Smoothed Complexity Theory

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Abstract. Smoothed analysis is a new way of analyzing algorithms introduced by Spielman and Teng (J. ACM, 2004). Classical methods like worst-case or average-case analysis have accompanying complexity classes, like P and Avg-P, respectively. While worst-case or average-case analysis give us a means to talk about the running time of a particular algorithm, complexity classes allows us to talk about the inherent difficulty of problems.

Smoothed analysis is a hybrid of worst-case and average-case analysis and compensates some of their drawbacks. Despite its success for the analysis of single algorithms and problems, there is no embedding of smoothed analysis into computational complexity, which is necessary to classify problems according to their intrinsic difficulty.

We propose a framework for smoothed complexity theory, define the relevant classes, prove some first results, and study some examples.

1 Introduction

Computational complexity theory is the science of classifying computational problems according to their intrinsic difficulty. While the analysis of algorithms is concerned with analyzing, say, the running time of a particular algorithm, complexity theory rather analyses the amount of resources that all algorithms need at least to solve a given problem.

Classical complexity classes, like P, reflect worst-case analysis of algorithms. A problem is in P if there is an algorithm whose running time on all inputs of length n is bounded by a polynomial in n. Worst-case analysis has been a success story: The bounds obtained are valid for every input of a given size, and, thus, we do not have to think about typical instances of our problem. If an algorithm has a good worst-case upper bound, then this is a very strong statement: The algorithm will perform well in practice. (For practical purposes, "good upper bound" of course also includes constants and lower order terms.)

However, some algorithms – like quicksort or the simplex algorithm for linear programming – work well in practice, despite having a provably high worst-case running time. The reason for this is that the worst-case running time can be dominated by a few pathological instances that rarely or never occur in practice. An alternative to worst-case analysis is average-case analysis. Many of the algorithms with poor worst-case but good practical performance have a good average running time. This means that the expected running time with instances drawn according to some fixed probability distribution is low.

In complexity-theoretic terms, P is the class of all problems that can be solved with polynomial worst-case running time. In the same way, the class Avg-P is the class of all problems that have polynomial average-case running time. Average-case complexity theory studies the structural properties of average-case running time. Bogdanov and Trevisan give a comprehensive survey of averagecase complexity [6].

While worst-case complexity has the drawback of being often pessimistic, the drawback of average-case analysis is that random instances have often very special properties with high probability. These properties of random instances distinguish them from typical instances. Since a random and a typical instance is not the same, a good average-case running time does not necessarily explain a good performance in practice. In order to get a more realistic performance measure, Spielman and Teng have proposed a new way to analyze algorithms called smoothed analysis [21]. In smoothed analysis, an adversary chooses an instance, and then this instance is subjected to a slight random perturbation. We can think of this perturbation as modeling measurement errors or random noise or the randomness introduced by taking, say, a random poll. The perturbation is controlled by some parameter ϕ , called the *perturbation parameter*. Spielman and Teng have proved that the simplex method has a running time that is polynomial in the size of the instance and the perturbation parameter [21]. (More precisely, for any given instance, the expected running time on the perturbed instance is bounded by a polynomial.) Since then, the framework of smoothed analysis has been applied successfully to a variety of algorithms. We refer to a recent survey for a broader picture of smoothed analysis [22]

However, with only few exceptions [3, 20], smoothed analysis has only been applied yet to single algorithms or single problems. Up to our knowledge, there is currently no attempt to formulate a smoothed complexity theory and, thus, to embed smoothed analysis into computational complexity.

This paper is an attempt to define a smoothed complexity theory, including notions of intractability, reducibility, and completeness. We define the class Smoothed-P (Section 2), which corresponds to problems that can be solved smoothed efficiently, we provide a notion of reducibility (Section 3), and define the class Dist-NP_{para}, which is a smoothed analogue of NP, and prove that it contains complete problems (Section 4). We continue with some basic observations (Section 5). We also add examples of problems in Smoothed-P (Sections 6 and 7) and discuss the relationship of smoothed complexity to semi-random models (Section 8). Finally, since this is an attempt of a smoothed complexity theory, we conclude with a discussion of extension, shortcomings, and difficulties of our definitions (Section 9).

2 Smoothed Polynomial Time and Smoothed-P

2.1 Basic Definitions

In the first application of smoothed analysis to the simplex method [21], the strength of the perturbation has been controlled in terms of the standard de-

viation σ of the Gaussian perturbation. While this makes sense for numerical problems, this model cannot be used for general (discrete problems). A more general form of perturbation models has been introduced by Beier and Vöcking [2]: Instead of specifying an instance that is afterwards perturbed (which can also be viewed as the adversary specifying the mean of the probability distribution according to which the instances are drawn), the adversary specifies the whole probability distribution. Now the role of the standard deviation σ is taken over by the parameter ϕ , which is an upper bound for the maximum density of the probability distributions. For Gaussian perturbation, we have $\sigma = \Theta(1/\phi)$. Because we do not want to restrict our theory to numerical problems, we have decided to use the latter model.

Let us now define our model formally. Our perturbation models are families of distributions $\mathcal{D} = (D_{n,x,\phi})$. The length of x is n (so we could omit the index n but we keep it for clarity). Note that length does not necessarily mean bit length, but depends on the problem. For instance, it can be the number of vertices of the graph encoded by x. For every n, x, and ϕ , the support of the distribution $D_{n,x,\phi}$ should be contained in the set $\{0,1\}^{\leq \text{poly}(n)}$. Let $S_{n,x} = \{y \mid D_{n,x,\phi}(y) > 0 \text{ for some } \phi\}$, and let $N_{n,x} = |S_{n,x}|$.

For all n, x, ϕ , and y, we demand $D_{n,x,\phi}(y) \leq \phi$. This controls the strength of the perturbation and restricts the adversary. We allow $\phi \in [1/N_{n,x}, 1]$. Furthermore, the values of ϕ are discretized, so that they can be described by at most poly(n) bits. $\phi = 1$ corresponds to the worst-case complexity; we can put all the mass on one string. $\phi = 1/N_{n,x}$ models the average case; here we usually have to put probability on an exponentially large set of strings. In general, the larger ϕ , the more powerful the adversary. We call such families $(D_{n,x,\phi})_{n,x,\phi}$ of probability distributions parameterized families of distributions.

Now we can specify what it means that an algorithm has smoothed polynomial running-time. The following definition can also be viewed as a discretized version of Beier and Vöcking's definition [3]. Note that we do not speak about expected running-time, but over expected running-time to some power ε . This is because the notion of expected running-time is not robust with respect to, e.g., quadratic slowdown. The corresponding definition for average-case complexity is due to Levin [16]. We refer to Bogdanov and Trevisan [6] for a thorough discussion of this issue.

Definition 2.1. An algorithm A has smoothed polynomial running time with respect to the family \mathcal{D} if there exists an $\varepsilon > 0$ such that, for all n, x, and ϕ , we have $\mathbb{E}_{y \sim D_{n,x,\phi}}(t_A(y; n, \phi)^{\varepsilon}) = O(n \cdot N_{n,x} \cdot \phi).$

This definition implies that (average-)polynomial time is only required if we have $\phi = O(\text{poly}(n)/N_{n,x})$. This seems to be quite generous at first glance, but it is in accordance with, e.g., Spielman and Teng's analysis of the simplex method [21] or Beier and Vöcking's analysis of integer programs [3]; they achieve polynomial time running time only if they perturb all but at most $O(\log n)$ digits: If we perturb a number with, say, a Gaussian of standard deviation $\sigma = 1/\operatorname{poly}(n)$, then we expect that the $O(\log n)$ most significant bits remain untouched, but the less significant bits are random.

In average-case complexity, one considers not decision problems alone, but decision problems together with a probability distribution. The smoothed analogue of this are tuples (L, \mathcal{D}) , where $L \subseteq \{0, 1\}^*$ is a decision problem and \mathcal{D} is a parameterized family of distributions. We call such problems parameterized distributional problems. As some examples for distributional problems, we will study the bounded halting problem, binary integer optimization, and graph problems under the smoothed extension of $G_{n,p}$. The notion of smoothed polynomial running-time (Definition 2.1) allows us to define what it means for a parameterized distributional problem to have polynomial smoothed complexity.

Definition 2.2. Smoothed-P is the class of all (L, D) such that there is a deterministic algorithm A with smoothed polynomial running time that decides L.

As an alternative, we can characterize smoothed polynomial running-time in terms of polynomial tail bounds.

Theorem 2.3. An algorithm A has smoothed polynomial running time if and only if there is an $\varepsilon > 0$ and a polynomial p such that for all n, x, ϕ , and t, $\Pr_{y \sim D_{n,x,\phi}}[t_A(y; n, \phi) \ge t] \le \frac{p(n)}{t^{\varepsilon}} \cdot N_{n,x} \cdot \phi.$

2.2 Heuristic Schemes

A different way to think about efficiency in the smoothed setting is via so-called heuristic schemes. This notion comes from average-case complexity [6], but can be adapted to our smoothed setting. The notion of a heuristic scheme comes from the observation that, in practice, we might only be able to run our algorithm for a polynomial number of steps. If the algorithms does not succeed within this time bound, then it "fails", i.e., it does not solve the given instance. The failure probability decreases polynomially with the running time that we allow. The following definition captures this.

Definition 2.4. Let (L, D) be a smoothed distributional problem. An algorithm A is an errorless heuristic scheme for (L, D) if there is a polynomial q such that

- 1. For every n, every x, every ϕ , every $\delta > 0$, and every $y \in \operatorname{supp} D_{n,x,\phi}$, we have $A(y; n, \phi, \delta)$ outputs either L(y) or \bot .
- 2. For every n, every x, every ϕ , every $\delta > 0$, and every $y \in \operatorname{supp} D_{n,x,\phi}$, we have $t_A(y; n, \delta) \leq q(n, N_{n,x}\phi, 1/\delta)$.
- 3. For every $n, x, \phi, \delta > 0$, and $y \in \operatorname{supp} D_{n,x,\phi}$, $\operatorname{Pr}_{y \sim D_{n,x,\phi}}[A(y; n, \phi, \delta) = \bot] \leq \delta$.

Theorem 2.5. $(L, D) \in$ Smoothed-P if and only if (L, D) has an errorless heuristic scheme.

2.3 Alternative Definition: Bounded Moments

At first glance, one might be tempted to use "expected running time" for the definition of Avg-P and Smoothed-P. However, as mentioned above, simply using

the expected running time does not yield a robust measure. This is the reason why the expected value of the running time raised to some (small) constant power is used. Röglin and Teng [19, Theorem 6.2] proved that for integer programming, the expected value indeed provides a robust measure. The reason for this is that all finite moments of the Pareto curve are polynomially bounded. Thus, a polynomial slow down does not cause the expected running time to jump from polynomial to exponential.

As far as we are aware, this phenomenon, i.e., the case that all finite moments have to be bounded by a polynomial, has not been studied yet in average-case complexity. Thus, for completeness, we define the corresponding average-case and smoothed complexity classes as an alternative to Avg-P and Smoothed-P.

- **Definition 2.6.** 1. An algorithm has robust smoothed polynomial running time with respect to \mathcal{D} if, for all fixed $\varepsilon > 0$ and for every n, x, and ϕ , we have $\mathbb{E}_{y \sim D_{n,x,\phi}}(t_A(y; n, \phi)^{\varepsilon}) = O(n \cdot N_{n,x} \cdot \phi)$. Smoothed-PBM is the class of all (L, \mathcal{D}) for which there exists a deterministic algorithm with robust smoothed polynomial running time. (The "PBM" stands for "polynomially bounded moments".)
- 2. An algorithm A has robust average polynomial running time with respect to \mathcal{D} if, for all fixed $\varepsilon > 0$ and for all n, we have $\mathbb{E}_{y \sim D_n}(t_A(y)^{\varepsilon}) = O(n)$. Avg-PBM contains all (L, \mathcal{D}) for which there exists a deterministic algorithm with robust smoothed polynomial running time.

From the definition, we immediately get Smoothed-PBM \subseteq Smoothed-P and Avg-PBM \subseteq Avg-P. From Röglin and Teng's result [19], one might suspect Avg-P = Avg-PBM and Smoothed-P = Smoothed-PBM, but this does not hold.

Theorem 2.7. Avg-PBM \subsetneq Avg-P and Smoothed-PBM \subsetneq Smoothed-P.

We conjecture that, assuming the exponential time hypothesis (ETH) [14], even an $L \in \mathsf{NP}$ exists to separate Avg-P from Avg-PBM and Smoothed-P from Smoothed-PBM. Given the ETH, 3SAT requires time $2^{\Omega(n)}$ in the worst case, thus also on average if we use the universal distribution. This holds even if we restrict 3SAT to O(n) clauses. However, n is here the number of variables, not the bit length of the input, which is roughly $\Theta(n \log n)$. Thus, a direct application of the ETH seems to be impossible here.

3 Disjoint Supports and Reducibility

The same given input y can appear with very high and with very low probability at the same time. What sounds like a contradiction has an easy explanation: $D_{n,x,\phi}(y)$ can be large whereas $D_{n,x',\phi}(y)$ for some other x' is small. But if we only see y, we do not know whether x or x' was perturbed. This causes some problems when one wants to develop a notion of reduction and completeness.

For a parameterized distributional problem (L, \mathcal{D}) , let

$$L_{\rm ds} = \{ \langle x, y \rangle \mid y \in L \text{ and } |y| \le \operatorname{poly}(|x|) \}.$$

The length of |y| is bounded by the same polynomial that bounds the length of the strings in any supp $D_{n,x,\phi}$. We will interpret a pair $\langle x,y\rangle$ as "y was drawn according to $D_{n,x,\phi}$ ". With the notion of L_{ds} , we can now define a reducibility between parameterized distributional problems. We stress that, although the definition below involves L_{ds} and L'_{ds} , the reduction is defined for pairs L and L' and neither of the two is required to be a disjoint-support language. This means that, for (L, \mathcal{D}) , the supports of $D_{n,x,\phi}$ for different x may intersect. And the same is allowed for (L', \mathcal{D}') .

Definition 3.1. Let (L, \mathcal{D}) and (L', \mathcal{D}) be two parameterized distributional problems. (L, \mathcal{D}) reduces to (L', \mathcal{D}') (denoted by " $(L, \mathcal{D}) \leq_{\text{smoothed}} (L', \mathcal{D}')$ ") if there is a polynomial time computable function f such that for every n, every x, every ϕ and every $y \in \operatorname{supp} D_{n,x,\phi}$ the following holds:

- 1. $\langle x, y \rangle \in L_{ds}$ if and only if $f(\langle x, y \rangle; n, \phi) \in L'_{ds}$. 2. There exist polynomials p and m such that, for every n, x, and ϕ and every $y' \in \operatorname{supp} D'_{m(n), f_1(\langle x, y \rangle; n, \phi), \phi}$, we have

$$\sum_{y:f_2(\langle x,y\rangle;n,\phi)=y'} D_{n,x,\phi}(y) \le p(n) D_{m(n),f_1(\langle x,y\rangle;n,\phi),\phi}(y'),$$

where $f(\langle x,y\rangle;n,\phi) = \langle f_1(\langle x,y\rangle;n,\phi), f_2(\langle x,y\rangle;n,\phi) \rangle.$

(Note that we could also allow that ϕ on the right-hand side is polynomially transformed. However, we currently do not see how to benefit from this.)

It is easy to see that \leq_{smoothed} is transitive. Ideally, Smoothed-P should be closed under this type of reductions. However, we can only show this for the related class of problems with disjoint support.

Definition 3.2. Smoothed- P_{ds} is the set of all distributional problems with disjoint supports such that there is an algorithm A for L_{ds} with smoothed polynomial running time.

Now, Smoothed- P_{ds} is indeed closed under the above type of reductions.

Theorem 3.3. If $(L, \mathcal{D}) \leq_{\text{smoothed}} (L', \mathcal{D}')$ and $(L'_{ds}, \mathcal{D}') \in \text{Smoothed-P}_{ds}$, then $(L_{ds}, \mathcal{D}) \in \mathsf{Smoothed}-\mathsf{P}_{ds}.$

With the definition of disjoint support problems, a begging question is how the complexity of L and L_{ds} are related. It is obvious that $(L, \mathcal{D}) \in \mathsf{Smoothed}-\mathsf{P}$ implies $(L_{ds}, \mathcal{D}) \in \mathsf{Smoothed}-\mathsf{P}_{ds}$. However, the converse is not so obvious. The difference between L and L_{ds} is that for L_{ds} , we get the x from which the input y was drawn. While this extra information does not seem to be helpful at a first glance, we can potentially use it to extract randomness from it. So this question is closely related to the problem of derandomization.

But there is an important subclass of problems in $\mathsf{Smoothed}-\mathsf{P}_{\mathrm{ds}}$ whose counterparts are in Smoothed-P, namely those which have an oblivious algorithm with smoothed polynomial running time. We call an algorithm (or heuristic scheme) for some problem with disjoint supports *oblivious* if the running time on $\langle x, y \rangle$ does not depend on x (up to constant factors). Let Smoothed-P^{obl}_{ds} be the resulting subset of problems in Smoothed-P_{ds} that have such an oblivious algorithm with smoothed polynomial running time.

Theorem 3.4. For any parameterized problem (L, D), $(L, D) \in \text{Smoothed-P}$ if and only if $(L_{ds}, D) \in \text{Smoothed-P}_{ds}^{obl}$.

Note that almost all algorithms, for which a smoothed analysis has been carried out, do not know the x from which y was drawn; in particular, there is an oblivious algorithm for them. Thus, we ask the question: Is there a problem $(L, \mathcal{D}) \notin \mathsf{Smoothed}\text{-P}$ but $(L_{ds}, \mathcal{D}) \in \mathsf{Smoothed}\text{-P}_{ds}$?

Note that in L_{ds} , each y is paired with every x, so there is no possibility to encode information by omitting some pairs. This prohibits attempts for constructing such a problem like considering pairs $\langle x, f(x) \rangle$ where f is some one-way function. For the classes Smoothed-BPP or Smoothed-P/poly, which can be defined in the obvious way, knowing x does not seem to help, since, given y, it should be possible to use the internal random bits (or the advice) to find an x'that is good enough.

4 Parameterized Distributional NP

In this section, we define the smoothed analogue of the worst-case class NP and the average-case class DistNP [13, 16]. First, we have to restrict ourself to "natural" distributions. This rules out, for instance, probability distributions based on Kolmogorov complexity that (the *universal distribution*), under which worst-case complexity equals average-case complexity for all problems [17]. We transfer the notion of computable ensembles to smoothed complexity.

Definition 4.1. A parameterized family of distributions is in PComp_{para} if the cumulative probability $F_{D_{n,x,\phi}} = \sum_{z \leq x} D_{n,x,\phi}$ is can be computed in polynomial time (given n, x and ϕ in binary).

With this notion, we can define the smoothed analogue of NP and DistNP.

Definition 4.2. Dist-NP_{para} = { $(L, D) \mid L \in NP \text{ and } D \in PComp_{para}$ }.

Having defined Dist-NP_{para}, we now prove that *bounded halting* – given a Turing machine, an input, and a running-time bound, does the Turing machine halt on this input within the given time bound – is complete for Dist-NP_{para}. Bounded halting is the canonical NP-complete language, and it has been the first problem that has been shown to be Avg-P-complete [16]. Formally, let

 $BH = \{ \langle g, x, 1^t \rangle \mid NTM \text{ with Gödel number } g \text{ accepts } x \text{ within } t \text{ steps} \}.$

Theorem 4.3. (BH, U^{BH}) is Dist-NP_{para}-complete under smoothed reductions for some $U^{BH} \in \mathsf{PComp}_{para}$.

The original DistNP-complete problem by Levin [16] was Tiling (see also Wang [23]): An instance of the problem consists of a finite set T of square tiles, a positive integer t, and a sequence $s = (s_1, \ldots, s_n)$ for some $n \leq t$ such that s_i matches s_{i+1} (the right side of s_i equals the left side of s_{i+1}). The question is whether S can be extended to tile an $n \times n$ square using tiles from T.

Theorem 4.4. (Tiling, U^{Tiling}) is Dist-NP_{para}-complete under \leq_{smoothed} for some $U^{\text{Tiling}} \in \mathsf{PComp}_{\text{para}}$.

5 Basic Relations to Worst-Case Complexity

Theorem 5.1. If $L \in \mathsf{P}$, then $(L, \mathcal{D}) \in \mathsf{Smoothed}\operatorname{-}\mathsf{P}$ for any \mathcal{D} . If $(L, \mathcal{D}) \in \mathsf{Smoothed}\operatorname{-}\mathsf{P}$ with $\mathcal{D} = (D_{n,x,\phi})_{n,x,\phi}$, then $(L, (D_{n,x_n,\phi})_n) \in \mathsf{Avg}\operatorname{-}\mathsf{P}$ for $\phi = O(\operatorname{poly}(n)/N_{n,x})$ and every sequence $(x_n)_n$ of strings with $|x_n| \leq \operatorname{poly}(n)$.

The observation that Avg-P, P, and Smoothed-P coincide for unary languages allows us to transfer the result that $DistNP \subseteq Avg-P$ implies NE = E [4] to smoothed complexity.

Theorem 5.2. If Dist-NP_{para} \subseteq Smoothed-P, then NE = E.

6 Tractability 1: Integer Programming

Now we will deal with tractable – in the smoothed sense – optimization problems: We will show that if a binary integer linear program can be solved in pseudo-polynomial time, then the corresponding decision problem belongs to Smoothed-P. This result is similar to Beier and Vöckings characterization [3].

A binary optimization problem is an optimization problem of the form "maximize $c^T x$ subject to $w_i^T x \leq t_i$ for $i \in [k]$ and $x \in S \subseteq \{0,1\}^n$ ". Here, $c^T x = \sum_{j=1}^n c_j x_j$ is the linear objective function and $w_i^T x = \sum_{j=1}^n w_{i,j} x_j \leq t_i$ are linear constraints. Furthermore, we have the constraint that the binary vector x must be contained in the set S. This set S should be viewed as containing the "structure" of the problem. Examples are that S contains all binary vectors representing spanning trees in a graph of n vertices or that S represents all paths connecting two given vertices or that S contains all vectors corresponding to perfect matchings of a given graph. Maybe the simplest case is k = 1 and $S = \{0,1\}^n$; then the binary program above represents the knapsack problem.

We assume that S is adversarial (i.e., non-random). Since we deal with decision problems in this paper rather than with optimization problems, we use the standard approach and introduce a threshold for the objective function. This means that the optimization problem becomes the question whether there is an $x \in S$ that fulfills $c^T x \ge b$ as well as $w_i^T x \le t_i$ for all $i \in \{1, \ldots, k\}$. In the following, we treat the budget constraint $c^T x \ge b$ as an additional linear constraint for simplicity. We call this type of problems *binary decision problems*.

For ease of presentation, we assume that we have just one linear constraint (whose coefficients will be perturbed) and everything else is encoded in the set S. This means that the binary decision problem that we want to solve is the following: Does there exist an $x \in S$ with $w^T x \leq t$?

The values w_1, \ldots, w_n are *n*-bit binary numbers. Thus, $w_i \in \{0, 1, \ldots, 2^n - 1\}$. While we can of course vary their length, we choose to do it this way as it conveys all ideas while avoiding another parameter.

We do not make any assumption about the probability distribution of any single coefficient. Instead, our result holds for any family of probability distribution that fulfills the following properties: w_1, \ldots, w_n are drawn according to independent distributions. The set S and the threshold t are part of the input and not subject to randomness. Thus, $N_{n,(S,w,t)} = 2^{n^2}$ for any instance (S, w, t) of size n. We assume that S can be encoded by a polynomially long string. (This is fulfilled for most natural optimization problems, like TSP, matching, shortest path, or knapsack.) The fact that w_1, \ldots, w_n are drawn independently means that the probability for one coefficient to assume a specific value is bounded from above by $\phi^{1/n}$. Since $N_{n,(S,w)} = 2^{n^2}$, the perturbation parameter ϕ can vary between 2^{-n^2} (for the average case) and 1 (for the worst case).

The proof of the following theorem is similar to the one by Beier and Vöcking [3]. However, at various places it gets slightly more technical because we have discrete rather than continuous probability distributions.

Theorem 6.1. If a binary decision problem can be solved in pseudo-polynomial time, then it is in Smoothed-P.

The main open problem concerning Smoothed-P and integer optimization is the following: Beier and Vöcking [3] have proved that (randomized) pseudopolynomiality and smoothed polynomiality are equivalent. The reason why we do not get a similar result is as follows: Our "joint density" for all coefficients is bounded by ϕ , and the density of a single coefficient is bounded by $\phi^{1/n}$. In contrast, in the continuous version, the joint density is bounded by ϕ^n while a single coefficient has a density bounded by ϕ .

However, our goal is to devise a general theory for arbitrary decision problems. This theory should include integer optimization, but it should not be restricted to integer optimization. The problem is that generalizing the concept of one distribution bounded by ϕ for each coefficient to arbitrary problems involves knowledge about the instances and the structure of the specific problems. This knowledge, however, is not available if we want to speak about classes of decision problems as in classical complexity theory.

7 Tractability 2: Graphs and Formulas

Graph Coloring and Smoothed Extension of $G_{n,p}$. The perturbation model that we choose is the smoothed extension of $G_{n,p}$ [22]: Given an adversarial graph G = (V, E) and an $\varepsilon \in (0, 1/2]$, we obtain a new graph G' = (V, E') on the same set of vertices by "flipping" each (non-)edge of G independently with a probability of ε . This means the following: If $e = \{u, v\} \in E$, then e is contained in E' with a probability of $1 - \varepsilon$. If $e = \{u, v\} \notin E$, then $\Pr(e \in E') = \varepsilon$. Transferred to our framework, this means the following: We represent a graph G on n vertices as a binary string of length $\binom{n}{2}$, and we have $N_{n,G} = 2^{\binom{n}{2}}$. The flip probability ε depends on ϕ : We choose $\varepsilon \leq 1/2$ such that $(1 - \varepsilon)^{\binom{n}{2}} = \phi$.

We will focus on graph coloring as a very simple example. k-Coloring is the decision problem whether the vertices of a graph can be colored with kcolors such that no pair of adjacent vertices get the same color. k-Coloring is NP-complete for any $k \ge 3$ [12, GT 4].

Theorem 7.1. For any $k \in \mathbb{N}$, k-Coloring \in Smoothed-P.

Remark 7.2. Bohman et al. [7] and Krivelevich et al. [15] consider a slightly different model for perturbing graphs: Given an adversarial graph, we add random edges to the graph to obtain our actual instance. No edges are removed.

They analyze the probability that the random graph thus obtained is guaranteed to contain a given subgraph H. By choosing H to be a clique of size k+1 and using a proof similar to Theorem 7.1's, we obtain that k-Coloring \in Smoothed-P also with respect to this perturbation model.

Unsatisfiability and Smoothed-RP. Feige [10] and Coja-Oghlan et al. [9] have considered the following model: We are given a (relatively dense) adversarial Boolean k-CNF formula. Then we obtain our instance by negating each literal with a small probability. It is proved that such smoothed formulas are likely to be unsatisfiable, and that their unsatisfiability can be proved efficiently. However, their algorithms are randomized, thus we do not get a result that kUNSAT (this means that unsatisfiability problem for k-CNF formulas) for dense instances belongs to Smoothed-P. However, it shows that kUNSAT for dense instance belongs to Smoothed-RP, where Smoothed-RP is the smoothed analogue of RP: A pair (L, \mathcal{D}) is in Smoothed-RP if there is a randomized polynomial algorithm A with the following properties:

- 1. For all $x \notin L$, A outputs "no" (independent of the perturbation).
- 2. For all $x \in L$, A outputs "yes" with a probability of at least 1/2. (This property is also independent of the perturbation.)
- 3. A has smoothed polynomial running time with respect to \mathcal{D} . (This property is independent of the internal randomness of A.)

Note that we have two sources of randomness in Smoothed-RP: The instance is perturbed, and the algorithm A is allowed to use randomness. Item 1 and 2 depend only on A's own randomness. Item 3 depends only on the perturbation \mathcal{D} . More formally, let kUNSAT_{β} be kUNSAT restricted to instances with at least βn clauses, where n denotes the number of variables. Let ε be the probability that a particular literal is negated. Feige [10] has presented a polynomial-time algorithm with the following property: If $\beta = \Omega(\sqrt{n \log \log n}/\varepsilon^2)$ and the perturbed instance of kUNSAT_{β} is unsatisfiable, which it is with high probability, then his algorithm proves that the formula is unsatisfiable with a probability of at least $1 - 2^{\Omega(-n)}$. The following result is a straightforward consequence.

Theorem 7.3. kUNSAT $_{\beta} \in$ Smoothed-RP for $\beta = \Omega(\sqrt{n \log \log n})$.

8 Smoothed Analysis vs. Semi-Random Models

Semi-random models for graphs and formulas exist even longer than smoothed analysis and can be considered as precursors to smoothed analysis. The basic concept is as follows: Some instance is created randomly that possesses a particular property. This property can, for instance, be that the graph is k-colorable. After that, the adversary is allowed to modify the instance without destroying the property. For instance, the adversary can be allowed to add arbitrary edges between the different color classes. Problems that have been considered in this model or a variant thereof are independent set [11] or graph coloring [5, 8, 11]. However, we remark that these results do not easily fit into a theory of smoothed analysis. The reason is that in these semi-random models, we first have the random instance, which is then altered by the adversary. This is in contrast to smoothed analysis in general and our smoothed complexity theory in particular, where we the adversarial decisions come before the randomness is applied.

9 Discussion

Our framework has many of the characteristics that one would expect. We have reductions and complete problems and they work in the way one expects them to work. To define reductions, we have to use the concept of disjoint supports. It seems to be essential that we know the original instance x that the actual instance y was drawn from to obtain proper domination. Although this is somewhat unconventional, we believe that this is the right way to define reductions in the smoothed setting. The reason is that otherwise, we do not know the probabilities of the instances, which we need in order to apply the compression function. The compression function, in turn, seems to be crucial to prove hardness results. Still, an open question is whether a notion of reducibility can be defined that circumvents these problems. Moreover, many of the positive results from smoothed analysis can be cast in our framework, like it is done in Sections 6 and 7.

Many positive results in the literature state their bounds in the number of "entities" (like number of nodes, number of coefficients) of the instance. However, in complexity theory, we measure bounds in the length (number of symbols) of the input in order to get a theory for arbitrary problems, not only for problems of a specific type. To state bounds in terms of bit length makes things less tight, for instance the reverse direction of integer programming does not work. But still, we think it is more important and useful to use the usual notion of input length such that smoothed complexity fits with average-case and worst-case complexity.

Finally, the results by Röglin and Teng [19] show that, for binary optimization problems, expected polynomial is indeed a robust measure. We have shown that this is in general not the case. To do this, we have used a language in E. The obvious question is now whether Avg-P and Avg-PBM as well as Smoothed-P and Smoothed-PBM coincide for problems in NP.

We hope that the present work will stimulate further research in smoothed complexity theory in order to get a deeper understanding of the theory behind smoothed analysis.

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A Proofs of Section 2

Theorem 2.3. An algorithm A has smoothed polynomial running time if and only if there is an $\varepsilon > 0$ and a polynomial p such that for all n, x, ϕ , and t,

$$\Pr_{y \sim D_{n,x,\phi}}[t_A(y; n, \phi) \ge t] \le \frac{p(n)}{t^{\varepsilon}} \cdot N_{n,x} \cdot \phi.$$

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Proof. Let A be an algorithm whose running time t_A fulfills

$$\mathbb{E}_{y \sim D_{n,x,\phi}} \left(t_A(y; n, \phi)^{\varepsilon} \right) = O\left(n \cdot N_{n,x} \phi \right).$$

The probability that the running time exceeds a certain value t can be bounded by Markov's inequality:

$$\mathbb{P}(t_A(y;n,\phi) \ge t) = \mathbb{P}(t_A(y;n,\phi)^{\varepsilon} \ge t^{\varepsilon})$$

$$\leq \frac{\mathbb{E}_{y \sim D_{n,x,\phi}}(t_A(y;n,\phi)^{\varepsilon})}{t^{\varepsilon}} = O\left(n \cdot N_{n,x}\phi \cdot t^{-\varepsilon}\right).$$

For the other direction, assume that

$$\Pr_{y \sim D_{n,x,\phi}}[t_A(y; n, \phi) \ge t] \le \frac{n^c}{t^{\varepsilon}} \cdot N_{n,x}\phi$$

for some constants c and ε . Let $\varepsilon' = \varepsilon/(c+2)$. Then we have

$$\mathbb{E}_{y \sim D_{n,x,\phi}} \left(t_A(y;n,\phi)^{\varepsilon'} \right) = \sum_t \mathbb{P}(t_A(y;n,\phi)^{\varepsilon'} \ge t)$$

$$\leq n + \sum_{t \ge n} \mathbb{P}(t_A(y;n,\phi) \ge t^{1/\varepsilon'})$$

$$\leq n + \sum_{t \ge n} t^{-2} \cdot N_{n,x}\phi = n + O(N_{n,x}\phi).$$

Theorem 2.5. $(L, \mathcal{D}) \in \mathsf{Smoothed}\text{-P}$ if and only if (L, \mathcal{D}) has an errorless heuristic scheme.

Proof. Let A be an algorithm for (L, \mathcal{D}) . By Theorem 2.3, the probability that

$$\mathbb{P}(t_A(y; n, \phi) \ge t) = O\left(n \cdot N_{n,x} \phi \cdot t^{-\varepsilon}\right).$$

We get an errorless heuristic scheme B from A as follows: Simulate A for $(n \cdot N_{n,x}\phi/\delta)^{1/\varepsilon}$ steps. If A stops within these number of steps, then output whatever A outputs. Otherwise, output \bot . By the choice of the parameters, the probability that B outputs \bot is bounded by δ .

For the other direction, let A be an errorless heuristic scheme for (L, \mathcal{D}) . We get an algorithm with smoothed polynomial running time by first running A with

 $\delta = 1/2$, then with $\delta = 1/4$, and in the *i*th iteration with $\delta = 1/2^i$. Whenever A does not answer \bot , B gives the same answer and stops. B will eventually stop, when $\delta < D_{n,x,\phi}(y)$. For *i* iterations, B needs

$$\sum_{j=1}^{i} q(n, N_{n,x}\phi, 2^j) \le \operatorname{poly}(n, N_{n,x}\phi) \cdot 2^{ci}$$

for some constant c. B stops after i iterations for all but a 2^{-i} fraction of the input. Thus B has smoothed polynomial running time. (Choose $\varepsilon < 1/c$.)

Theorem 2.7. Avg-PBM \subsetneq Avg-P and Smoothed-PBM \subsetneq Smoothed-P.

Proof. We only prove the theorem for average-case complexity. The proof for the smoothed complexity case is almost identical.

By the time hierarchy theorem [1], there is a language $L' \in \mathsf{DTime}(2^n)$ such that $L' \notin \mathsf{DTime}(2^{o(n)})$. Consider the following language $L = \{x0^n \mid |x| = n, x \in L'\}$. Let $\mathcal{D}' = (D'_n)$ be a hard probability distribution for L', i.e., (L', \mathcal{D}') is as hard to solve as L' in the worst case [17, 18]

Let $\mathcal{D} = (D_n)$ be given as follows:

$$D_n(xy) = \begin{cases} 2^{-n} \cdot D'_n(x) & \text{if } |x| = n \text{ and } y = 0^n \text{ and} \\ 2^{-2n} & \text{otherwise.} \end{cases}$$

Since $L' \in \mathsf{DTime}(2^n)$, we have $(L, \mathcal{D}) \in \mathsf{Avg-P}$: L can be decided in expected time time $2^{-n} \cdot 2^n + O(n) = O(n)$. Now we prove that $(L, \mathcal{D}) \notin \mathsf{Avg-PBM}$. If $(L, \mathcal{D}) \in \mathsf{Avg-PBM}$ were true, then $2^{-n} \cdot \mathbb{E}_{x \sim D'_n}(t(x)^c)$ would be bounded by a polynomial for all fixed c. Here, t is the time needed to solve the L' instance x.

Our choice of \mathcal{D}' , Jensen's inequality, and the fact that $L' \notin \mathsf{DTime}(2^{o(n)})$ implies that $\mathbb{E}_{x \sim D'_n}(t(x)^c) \geq \mathbb{E}_{x \sim D'_n}(t(x))^c = 2^{c \cdot \Omega(n)}$. Thus, for some sufficiently large $c, 2^{-n} \cdot \mathbb{E}_{x \sim D'_n}(t(x)^c)$ exceeds any polynomial.

Theorem 3.3. If $(L, D) \leq_{\text{smoothed}} (L', D')$ and $(L'_{\text{ds}}, D') \in \text{Smoothed-P}_{\text{ds}}$, then $(L_{\text{ds}}, D) \in \text{Smoothed-P}_{\text{ds}}$.

Proof. Let A' be a an errorless heuristic scheme for (L'_{ds}, \mathcal{D}') . Let f be the reduction from (L, \mathcal{D}) to (L', \mathcal{D}') and let p and m be the corresponding polynomials.

We claim that $A(\langle x, y \rangle; n, \phi, \delta) = A'(f(\langle x, y \rangle; n, \phi); m(n), \phi, \delta/p(n))$ is an errorless heuristic scheme for (L_{ds}, \mathcal{D}) . To prove this, let

$$B = \{ y' \in \operatorname{supp} D'_{m(n), f_1(\langle x, y \rangle; n, \phi), \phi} \mid A'(\langle f(\langle x, y \rangle; n, \phi), y' \rangle; m(n), \phi, \delta/p(n)) = \bot \}$$

be the set of string on which A' fails.

Because A' is a heuristic scheme, we have $D'_{m(n),f_1(\langle x,y\rangle;n,\phi),\phi}(B) \leq \delta/p(n)$. Therefore,

$$\begin{aligned} &\Pr_{y\sim D_{n,x,\phi}}\left(A(\langle x,y\rangle;n,\phi,\delta)=\bot\right)\\ &= \Pr_{y\sim D_{n,x,\phi}}\left(A'(f(\langle x,y\rangle;n,\phi);m(n),\phi,\delta/p(n)=\bot)\right)\\ &= \sum_{y:f_2(\langle x,y\rangle;n,\phi)\in B}D_{n,x,\phi}(y)\\ &\leq \sum_{y'\in B}p(n)D'_{m(n);f_1(\langle x,y\rangle;n,\phi);\phi}(y')\\ &= p(n)D'_{m(n);f_1(\langle x,y\rangle;n,\phi);\phi}(B)\leq\delta. \end{aligned}$$

Thus, $(L_{ds}, \mathcal{D}) \in \mathsf{Smoothed}-\mathsf{P}_{ds}$.

Β **Proofs of Section 3**

Theorem 3.4. For any parameterized problem $(L, \mathcal{D}), (L, \mathcal{D}) \in \mathsf{Smoothed-P}$ if and only if $(L_{ds}, \mathcal{D}) \in \mathsf{Smoothed}-\mathsf{P}_{ds}^{\mathrm{obl}}$.

Proof. Let A be an oblivious algorithm with smoothed polynomial running time for L_{ds} . Since A is oblivious, we get an algorithm for L with the same running time (up to constant factors) by running A on $\langle x_0, y \rangle$ on input y, where x_0 is an arbitrary string of length n.

\mathbf{C} **Proofs of Section 4**

For the completeness result of BH, we first need a "compression function" according to the following lemma.

Lemma C.1. Let $\mathcal{D} = (D_{n,x,\phi}) \in \mathsf{PComp}_{\text{para}}$ be an ensemble. There exists a deterministic algorithm C such that the following holds:

- 1. $C(y; n, x, \phi)$ runs in time polynomial in n and ϕ for all $y \in \operatorname{supp} D_{n,x,\phi}$.
- 2. For every $y, y' \in \text{supp } D_{n,x,\phi}$, $C(y; n, x, \phi) = C(y'; n, x, \phi)$ implies y = y'.
- 3. If $D_{n,x,\phi}(y) < 2^{-|y|}$, then $|C(y; n, x, \phi)| = 1 + |y|$. Else, $|C(y; n, x, \phi)| = \log \frac{1}{D_{n,x,\phi}(y)} + c \cdot \log n + 1$ for some constant c.

Proof. Consider any string $y \in \text{supp}(D_{n,x,\phi})$. If $D_{n,x,\phi}(y) \leq 2^{-|y|}$, then we let $C(y; n, x, \phi) = 0y$. If $D_{n,x,\phi}(y) > 2^{-|x|}$, then let y' be the string that precedes y in lexicographic order, and let $p = F_{D_{n,x,\phi}}(y')$. Then we set $C(y; n, x, \phi) = 1a$, where a is the longest common prefix of the binary representation of p and $F_{D_{n,x,\phi}}(y) = p + D_{n,x,\phi}(y)$. Since $\mathcal{D} \in \mathsf{PComp}_{\text{para}}$, the string z can be computed in polynomial time.

We have $D_{n,x,\phi}(y) \leq 2^{-|a|}$, since adding $D_{n,x,\phi}(y)$ leaves the first |a| bits of p unchanged.

Let z be another string, z' its predecessor and b the longest common prefix of $q = F_{D_{n,x,\phi}}(z')$ and $q + D_{n,x,\phi}(z')$. The intervals $[p, p + D_{n,x,\phi}(y))$ and $[q, q + D_{n,x,\phi}(y))$ are disjoint by construction. Therefore, a and b have to be different, because otherwise these intervals would intersect.

Let c be such that $|y| \leq n^c$ for all $y \in \text{supp}(D_{n,x,\phi})$. We set

$$C(y; n, x, \phi) = 1 \operatorname{bin}(|a|) a 0^{\log \frac{1}{D_{n,x,\phi}(y)} - |a|}$$

(Note that $\log \frac{1}{D_{n,x,\phi}(y)} \ge |a|$.) Here bin(|a|) is a fixed length binary encoding of a. We can bound this length by $c \log n$. The total length of $C(y; n, x, \phi)$ is

$$|C(y; n, x, \phi)| = 1 + c \log n + \log \frac{1}{D_{n, x, \phi}(y)}$$

It remains to be proved that C is injective. Let $C(y; n, x, \phi) = C(z; n, x, \phi)$. If $C(y; n, x, \phi)$ starts with a 0, then obviously y = z. If $C(y; n, x, \phi)$ starts with a 1, then the prefixes a and b are the same. Therefore y = z by the consideration above.

The instances of BH are triples $\langle g, x, 1^t \rangle$ of length $2 \log |g| + 2 \log |x| + |x| + |g| + t + \Theta(1)$. Let

$$U_{N,\langle g,x,1^t\rangle,\phi}^{\text{BH}}(\langle g',x',1^{t'}\rangle) = \begin{cases} c_{\phi}2^{-|x'|} & \text{if } g = g', N = |\langle g',x',1^{t'}\rangle|, |x'| \ge \log \frac{1}{\phi}, \\ 0 & \text{otherwise.} \end{cases}$$

Above, c_{ϕ} is an appropriate scaling factor. More precisely, c_{ϕ} is the reciprocal of the number of possible lengths for a string x', i.e., it is of order $\frac{1}{N-\log \phi}$. In particular, $U_{N,\langle g,x,1^t\rangle,\phi}^{\text{BH}}(y) \leq \phi$ for all y.

Theorem 4.3. (BH, U^{BH}) is Dist-NP_{para}-complete under polynomial-time smoothed reductions.

Proof. Let $(L, \mathcal{D}) \in \text{Dist-NP}_{\text{para}}$ be arbitrary. Let p(n) be an upper bound for the length of the strings in any $\text{supp}(D_{n,x,\phi})$. Let M be a nondeterministic machine that accepts an input a if and only if there is a string $y \in L$ with $C(y; n, x, \phi) = a$. Let q be an upper bound on the running time of M. Let g be the Gödel number of M. Our reduction maps a string y to

$$f(\langle x, y \rangle; n, x, \phi) = \left\langle \langle g, C(x; n, x, \phi), 1^t \rangle, \langle g, C(y; n, x, \phi), 1^{t'} \rangle \right\rangle$$

where t and t' chosen in such a way that they are larger than q(p(|x|)). (And t and t' should be chosen in such a way that all tuples have the same length N(n).)

By construction, $\langle x, y \rangle \in L_{ds}$ if and only if $f(\langle x, y \rangle; n, x, \phi) \in BH_{ds}$.

Domination remains to be verified. Since C is injective, at most one y is mapped to $\langle g, a, 1^t \rangle$ given n, x, and ϕ . We have

$$U_{N,\langle g,C(x;n,x,\phi),1^t\rangle,\phi}^{\mathrm{BH}}(\langle g,C(y;n,x,\phi),1^{t'}\rangle) = c_{\phi} \cdot 2^{-|C(y;n,x,\phi)|}.$$

If $|C(y; n, x, \phi)| \le \log \frac{1}{D_{n, x, \phi}(y)} + c \log n + 1$, then

$$U_{N,\langle g,C(x;n,x,\phi),1^t\rangle,\phi}^{\mathrm{BH}}(\langle g,C(y;n,x,\phi),1^{t'}\rangle) \leq c_{\phi}\cdot\frac{D_{n,x,\phi}(x)}{2n^c}$$

and domination is fulfilled. If $|C(y; n, x, \phi)| = 1 + |y|$, then

$$U_{N,\langle g,C(x;n,x,\phi),1^t\rangle,\phi}^{\mathsf{BH}}(\langle g,C(y;n,x,\phi),1^{t'}\rangle) \le c_{\phi} \cdot 2^{-|y|-1} \le 2c_{\phi} \cdot D_{n,x,\phi}(y).$$

This completes the hardness proof. The completeness follows since (BH, U^{BH}) is indeed contained in Dist-NP_{para}.

We use the following probability distribution for Tiling:

$$U_{N,\langle T,s,1^t\rangle,\phi}^{\texttt{Tiling}}(\langle T',s',1^{t'}\rangle) = \begin{cases} c_{\phi}a^{-|s'|} & \text{if } T = T', \ N = |\langle T',s',1^{t'}\rangle|, \ |T'| \ge \log\frac{1}{\phi}, \\ 0 & \text{otherwise.} \end{cases}$$

Here, a is the number of possible choices in T for each initial tile s_i .

Theorem 4.4. (Tiling, U^{Tiling}) is Dist-NP_{para}-complete under \leq_{smoothed} .

Proof. By construction, we have $(\texttt{Tiling}, U^{\texttt{Tiling}}) \in \mathsf{Dist-NP}_{para}$. For simplicity, we assume that the set T of tiles always contains two tiles encoding the input bits "0" and "1" and that these are the only possible tiles for the initial tiling (s_1, \ldots, s_n) . (The problem does not become easier without this restriction, but the hardness proof becomes more technical.)

For the hardness, (BH, U^{BH}) reduces to $(Tiling, U^{Tiling})$ because the Turing machine computations can be encoded as tiling problems in a straightforward way [23] (the Gödel number g maps to some set T of tiles, and the input x maps to the initial tiling s). Finally, Item 2 of the reduction (Definition 3.1) is fulfilled can be seen to be satisfied because of the similarity between the two probability distributions.

D Proofs of Section 7

Theorem 7.1. For any $k \in \mathbb{N}$, k-Coloring \in Smoothed-P.

Proof. To show that k-Coloring \in Smoothed-P, we analyze the following simple algorithm: First, we check whether the input graph contains a clique of size k + 1. This can be done easily in polynomial time. If yes, we output no. If no, we perform exhaustive search.

The analysis is similar to Wilf's analysis [24] of the coloring problem: First, we check whether the input graph contains a clique of size k + 1. This can be done easily in polynomial time. If yes, we output no. If no, we perform exhaustive search. The correctness of the algorithm is obvious.

A graph is k-colorable only if it does not contain a clique of size k + 1. The probability that a specific set of k + 1 vertices form a k + 1 clique is at least $\varepsilon^{\binom{k+1}{2}}$. Thus, the probability that a graph G on n vertices does not contain a k+1 clique is at most $\left(1-\varepsilon^{\binom{k+1}{2}}\right)^{\frac{n}{k+1}}$.

We distinguish two cases: First, $\varepsilon \geq 0.1$. In this case, $\left(1 - \varepsilon^{\binom{k+1}{2}}\right)^{\frac{n}{k+1}}$ can be bounded from above by c^n for some positive constant c < 1 that depends on k. Brute-force testing whether a graph can be k-colored can be done in time $\operatorname{poly}(n) \cdot k^n$. The probability that we need brute force is at most c^n . Thus, the expected running-time, raised to the power $\varepsilon = \log_k(1/c)$, is bounded from above by a polynomial.

Second, $\varepsilon < 0.1$. Then we have $\phi = (1 - \varepsilon)^{\binom{n}{2}} \ge 0.9^{\binom{n}{2}}$. The allowed runningtime (raised to some constant power) is $N_{n,G}\phi n = 2^{\binom{n}{2}}\phi n \ge 1.8^{\binom{n}{2}}$. Thus, we can afford exhaustive search in every run.

E Proofs and Details of Section 6

Technical Lemmas and Main Theorem. The following simple lemma bounds the probability that a certain coefficient assumes a value in a given small interval.

Lemma E.1. Let $\delta, z \in \mathbb{N}$. Let a be an n-bit coefficient drawn according to some discrete probability distribution bounded from above by $\phi^{1/n}$. Then $\Pr(a \in [z, z + \delta)) \leq \phi^{1/n} \delta$.

Proof. There are exactly δ outcomes of a that lead to $a \in [z, z + \delta)$. Thus, $\Pr(a \in [z, z + \delta)) \leq \phi^{1/n} \varepsilon$.

Our goal is to show that $O(\log(n\phi^{1/n}2^n))$ bits for each coefficient suffice to determine whether a solution exists. (For the average case, we have $\phi = 2^{-n^2}$, thus $O(\log n)$ bits per coefficient.) To do this, it is not sufficient for an $x \in S$ to just satisfy $w^T x \leq t$: Because of the rounding, we might find that x is feasible with respect to the rounded coefficients whereas x is infeasible with respect to the true coefficients. Thus, what we need is that $w^T x$ is sufficiently smaller than t. Then the rounding does not affect the feasibility. Unfortunately, we cannot rule out the existence of solutions $x \in S$ that are very close to the threshold (after all, there can be an exponential number of solutions, and it is likely that some of them are close to the threshold). But it is possible to prove the following: Assume that there is some ranking among the solutions $x \in S$. Let the winner be the solution $x^* \in S$ that fulfills $w^T x^* \leq t$ and is ranked highest among all such solutions. Then it is likely that $t - w^T x^*$ is not too small. Now, any solution that is ranked higher than x^* must be infeasible because it violates the linear constraint $w^T x \leq t$. Let \hat{x} be the solution that minimizes $w^T x - t$ among all solutions ranked higher than x^* . Then it is also unlikely that $w^T \hat{x} - t$ extremely small, i.e., that \hat{x} violates the linear constraint by only a small margin.

Remark E.2. In Beier and Vöcking's analysis [3], the ranking was given by the objective function. We do not have an objective function here because we deal

with decision problems. Thus, we have to introduce a ranking artificially. In the following, we use the lexicographic ordering (if not mentioned otherwise), which satisfies the following *monotonicity* property that simplifies the proofs: if $x \in S$ is ranked higher than $y \in S$, then there is an *i* with $x_i = 1$ and $y_i = 0$.

Now let x^* be the winner (if it exists), i.e., the highest ranked (with respect to lexicographic ordering) solution among all feasible solutions. Then we define the *winner gap* as

$$\Gamma(t) = \begin{cases} t - w^T x^* & \text{if there exists a feasible solution and} \\ \bot & \text{otherwise.} \end{cases}$$

The goal is to show that it is unlikely that Γ is small. In order to analyze Γ , it is useful to define also the loser gap Λ . The loser $\hat{x} \in S$ is a solution that is ranked higher than x^* but cut off by the constraint $w^T x \leq t$. It is the solution with minimal $w^T x - t$ among all such solutions. (If there is a tie, which can happen because we have discrete probability distributions, then we take the highest-ranked solution as the loser.) We define

$$\Lambda(t) = \begin{cases} w^T \hat{x} - t & \text{if there exists a loser } \hat{x} \text{ and} \\ \bot & \text{otherwise.} \end{cases}$$

The probability that Λ or Γ is smaller than some value δ is bounded by $\delta \phi^{1/n} n$, which we will prove in the following.

The following lemma states that it suffices to analyze Λ in order to get bounds for both Λ and Γ . In fact, for the setting with just one linear constraint with non-negative coefficients, we do not even need the winner gap. But the winner gap is needed for more general cases, which we discuss briefly below but do not treat in detail for conciseness.

Lemma E.3 (discrete version of [3, Lemma 7]). For all t and δ , we have $\Pr(\Gamma(t) < \delta) = \Pr(\Lambda(t - \delta) \le \delta)$.

Proof. A solution $x \in S$ is called *Pareto-optimal* if there is no other solution $x' \in S$ such that $w^T x' \leq w^T x$ and x' is ranked higher than x. Let us make two observations. First, we observe that both winners and losers are Pareto-optimal. Second, for every Pareto-optimal solution x, there exists a threshold t such that x is the loser for this particular threshold. To see this, simply set $t = w^T x - 1$.

Let $P\subseteq S$ be the set of Pareto-optimal solutions. Then

$$\Gamma(t) = \min\{t - w^T x \mid x \in P, w^T x \le t\} \text{ and }$$

$$\Lambda(t) = \min\{w^T x - t \mid x \in P, w^T x > t\} = \min\{w^T x - t \mid x \in P, w^T x \ge t + 1\}.$$

Now $\Gamma(t) < \delta$ if and only if there is an $x \in P$ with $t - w^T x \in \{0, \dots, \delta - 1\}$. This is equivalent to $w^T x - t \in \{-\delta + 1, \dots, 0\}$ and to $w^T x - (t - \delta) \in \{1, \dots, \delta\}$. In turn, this is equivalent to $\Lambda(t - \delta) \leq \delta$.

Now we analyze $\Lambda(t)$. The following lemma makes this rigorous. It is a discrete counterpart to Beier and Vöcking's separating lemma [3, Lemma 5]. We have to assume that the all-zero vector is not contained in S. The reason for this is that its feasibility does not depend on any randomness.

Lemma E.4 (separating lemma). Suppose that $(0, ..., 0) \notin S$. For every $\delta, t \in \mathbb{N}$, we have $\Pr(\Gamma(t) < \delta) \leq \delta \phi^{1/n} n$ and $\Pr(\Lambda(t) \leq \delta) \leq \delta \phi^{1/n} n$.

If we use a non-monotone ranking, then the bounds for the probabilities become $\delta \phi^{1/n} n^2$.

Proof. Because of Lemma E.3, it suffices to analyze the loser gap Λ . We only give a proof sketch for monotone rankings as that emphasis the differences to the continuous counterpart [3, Lemma 5].

the continuous counterpart [3, Lemma 5]. Let $S_i = \{x \in S \mid x_i = 1\}$, and let $\overline{S}_i = S \setminus S_i = \{x \in S \mid x_i = 0\}$. Let $x^{\star i} \in \overline{S}_i$ be the winner from \overline{S}_i : $x^{\star i}$ is ranked highest in \overline{S}_i and satisfies the linear constraint $w^T x^{\star i} \leq t$. Let $\hat{x}^i \in S_i$ be the loser with respect to $x^{\star i}$, i.e., a solution that is ranked higher than $x^{\star i}$ and minimizes $w^T \hat{x}^i - t$ (if such a solution exists). Let

$$\Lambda_i = \begin{cases} w^T \hat{x}^i - t & \text{if } \hat{x}^i \text{ exists and} \\ \bot & \text{otherwise.} \end{cases}$$

Note that \hat{x}^i can be feasible and, thus, Λ_i can be negative.

To analyze Λ_i , we assume that all w_j with $j \neq i$ are fixed by an adversary. The winner $x^{\star i}$ does not depend w_i because all solutions $x \in \overline{S}_i$ have $x_i = 0$. Once $x^{\star i}$ is fixed, also \hat{x}^i is fixed. Because w_j for $j \neq i$ is fixed and $\hat{x}_i^i = 1$, we can rewrite $w^T \hat{x}^i - t = z + w_i$. Now $\Lambda_i \in \{1, \ldots, \delta\}$ if w_i assumes a value in some interval of length δ , which happens with a probability of at most $\delta \phi^{1/n}$.

Furthermore, if $\Lambda \neq \bot$, then there exists an *i* with $\Lambda_i = \Lambda$ [3, Claim B]. Thus, a union bound over all *n* possibilities for *i* yields $\Pr(\Lambda(t) \leq \delta) \leq \delta \phi^{1/n} n$.

For the probabilistic constraint $w^T x \leq t$, it is not sufficient for an x to satisfy it. Instead, we want that only a few bits of each coefficient of w suffice to find an x that satisfies that constraint. Here, "few" means roughly $O(\log(n\phi^{1/n}2^n))$. (Note that this is roughly $O(\log n)$ if we are close to the average case, where $\phi \approx 2^{-n^2}$.) Different from Beier and Vöcking's continuous case (where the realvalued coefficients where revealed by an oracle), we have the true coefficients at hand. Thus, we do not need their certificates that a solution is indeed feasible, but we can simply test with the true coefficients. Clearly, this testing can be done in polynomial time.

For an *n*-bit natural number *a* and $b \in \mathbb{N}$, let $\lfloor a \rfloor_b =$ be the number obtained from *a* by only taking the *b* most significant bits. This means that $\lfloor a \rfloor_b = 2^{n-b} \cdot \lfloor a/2^{n-b} \rfloor$.

In order to show that pseudo-polynomiality implies smoothed polynomial complexity, we use a pseudo-polynomial algorithm as a black box in the following way: We run the pseudo-polynomial algorithm with the highest $O(\log n)$ bits. (To do this, we scale the rounded coefficients of w down. Furthermore, we also have to scale t down appropriately.) If we find a solution, then we check it against the true coefficients of w. If it remains feasible, we output "yes". If it becomes unfeasible, then we take one more bit for each coefficient and continue. The following lemma gives a tail bound for how long this can go on.

Lemma E.5. Assume that we use b bits for each coefficient of w. Let x^* be the winner (with respect to the true w without rounding). The probability that solving the problem with b bits for each coefficient yields a solution different from x^* is bounded from above by $2^{n-b}\phi^{1/n}n^2$.

Proof. We only get a solution different from x^* if there is a solution \hat{x} ranked higher than x^* that is feasible with respect to the rounded coefficients. By rounding, we change each coefficient by at most 2^{n-b} . Thus, $w^T \hat{x} - \lfloor w \rfloor_b^T \hat{x} \leq 2^{n-b}n$.

We can conclude that we find \hat{x} instead of x^* only if the loser gap Λ is at most $2^{n-b}n$, which happens with a probability of at most $2^{n-b}\phi^{1/n}n^2$ (or $2^{n-b}\phi^{1/n}n^3$ if the ranking is not monotone).

With this preparation, we can prove the main result of this section.

Theorem 6.1. If a binary decision problem can be solved in pseudo-polynomial time, then it is in Smoothed-P.

Proof. We have to show that the running time of the algorithm sketched above, which uses the pseudo-polynomial algorithm as a black box, fulfills Theorem 2.3.

If b bits for each coefficient are used, the running time of the pseudo-polynomial algorithm is bounded from above by $O((n2^b)^c)$ for some constant c. (Even the total running time summed over all iterations up to b bits being revealed is bounded by $O((n2^b)^c)$, because it is dominated by the last iteration.)

The probability that more than time $t = O((n2^b)^c)$ is needed is bounded from above by $2^{n-b}\phi^{1/n}n^2$ according to Lemma E.5. We can rewrite this as

$$2^{n-b}\phi^{1/n}n^2 = n^2 2^{-b} \left(2^{n^2}\phi\right)^{1/n} = \frac{n^3}{O(t^{1/c})} \cdot \left(2^{n^2}\phi\right)^{1/n} \le \frac{n^3}{O(t^{1/c})} \cdot 2^{n^2}\phi.$$

The last inequality holds because $\phi \geq 2^{-n^2}$. The theorem is proved because this tail bound for the running time is strong enough according to Theorem 2.3.

Examples of problems in Smoothed-P are the decision problems associated with the following NP-hard optimization problems:

- knapsack, where the goal is to find a subset of a given collection of items that maximizes the profit while obeying a budget for its weight;
- constrained shortest path, where the goal is to find a path of minimum length that obeys a certain a budget;
- constrained minimum-weight spanning tree.

These problems can be solved in pseudo-polynomial time using dynamic programming, even if we insist on a lexicographically maximal solution (as we have to for Lemma E.4).

Let us now discuss some extensions of the model. We have restricted ourselves to deterministic pseudo-polynomial algorithms, which yield smoothed polynomial complexity. These deterministic algorithms can be replaced without any complication by randomized errorless algorithms that have expected pseudopolynomial running time.

So far, we have not explicitly dealt with constraints of the form " $w^T x \ge t$ ". But they can be treated in the same way as " $w^T x \le t$ ", except that winner and loser gap change their roles. Furthermore, we did not include the case that coefficients can be positive or negative. This yields additional technical difficulties (we have to round more carefully and take both winner and loser gap into account), but we decided to restrict ourselves to the simpler form with non-negative coefficients for the sake of clarity. Moreover, we have not considered the case of multiple linear constraints [3, Section 2.3] for the same reason. Finally, Röglin and Vöcking [20] have extended the smoothed analysis framework to integer programming. We believe that the same can be done for our discrete setting.