shen, and Westrick [231] demonstrates that its approach via the density of the points of agreement is exactly what is needed to settle an important question in effective dimension and algorithmic randomness. We say that two sets are *coarsely equivalent* if they agree on a set of density 1. We first note a preliminary result.

PROPOSITION 4.5.13. *If two sequences are coarsely equivalent, they have the same effective Hausdorff dimension.*

THEOREM 4.5.14 ([231]). *A sequence has effective Hausdorff dimension 1 if and only if it is coarsely equivalent to a 1-random sequence.*

PROOF. Let X be 1-random. By Proposition 3.4.7, it has effective Hausdorff dimension 1, and by Proposition 4.5.13, any Y coarsely equivalent to X must have the same effective Hausdorff dimension. The second direction is more involved.

Assume X has effective Hausdorff dimension 1, and let

$$s_m = \dim(X \upharpoonright_{[2^m, 2^{m+1})} | X \upharpoonright_{2^m}).$$

Let

$$P = \{Y : \forall n \ K(Y \upharpoonright_n) \geq n\},$$

and let \mathbb{P} denote the set of finite initial segments of elements of P . Let d denote the Hamming distance, and let E be the set of all sequences $(e_m : m \in \mathbb{N})$ such that $e_0 = 1$ and such that for all k either $e_k \in \{e_{k-1}, \frac{e_{k-1}}{2}\}$. For each $\bar{e} \in E$, we define a relation $\sim_{\bar{e}}$ on strings so that $\sigma \sim_{\bar{e}} \tau$ if and only if for all k we have

$$d(\sigma \upharpoonright_{[2^{k-1}, 2^k)}, \tau \upharpoonright_{[2^{k-1}, 2^k)}) \leq e_k.$$

We now define a sequence $(\epsilon_m : m \in \mathbb{N}) \in E$ such that for all m we have

- (1) $\lim_{m \rightarrow \infty} \epsilon_m = 0$, and
- (2) There is some $\tau \in \mathbb{P}$ of length 2^m with $\tau \sim_{\epsilon_0, \dots, \epsilon_m} X \upharpoonright_{2^m}$.

We claim that for each m there is $\nu \in \mathbb{P}$ of length 2^{m+1} such that

$$\nu \sim_{(\epsilon_0, \dots, \epsilon_m)} X \upharpoonright_{2^{m+1}}.$$

To this end, let A be the set of strings η of length 2^m such that for some $\hat{\tau} \in \mathbb{P}$ we have

$$\hat{\tau} \sim_{(\epsilon_0, \dots, \epsilon_{m-1})} X \upharpoonright_{2^m}$$

and $\hat{\tau}\eta \in \mathbb{P}$. Since \mathbb{P} is co-c.e., it follows that A is co-c.e. It can be shown that there is some $q < 1$ such that if there are at most 2^{q2^m} strings π whose distance from A is greater than ϵ_m . Call the computably enumerable set of all such strings B . Then, conditioned on $X \upharpoonright_{2^m}$, each $\pi \in B$ has a description of length $q2^m + m + O(1)$. Consequently, for sufficiently large m , every $\pi \in B$ will satisfy

$$\dim(\pi | X \upharpoonright_2^m) < s_m$$

so that $X \upharpoonright_{[2^m, 2^{m+1})} \notin B$. In that case, there must be some $\eta \in A$ such that

$$d(\eta, X \upharpoonright_{[2^m, 2^{m+1})}) \leq \epsilon_m.$$

Let $\hat{\tau}$ witness that $\eta \in A$, and take $\nu = \hat{\tau}\eta$.

We now take the set Q_m of all such strings $\nu \in \mathbb{P}$ of length 2^{m+1} such that

$$\nu \sim_{(\epsilon_0, \dots, \epsilon_m)} X \upharpoonright_{2^{m+1}}.$$

We have shown that these sets are nonempty, and every element of Q_m has a prefix from Q_{m-1} . By compactness, there is $Y \in P$ such that for all m , we have $Y \upharpoonright_{2^m} \in Q_m$. Since the differences between X and Y on a sequence of intervals whose length grows exponentially tends to 0, we have Y coarsely equivalent to X . \square

Again, as in the case of generic computability, one can loosen the requirement of density 1 in the definition. Although the idea was implicit in [159], this was first done explicitly in [256].

DEFINITION 4.5.15. Let $A \subseteq \mathbb{N}$, and $r \in [0, 1]$. We say that A is coarsely computable at density r if there is a computable set B such that the lower density of the set of all n on which A and B agree is at least r .

DEFINITION 4.5.16. Let $A \subseteq \mathbb{N}$. Then the *coarse computability bound* of A , denoted $\gamma(A)$, is the supremum of all r such that A is coarsely computable at density r .

As we will see in Theorem 4.5.20, there are sets A with $\gamma(A) = 1$ which are nevertheless not coarsely computable.

4.5.3. Relationships between Generic and Coarse. It is natural to think that generic and coarse computability might be closely related, although intuition, by its nature, may vary from person to person. In fact, the two are nearly orthogonal. The first several results we consider in this area (as well as stronger forms of some of them) are from [281].

PROPOSITION 4.5.17. *There is a computably enumerable set which is coarsely computable but not generically computable.*

PROOF. Consider the simple set S of density 0 constructed in the proof of Theorem 4.5.8. Since S has density 0, it is coarsely computable. However, if machine T witnesses that S is generically computable, take $C_i = \{n : T(n) = i\}$, as before, and note that $C_0 \cup C_1$ has density 1. Since C_1 is contained in a set of density 0, it must also have density 0, leaving C_0 an infinite computably enumerable subset of the complement of S . \square

PROPOSITION 4.5.18. *There is a generically computable computably enumerable set which is not coarsely computable.*

PROOF. If A_0, A_1 are disjoint computably enumerable sets whose union has density 1, both A_0 and A_1 are generically computable; indeed, a Turing machine can enumerate each set, and can give output exactly on $A_0 \cup A_1$. If we can construct such sets where A_1 is not coarsely computable, then the theorem will hold.

We recall the definition of the sets R_e from Proposition 4.5.4, and, as usual, let W_e denote the domain of machine T_e . We initialize $A_{0,0} = A_{1,0} = \emptyset$ and $r(e, 0) = \min(R_e)$. At stage s , for each $e \leq s$, we check whether $R_e \upharpoonright_{r(e,s)} \subseteq W_{e,s+1}$.

If so, then we let

$$F = R_e \upharpoonright_{r(e,s)} - (A_{0,s} \cup A_{1,s}).$$

For each $n \in F$, we include n in $A_{0,s+1}$ if $T_e(n) = 1$, and in $A_{1,s+1}$ otherwise. We set $r(e, s+1)$ so that at most half of the elements of $R_e \upharpoonright_{r(e,s+1)}$ are in $A_{0,s+1} \cup A_{1,s+1}$. Consequently, T_e and $A_{1,s+1}$ will differ on every element of F .

Otherwise, we find the least z in R_e which is greater than $r(e, s)$ with $z \notin A_{1,s}$. We then set $A_{1,s+1} = A_{1,s} \cup \{z\}$ and leave $A_{0,s+1} = A_{0,s}$ and $r(e, s+1) = r(e, s)$.

Let $A_i = \bigcup_{j \in \mathbb{N}} A_{i,j}$. Now if T_e is total, then we have constructed a set of positive measure on which T_e and A_1 disagree, so that A_1 cannot be coarsely computable by any T_e . On the other hand, for each e , we will have almost every element of R_e enumerated into either A_0 or A_1 at some stage. Consequently, $A_0 \cup A_1$ has density 1, as required. \square

PROPOSITION 4.5.19. *There is a computably enumerable set which is neither coarsely computable nor generically computable.*

PROOF. Recall from Proposition 4.5.4 the definition of the sets R_e , as before. Now we define

$$A = \bigcup_{e \in \mathbb{N}} (W_e \cap R_e).$$

Now for each e , we know that A differs from the complement of W_e on all of R_e . Consequently, A cannot be coarsely computable, since if T witnesses that A is coarsely computable, the set $N = \{n : T(n) = 0\}$ is computably enumerable, and A can differ from the complement of this set only with density 0.

Now, toward contradiction, let T witness that A is generically computable. Let $C_i = \{n : T(n) = i\}$, and note, as usual, that C_0 is contained in the complement of A , that C_1 is contained in A , and that $C_0 \cup C_1$ has density 1. Then A can differ from the complement of C_0 only on elements outside $C_0 \cup C_1$, a set of density 0. However, C_0 is computably enumerable, and A must differ from it on a set of positive density. \square

In view of these limitations, the following results on partial and coarse computability bounds, from a paper of Hirschfeldt, Jockusch, McNicholl, and Schupp [256], seem striking.

THEOREM 4.5.20. *For any $A \subseteq \mathbb{N}$, we have $\alpha(A) \leq \gamma(A)$.*

PROOF. Let $\epsilon > 0$, and let T be a partial Turing machine such that T agrees with A on its domain, and such that the domain of T has density $\alpha(A) - \frac{\epsilon}{2}$. We will find a computable computable C within the domain of T with lower density at most $\frac{\epsilon}{2}$ less than the lower density δ of the domain of T . Given such a C , we can define a Turing machine U to accept those n for which $n \in C$ and $T(n) = 1$, and to reject all others. Now U agrees with T on all of C , but U is total, so that A is coarsely computable at density $\alpha(A) - \epsilon$.

We now construct C . Let q be a rational between $\delta - \frac{\epsilon}{2}$ and δ . There is some n_0 so that the density of the domain of T up to n is at least q for all $n > n_0$. Now we define C to be the subset of the domain of T consisting of all elements k less than the maximum over all n between n_0 and k^2 of the least s such that the density up to n of the domain of T_s is greater than q . This set is computable. On the other hand, $\rho_n(B) \geq q - \frac{1}{n}$, so that the lower density of C is at least $\delta - \frac{\epsilon}{2}$. \square

This inequality gives us, as a consequence, a set which has coarse computability bound 1, but which is not coarsely computable, since we know that there is a set which is generically computable but not coarsely computable. The two bounds need not be close.

PROPOSITION 4.5.21. *There is a set whose partial computability bound is 0, but whose coarse computability bound is $\frac{1}{2}$.*

PROOF. Let $I_n = [n!, (n+1)!)$, and define $\mathcal{I}(A) = \bigcup_{n \in A} I_n$. Now if T witnesses that $\mathcal{I}(A)$ is coarsely computable at density greater than $\frac{1}{2}$, then we can compute A in the following way: for some n_0 and for all $n > n_0$, the machine T must agree with $\mathcal{I}(A)$ on more than half of the elements of I_n . In that case, a majority vote decides whether $n \in A$. On the other hand, $\mathcal{I}(A) \equiv_T A$, so $\mathcal{I}(A)$ is coarsely computable just in case A was computable.

On the other hand, any set of positive lower density must intersect almost all of the sets I_n , so that if $\mathcal{I}(A)$ is partially computable at density $r > 0$, we again have A computable. \square

Moreover, below either a 1-generic or a weak 2-random, the inequality of Theorem 4.5.20 becomes equality, so that all generically computable sets become partially computable. More exactly,

THEOREM 4.5.22. *Let $A \subseteq \mathbb{N}$, and $r \in [0, 1]$. Let $B \leq_T A$. If A is either 1-generic or weakly 2-random, then if B is partially computable at density r , then B is coarsely computable at density r .*

PROOF. We first let $B = \Phi^A$, with T witnessing that B is partially computable at density r .

Consider first the case that A is 1-generic. We define a notion of forcing

$$P = \{\sigma \in 2^{<\omega} : \Phi^\sigma \perp T\}.$$

Note that P is computably enumerable. If there is n such that $A \upharpoonright_n \in P$, then B disagrees with T somewhere. Otherwise, since A is 1-generic, we have some n such that all τ extending $A \upharpoonright_n$ are outside P . In that case, we will define a total Turing machine U that agrees with B on a set of density at least r .

To define the behavior of U on input m , find $\sigma \supseteq A \upharpoonright_n$ such that $\Phi^\sigma(m) \downarrow$, and set $U(m) = \Phi^\sigma(m)$. Since $A \supseteq A \upharpoonright_n$, some such σ can always be found. However, since σ was not in S , we know that $U(m)$ must be compatible with B , so that U agrees with B throughout the domain of T , which has density at least r .

Now if A is weakly 2-random, we define a Π_2^0 class P so that $A \in A$. We set

$$P = \{X : \Phi^X \text{ is total and compatible with } T\}.$$

Since A is weakly 2-random and included in P , we know that P must have positive measure. The Lebesgue density theorem provides that there is some string γ such that P includes at least relative measure .8 of the extensions of γ , so that for each n there is some $i_n \in \{0, 1\}$ so that

$$\mu(\{X : \Phi^X(n) = i_n\}) \geq .4.$$

Then we define U so that $U(n) = i_n$. Now U is a total Turing machine that agrees with B throughout the domain of T , so that B is coarsely computable at density r . \square

4.5.4. Beginnings of Generically and Coarsely Computable Structure

Theory. The origin of dense computability is in structure theory; in particular, in word problems for groups. It is natural, then, that its further application to structure theory be explored. This study is ongoing, but some recent work by the present author with Cenzer, Harizanov, and Gonzalez shows some promise in the field.

There is an initial issue of how to handle the dense computability of a set with arity greater than one, which are necessary to represent structures with functions, or with binary relations. The following result represents straightforward calculations.

- LEMMA 4.5.23 ([96]).** (1) *Let $A \subseteq \omega$. Then A has asymptotic density δ if and only if A^r has asymptotic density δ^r in ω^r . In particular, A is asymptotically dense in ω iff A^r is asymptotically dense in ω^r .*
- (2) *There is a computable dense $C \subseteq \omega^2$ such that for any infinite c.e. set $A \subseteq \omega$, the product $A \times A$ is not a subset of C .*

These observations inform the definition of densely computable structures. In particular, we define them by density in the (one-dimensional) domain.

- DEFINITION 4.5.24 ([96]).** (1) A structure \mathcal{A} is said to be *generically computable* if and only if there is a substructure \mathcal{D} whose universe is a c.e. dense set D , where for every k -ary function f and every k -ary relation R ,

both $f \upharpoonright_{D^k}$ and $\chi_R \upharpoonright_{D^k}$ are restrictions to D^k of some partial computable function.

- (2) A structure \mathcal{A} is said to be *coarsely computable* if and only if there are a computable structure \mathcal{E} and a dense set D such that the structure \mathcal{D} with universe D is a substructure of both \mathcal{A} and \mathcal{E} , and all relations and functions of \mathcal{A} and \mathcal{E} agree on D .

The first observation about these notions is concerning.

THEOREM 4.5.25 ([96]). *Every countable equivalence structure has a generically computable copy.*

The proof of this theorem arises from the fact that it is only necessary for the structure to have an infinite substructure simple enough to have a computable copy. An early approach to rectifying this was to demand that D be closed under equivalence, but this eventually gave way to the much more productive application of graduated elementarity conditions.

DEFINITION 4.5.26. Let $n \geq 0$, and \mathcal{A} a structure. Then a substructure \mathcal{B} is a Σ_n elementary substructure of \mathcal{A} if and only if for any infinitary Σ_n formula $\varphi(\bar{x})$ and any $\bar{b} \in \mathcal{B}$, we have

$$\mathcal{A} \models \varphi(\bar{b}) \Leftrightarrow \mathcal{B} \models \varphi(\bar{b}).$$

We can then say that \mathcal{A} is Σ_n -generically c.e. if and only if the subset D can be chosen to be Σ_n elementary. At this point the trivialization disappears, and one frequently (now across several classes of structures) sees a progression from all countable structures having generically computable copies to some α at which exactly the structures with computable copies have Σ_α -generically c.e. copies. Similar situations hold for coarsely computable structures. Ongoing work in this area, including densely computable morphisms and application to several classes of structures, can be found in [96, 97, 98, 95].

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