ABSTRACT

We consider the problem of clustering mixtures of mean-separated Gaussians in high dimensions. We are given samples from a mixture of $k$ identity covariance Gaussians, so that the minimum pairwise distance between any two pairs of means is at least $\Delta$, for some parameter $\Delta > 0$, and the goal is to recover the ground truth clustering of these samples. It is folklore that separation $\Delta = \Theta(\sqrt{\log k})$ is both necessary and sufficient to recover a good clustering (say with constant or $1/\text{poly}(k)$ error), at least information theoretically. However, the estimators which achieve this guarantee are inefficient. We give the first algorithm which runs in polynomial time, and which almost matches this guarantee. More precisely, we give an algorithm which takes polynomially many samples and time, and which can successfully recover a good clustering, so long as the separation is $\Delta = \Omega((\log k)^{1/2+\epsilon})$, for any $\epsilon > 0$. Previously, polynomial time algorithms were only known for this problem when the separation was polynomial in $k$, and all algorithms which could tolerate poly log $k$ separation required quasipolynomial time.

We also extend our result to mixtures of translations of a distribution which satisfies the Poincaré inequality, under additional mild assumptions. Our main technical tool, which we believe is of independent interest, is a novel way to implicitly represent and extract important information about high-degree moments without ever writing down the full moment tensors explicitly.

1 INTRODUCTION

Gaussian mixture models are some of the most popular generative models in both theory and practice. A $k$-Gaussian mixture model $\mathcal{M}$ (henceforth $k$-GMM) is a distribution specified by $k$ non-negative mixing weights $w_1, \ldots, w_k$ which sum to 1, and $k$ component Gaussians $N(\mu_1, \Sigma_1), \ldots, N(\mu_k, \Sigma_k)$, and is given by the probability density function

$$M = \sum_{i=1}^{k} w_i N(\mu_i, \Sigma_i).$$

In other words, to draw a sample from $\mathcal{M}$, we select the $i$-th component with probability $w_i$, and then draw an independent sample from $N(\mu_i, \Sigma_i)$. An important special case of this model is the isotropic (also sometimes referred to as spherical) case, where $\Sigma_i = I$ for all $i = 1, \ldots, k$.

One of the most important and well-studied problems for GMMs and isotropic GMMs in particular is clustering. Given $n$ independent samples $X_1, \ldots, X_n$ drawn from an unknown $k$-GMM, the goal of clustering is to recover a partition of the data points into $k$ parts $S_1, \ldots, S_k$ so that (a) almost all the points in every cluster are drawn from the same component Gaussian, and (b) for every component, there is some cluster which contains almost all of the samples drawn from it. Of course, not every isotropic GMM is clusterable: for instance, if two components are identical, then it is information-theoretically impossible to detect which component a sample came from, and so recovering a good clustering is also impossible. In the isotropic setting, a necessary and sufficient condition for the GMM to be clusterable is some amount of mean separation, namely, that $\|\mu_i - \mu_j\|_2 \geq \Delta$ for all $i \neq j$, for some parameter $\Delta$.

The question then becomes: how small can we take $\Delta$ so that we can still cluster? For simplicity, for the remainder of this section, let us assume that $w_i \geq 1/\text{poly}(k)$ for all $i = 1, \ldots, k$, that is, all the components have a nontrivial amount of weight. Information-theoretically, it is well-known (see e.g. [40]) that $\Delta = \Theta(\sqrt{\log k})$ separation is necessary and sufficient to achieve a clustering which correctly clusters more than an $1 - \epsilon$ fraction of the points with high probability, for all $\epsilon > 0$. Unfortunately, all known efficient algorithms for clustering at this separation rely on brute force methods, and run in exponential time.

If we restrict our attention to efficient estimators, the state of affairs is a bit more complicated. In fact, for over 20 years, the smallest separation that polynomial time algorithms could tolerate was $\Delta = \Omega(k^{1/4})$ [44]. It was not until recently that three concurrent papers [21, 26, 31] gave algorithms which could handle separation $\Delta = \Omega(k^{1/3})$ with runtime and sample complexity $O(d\text{poly}(1/\epsilon'))$. In particular, whenever the separation is some fixed polynomial in $k$, these algorithms run in polynomial time. Unfortunately, if we
wish to achieve the optimal separation of $\Omega(\sqrt{\log k})$—or indeed, any polylogarithmic amount of separation—these algorithms would require quasipolynomial time and sample complexity.

The barrier at quasipolynomial time. These algorithms all get stuck at quasipolynomial time when $\Delta = \Theta(\text{poly log } k)$ for the same reason. Fundamentally, all current algorithmic approaches to this problem rely on the following geometric identifiability fact:

Given enough samples from an isotropic k-GMM with separation $\Delta = \Omega(k^{\gamma})$, then sufficiently large subset of samples whose empirical $t = \Theta(1/\gamma)$ moments approximately match those of a Gaussian along all projections must essentially recover one of the true clusters.

Algorithmically, this amounts to finding a subset of points whose empirical $t$-th moment tensor is close to that of a Gaussian in the appropriate norm. Since this results in a hard optimization problem whenever $t > 2$, these algorithms often solve some suitable relaxation of this using something like the Sum of Squares hierarchy. But, as the separation decreases, the relaxations must match more and more moments. In particular, to achieve $\Delta = \Theta(\text{poly log } k)$, one must set $t = \text{poly log } k$, that is, one must match polynologically many moments. However, even writing down the degree $t = \text{poly log } k$ moment tensor requires quasipolynomial time, and guaranteeing that the empirical moment tensor concentrates requires quasipolynomially many samples. As a result, the aforementioned algorithms all require quasipolynomial time and sample complexity, as they need to not only write down the moment tensor, but perform some fairly complex optimization tasks on top of it.

On the flip side, there is no concrete reason for pessimism either. While there are lower bounds against large classes of efficient algorithms for clustering mixtures of arbitrary Gaussians, see e.g. [10, 20], none of these apply when the components are isotropic. In particular, this leaves open the appealing possibility that one could even cluster down to separation $\Delta = \Omega(\sqrt{\log k})$ in polynomial time. Stated another way, the question becomes:

Can we cluster any clusterable mixture of isotropic Gaussians in polynomial time?

1.1 Our Results

In this paper, we (almost) resolve this question in the affirmative. Namely, for all constants $c > 0$, we give an algorithm which takes polynomially many samples and time, and which can cluster with high probability, so long as the separation satisfies $\Delta = \Omega(\log^{1/2+c} k)$. In other words, our algorithm can almost match the information theoretically optimal separation, up to sub-polynomial factors. Our main theorem is:

Theorem 1.1 (informal, see Theorem 2.5, Corollary 2.6). Let $c > 0$ be fixed but arbitrary. Let $\mathcal{M}$ be a mixture of $k$ isotropic Gaussians with minimum mixing weight lower bounded by $1/\text{poly}(k)$, and minimum mean separation at least $\Delta = \Omega(\log^{1/2+c} k)$. Then, there is an algorithm which, given samples $X_1, \ldots, X_n \sim \mathcal{M}$ for $n = \text{poly}(k, d)$, outputs a clustering which is correct for all of the points, with high probability. Moreover, this algorithm runs in time $\text{poly}(d, k)$.

We briefly remark that we can handle arbitrary mixing weights as well, but for simplicity we only state the theorem here in the more natural regime where all the mixing weights are not too small. We also remark that a simple corollary of this is that we can also estimate the parameters of $\mathcal{M}$ to good accuracy in polynomial time. In fact, by using our algorithm as a warm start for the method proposed in [40], we can achieve arbitrarily good accuracy for parameter estimation, in polynomial time. It is known that $\Delta = \Omega(\sqrt{\log k})$ is also necessary to achieve nontrivial parameter estimation with polynomially many samples [40], so our results for parameter estimation are also almost-optimal, in terms of the separation that they handle. We note that our formal theorems are actually stated for parameter estimation, rather than clustering, however, in light of [40], these two problems are equivalent in the regime we consider.

Our main technique (as we will discuss in more detail later) extends to beyond Gaussians, and in fact also allows us to cluster any mixture of translations of a distribution $\mathcal{D}$, so long as this distribution satisfies the Poincaré inequality, under a mild technical condition. Recall that a distribution $\mathcal{D}$ over $\mathbb{R}^d$ is said to be $\sigma$-Poincaré if for all differentiable functions $f : \mathbb{R}^d \rightarrow \mathbb{R}$, we have

$$\text{Var}_{X \sim \mathcal{D}}(f(X)) \leq \sigma^2 \cdot \mathbb{E}_{X \sim \mathcal{D}}(\|\nabla f(X)\|_2^2).$$

This condition is widely studied in probability theory, and is satisfied by many natural distribution classes. For instance, isotropic Gaussians are 1-Poincaré, and any isotropic logconcave distribution is $\psi$-Poincaré, where $\psi$ is the value of the KLS constant (see e.g. [43] for an overview of the KLS conjecture).

In fact, the family of Poincaré distributions is the most general family of distributions for which the previously mentioned Sum of Squares-based methods for clustering are known to work. For any well-separated mixture of 1-Poincaré distributions, [31] shows that one can recover the same guarantees as mentioned above: if the minimum mean separation is $\Delta = \Omega(k^{\gamma})$, then their algorithm successfully clusters the points in time $\mathcal{O}(d^{\text{poly}(1/\gamma)})$. As before, when the separation is polylogarithmic, the runtime and sample complexity of their method is once again quasipolynomial.

We show that one can improve the runtime and sample complexity to polynomial time, under two additional assumptions: first, the mixture must consist of translated versions of the same Poincaré distribution which we can get samples from, and second, the maximum and minimum separations between any two components must be polynomially related. More concretely, we show:

Theorem 1.2 (informal, see Theorem 2.3, Corollary 2.4). Let $c > 0$ be fixed but arbitrary. Let $\mathcal{D}$ be a fixed distribution with mean zero over $\mathbb{R}^d$ which is 1-Poincaré. Let $\mathcal{M}$ be a mixture of $k$ distributions where each component is of the form $\mu_i + \mathcal{D}$. Assume that the minimum mixing weight in this distribution is at least $1/\text{poly}(k)$, and moreover, assume that

$$\min_{i \neq j} \|\mu_i - \mu_j\|_2 \geq \Omega(\log^{1/2+c} k),$$

$$\max_{i \neq j} \|\mu_i - \mu_j\|_2 \leq \text{poly}\left(\min_{i \neq j} \|\mu_i - \mu_j\|_2\right).$$

Then, there is an algorithm which, given samples $X_1, \ldots, X_n \sim \mathcal{M}$ and samples $z_1, \ldots, z_n \sim \mathcal{D}$ for $n = \text{poly}(k, d)$, outputs a clustering of $X_1, \ldots, X_n$ which is correct for all of the points, with high probability. Moreover, this algorithm runs in time $\text{poly}(d, k)$. 

1249
Remark. Note that we only need the samples from $\mathcal{D}$. We do not need to actually know the distribution or access the p.d.f.

We note that separation $\Omega(\log k)$ is optimal for general Poincaré distributions, as Poincaré distributions include some distributions with exponential tails, for which $\Omega(\log k)$ separation is necessary to cluster. Thus, the separation that we require is almost optimal for general Poincaré distributions. We also note that one immediate consequence of this theorem, along with Chen’s recent breakthrough result [13] for KLS that $\psi \leq \exp(C\sqrt{\log d \log \log d})$ for some universal constant $C$, and a simple application of PCA, is that we can cluster a mixture of translates of an isotropic logconcave distribution in polynomial time, as long as the separation is $\Omega(\exp(C\sqrt{\log k \log \log k}))$.

1.2 Our Techniques

In this section, we describe how our techniques work at a high level. Our goal will be to devise a method which can, given samples $X, X' \sim \mathcal{M}$, detect whether or not $X$ and $X'$ are from the same components or from different ones.

We first make the following reduction. Notice that if $\mathcal{M}$ is a mixture with separation $\Delta$, then $(X - X')/\sqrt{\Delta}$ can be thought of as a sample from the difference mixture $\mathcal{M}'$. This is a mixture with $k$ components, each with covariance $I$. It has one component with mean zero, and the means of the remaining components all have norm at least $\sqrt{\Delta}$. Moreover, given $X, X' \sim \mathcal{M}$, we have that $X - X'$ is drawn from the zero component of the difference mixture if and only if $X, X'$ were drawn from the same component in the original mixture. Thus, to check if two samples from the original mixture $X, X'$ are from the same component, it suffices to be able to detect, given a sample from the difference mixture, whether or not this sample comes from the mean zero component or not.

In the remainder of this section, in a slight abuse of notation, we will let $\mathcal{M}$ denote the difference mixture, we will assume it has $k$ components with nonzero mean, and we will assume that all nonzero means of the difference mixture have norm at least $\Delta$. We will henceforth refer to the components with nonzero mean as the nonzero components of the mixture. This reparameterization of the problem only changes things by polynomial factors, which do not impact our qualitative results.

1.2.1 Rough clustering via implicit moment tensors. The main conceptual contribution of our paper is a novel way to implicitly access degree $t = O(\log k / \log \log k)$ moment information with polynomially many samples and time. We do so by carefully constructing an implicitly maintained projection map from $\mathbb{R}^{d'}$ down to a subspace of dimension $k$, which still preserves meaningful information about the nonzero components. For now, let us first focus on the case where the maximum norm of any mean in the difference mixture is upper bounded by $\operatorname{poly}(t)$, or equivalently, the maximum separation between any two components in the original mixture is at most polynomially larger than the minimum separation.

Low rank estimators for Hermite polynomials. Central to our methods is the $t$-th Hermite moment tensor, a classical object in probability theory, which we denote $h_t : \mathbb{R}^d \to \mathbb{R}^{d'^t}$. These are explicit polynomials, and are the natural analog of the univariate Hermite polynomials to high dimensions. A well-known fact about the Hermite polynomial tensor is that

$$\mathbb{E}_{X \sim \mathcal{N}(\mu, I)} [h_t(X)] = \mu^\otimes t.$$  

(1)

Throughout the introduction, we will treat all tensors as flattened into vectors in $\mathbb{R}^{d'}$ (in a canonical way) e.g. we view the RHS of the above as a vector in $\mathbb{R}^{d'}$. One simple but important implication of (1) is that

$$\mathbb{E}_{X \sim \mathcal{M}} [h_t(X)] = \sum_{i=1}^k w_i \mu_i^\otimes t.$$  

(2)

This fact will be crucial for us going forward.

However, a major bottleneck for algorithmically using the Hermite polynomial tensors is that we cannot write down $h_t(X)$ in polynomial time when $t$ gets large, e.g. when $t = \Omega(\log k / \log \log k)$, which is the regime we will require.

To get around this, we will use a modification of the Hermite polynomial tensors that can still be used to estimate the RHS of (2) but can also be easily manipulated implicitly. In particular, we will construct a random polynomial $R_t : \mathbb{R}^d \to \mathbb{R}^{d'}$ i.e. we can imagine $R_t$ is actually a polynomial in $x$ whose coefficients are randomly chosen. The polynomial $R_t$ satisfies two key properties.

1. (1) $R_t(x)$ is an unbiased estimator for $h_t(x)$ with bounded variance i.e. for a fixed $x$, $\mathbb{E}[R_t(x)] = h_t(x)$ where the expectation is over the random coefficients of $R_t$.

2. (2) For any choice of the randomness in the coefficients, the polynomial $R_t(x)$ can be written as a sum of polynomially many rank-1 tensors i.e. tensors of the form

$$v = v_1 \otimes \ldots \otimes v_t$$

where $v_i \in \mathbb{R}^{d'}$ for all $i = 1, \ldots, t$.

Note that the first property implies that

$$\mathbb{E}_{R_t X \sim \mathcal{N}(\mu, I)} [R_t(X)] = \mu^\otimes t,$$

and

$$\mathbb{E}_{R_t X \sim \mathcal{M}} [R_t(X)] = \sum_{i=1}^k w_i \mu_i^\otimes t.$$  

(3)

The second property is the main motivation behind the definition of $R_t$, as it implies that we can have efficient, but restricted access to it. The key point is that if our algorithm can be implemented with techniques that only require accesses to rank-1 tensors, we can implicitly access $R_t$ in polynomial time.

Implicitly finding the span of the tensorized means. Motivated by the above discussion, our goal will be to find a projection matrix $\Pi : \mathbb{R}^{d'} \to \mathbb{R}^k$ with the following properties:

1. (Efficient projection) If $v \in \mathbb{R}^{d'}$ is a rank-1 tensor, then $\Pi v$ can be evaluated in polynomial time.

2. (Zero component is small) If $X \sim \mathcal{N}(0, I)$, then $\|\Pi R_t(X)\|_2 < k^{50}$ with high probability.

3. (Nonzero components are large) If $X \sim \mathcal{M}$ is a sample from a nonzero component of the difference mixture, then $\|\Pi R_t(X)\|_2 \geq k^{100}$ with high probability.

Given such a projection map, our clustering procedure is straightforward: given two samples $X, X'$ from the original mixture, we apply the projection map to many copies of $R_t((X - X')/\sqrt{\Delta})$, and cluster them in the same component if and only if their projected norm is small on average. We show that the above properties, as
well as some facts about the concentration of $R_t$, imply that this clustering algorithm succeeds with high probability, assuming we have access to $\Pi$.

It thus remains how to construct $\Pi$. In fact, there is a natural candidate for such a map. Let $\mu_1, \ldots, \mu_k$ denote the means of the nonzero components, and let

$$S_t = \text{span} \left\{ \{\mu_i^1 \}_{i=1}^k \right\}.$$  

If we could find the projection $\Gamma_t : \mathbb{R}^{d^t} \rightarrow \mathbb{R}^k$ onto the subspace $S_t$, it can be verified that this projection map would satisfy Conditions (2) and (3).

Moreover, there is a natural estimator for this subspace. In particular (5) implies that $\mathbb{E}[R_{2s}(X)]$ rearranged as a $d^t \times d^t$ matrix in a canonical way is exactly

$$\sum_{i=1}^{k} w_i (\mu_i^{(s)}) (\mu_i^{(s)})^T.$$  

Notice that this matrix is rank $k$, and moreover, the span of its nonzero eigenvectors is exactly $S_t$. Consequently, if we could estimate this quantity, then find the projection onto the span of the top $k$ eigenvectors, we would be done.

As alluded to earlier, the difficulty is that doing this naively would not be efficient; writing down any of these objects would take quasipolynomial time. Instead, we seek to find an implicit representation of $\Gamma_t$ with the key property that it can be applied to rank-1 tensors in polynomial time.

We will do so by iteratively building an approximation to this subspace. Namely, we give a method which, given a good approximation to $\Gamma_{t-1} : \mathbb{R}^{d^{t-1}} \rightarrow \mathbb{R}^k$ which can be efficiently applied to flattenings of rank-1 tensors, constructs a good approximation to $\Gamma_{t}$ with the same property. We do so by obtaining a good approximation to the $dk \times dk$ sized matrix

$$M_t = \sum_{i=1}^{k} w_i (\mu_i \otimes \Gamma_{t-1} - \mu_i^{(s-1)}) (\mu_i \otimes \Gamma_{t-1} - \mu_i^{(s-1)})^T. \quad (4)$$

Notice that $M_t$ has rank $k$, and moreover, the span of its $k$ largest eigenvectors is equal to the span of $\{\mu_i \otimes \Gamma_{t-1} - \mu_i^{(s-1)}\}_{i=1}^{k}$. Therefore, if we let $\Pi_t : \mathbb{R}^{dk} \rightarrow \mathbb{R}^k$ denote the projection onto the span of the $k$ largest eigenvectors of $M_t$, then one can easily verify that

$$\Gamma_t = \Pi_t (I \otimes \Gamma_{t-1}). \quad (5)$$

Moreover, if $\Gamma_{t-1}$ can be efficiently applied to flattenings of rank-1 tensors in $\mathbb{R}^{d^{t-1}}$, then the form of (3) immediately implies that $\Gamma_t$ can also be applied efficiently to flattenings of rank-1 tensors in $\mathbb{R}^{d^t}$.

It remains to demonstrate how to efficiently approximate $M_t$, given $\Gamma_{t-1}$. There is again a fairly natural estimator for this matrix. Namely, each component of the sum in (4) can be formed by rearranging the length-$(dk)^2$ vector

$$(I \otimes \Gamma_{t-1})^{\otimes^2} \mu_i^{\otimes^2} = \mathbb{E}_{R_{2s}X-N(\mu_i,t)} [(I \otimes \Gamma_{t-1})^{\otimes^2} R_{2s}(X)].$$

This implies that the overall matrix is the rearrangement of the length-$(dk)^2$ vector

$$\sum_{i=1}^{k} w_i (I \otimes \Gamma_{t-1})^{\otimes^2} \mu_i^{\otimes^2} = \mathbb{E}_{R_{2s}X-M} [(I \otimes \Gamma_{t-1})^{\otimes^2} R_{2s}(X)] \quad (6)$$

into a $dk \times dk$ sized matrix in the canonical way. In particular, this implies that $\mathbb{E}[R_{2s}(X)]$ rearranged as a $d^t \times d^t$ matrix in a canonical way is exactly

$$\mathbb{E}[R_{2s}(X)] = \mathbb{E}_{R_{2s}X-M} [(I \otimes \Gamma_{t-1})^{\otimes^2} R_{2s}(X)].$$

Putting it all together, this allows us to approximate $M_t$ efficiently given $\Gamma_{t-1}$, which, by (5), gives us the desired expression for $\Gamma_t$. Iterating this procedure gives us a way to estimate $\Gamma_t$ as a sequence of nested projection maps, i.e.

$$\Gamma_t = \Pi_t (I \otimes \Gamma_{t-1} (I \otimes \ldots )).$$

where we can compute $\Pi_1, \ldots, \Pi_t$ efficiently, given samples from $M$. This form allows us to evaluate $\Gamma_t$ efficiently on any flattening of a rank-1 tensor, thus satisfying Condition (1), and we previously argued that $\Gamma_t$ satisfies Conditions (2) and (3). Combining all of these ingredients gives us our clustering algorithm, when the minimum and maximum separations are at most polynomially separated.

**Implicit moment tensors for Poincaré distributions.** So far, we have only discussed how to do this implicit moment estimation for isotropic Gaussians. It turns out that all of the quantities discussed above have very natural analogues for any Poincaré distribution. For instance, given any Poincaré distribution $D$ with zero mean, there is an explicit polynomial tensor we call the $D$-adjusted polynomial $\Gamma_{t,D}$ (see Section 4) that essentially satisfies all the same properties that we needed above. If we use these polynomials instead of the Hermite polynomial tensor, it turns out that all of these proofs directly lift to any Poincaré distribution. In fact, in the actual technical sections, we directly work with arbitrary Poincaré distributions, as everything is stated very naturally there. The resulting clustering algorithm immediately gives us Theorem 2.3.

**Sample complexity.** Previous approaches always needed to estimate high degree moment tensors, and as a result, their sample complexity was quasipolynomial. While we may also need to estimate fairly high degree polynomials, notice that all quantities that we will deal with will be polynomially bounded. This is because we can terminate our procedure at any $t$ which satisfies Conditions (2) and (3). Therefore, all the quantities that we need are polynomially large. As a result, one can verify that the polynomials we construct will also only ever have polynomially large range, with high probability. Therefore, all quantities we need to estimate can be estimated using polynomially many samples. We defer the detailed proofs outlined in this discussion to Sections 4, 5, 6 and 7. Note that in those sections, we work with a general Poincaré distribution but the outline follows the description here.

1.2.2 **Fine-grained clustering for Gaussians.** We now discuss how to handle general mixtures of isotropic Gaussians, without any assumption on the maximum separation. The problem with applying the implicit moment estimation method outlined above to this general setting is that the signal from the components in the
difference mixture with relatively small mean will be drowned out by the signal from the components with much larger norm. Consequently, we can reliably cluster points from the components with large mean, but we could obtain an imperfect clustering for some components with somewhat smaller mean, and we will be unable to detect components with very small mean.

To overcome this, we devise a recursive clustering strategy. One somewhat simple approach is as follows. We first use our rough clustering algorithm described above to find a “signal direction” $v \in \mathbb{R}^d$. This direction will have the property that there is a pair of well-separated means along this direction. Thus, if we project the data points on this direction, and take only points which lie within a randomly chosen small interval on this interval, we can guarantee that with reasonable probability, we only accept points from at most half of the components of the mixture. Of course, after restricting to this interval, the resulting distribution is no longer a mixture of Gaussians. However, if we consider the projection of these accepted points to the subspace orthogonal to $v$, the resulting distribution will again be a mixture of fewer isotropic Gaussians. We can then recurse on this mixture with fewer components. Here, we are crucially using the fact that isotropic Gaussians remain isotropic Gaussians after slicing and projecting orthogonally.

While this strategy described above, when implemented carefully, would work down to $\Delta = \text{poly}(\log k)$, it would not be able to achieve the nearly optimal separation in Theorem 2.5. To achieve the nearly optimal separation, there are several more technical details that need to be dealt with and thus there will be several additional steps in the algorithm. Due to space constraints, the fine-grained clustering steps are omitted here and deferred to the full version.

1.3 Related Work

The literature on mixture models—and Gaussian mixture models in particular—dates back to seminal work of Pearson [38] and is incredibly vast. For conciseness, we will focus only on the most related papers here. Our results are most closely related to the aforementioned line of work on studying efficient algorithms for clustering and parameter estimation under mean-separation conditions [5, 14, 15, 19, 21, 26, 31, 40, 44]. However, none of these algorithms can handle polylogarithmic separation in polynomial time.

We also note that a number of papers also generalize from mixtures of Gaussians to mixtures of more general classes of distributions [3, 26, 31, 32, 36]. These algorithms fall into two classes: either they require separation which is at least $\Omega(k^{1/2})$ or even larger, but they can handle general subgaussian distributions, or they require more structure on the higher moments of the distribution, but they can tolerate much less separation. The most general condition under which the latter is known to work is the condition commonly referred to as certifiable hypercontractivity, which roughly states that the Sum-of-Squares hierarchy can certify that the distribution has bounded tails. While there is no complete characterization of what distributions satisfy this condition, the most general class of distributions for which it is known to hold is the class of Poincaré distributions [31], which is also the class of distributions we consider here.

Another line of work focuses on parameter estimation for mixtures models without separation conditions [7, 19, 25, 30, 37]. These papers typically make much weaker assumptions on the components, namely, that they are statistically distinguishable, and the goal is to recover the parameters of the mixture, even in settings where clustering is impossible. However, typically, these algorithms incur a sample complexity and runtime which is exponential in the number of components; indeed, [25] demonstrate that this is tight, even in one dimension, when the Gaussians can have different variances.

To circumvent this, researchers have also considered the easier notion of proper or semi-proper learning, where the goal is to output a mixture of Gaussians which is close to the unknown mixture in statistical distance. A learning algorithm is said to be proper if its output is a mixture of $k$ Gaussians, where $k$ is the number of components in the true mixture, and semi-proper if it outputs a mixture of $k' \geq k$ Gaussians. While the sample complexity of proper learning is polynomial in all parameters, all known algorithms still incur a runtime which is exponential in $k$, even in the univariate setting [2, 6, 16, 22, 34]. When the hypothesis is only constrained to be semi-proper, polynomial time algorithms are known in the univariate setting [9, 18, 33], but these do not extend to high dimensional settings. In the more challenging high dimensional regime, a remarkable recent result of [19] demonstrate that for a mixture of isotropic Gaussians, one can achieve semi-proper learning with quasipolynomial sample complexity and runtime. They do this by explicitly constructing a small cover for the candidate means, by techniques inspired by algebraic geometry.

An even weaker goal that has been commonly considered is that of density estimation, where the objective is to output any hypothesis which is close to the unknown mixture in statistical distance. Efficient (in fact, nearly optimal) algorithms are again known for this problem in low dimensions, see e.g. [1, 11, 12, 18], but as was the case with proper learning, these techniques do not extend nicely to high dimensional settings.

An alternate assumption that has been considered in the literature is that the means satisfy some algebraic non-degeneracy assumptions. For instance, such assumptions are satisfied in smoothed analysis settings [4, 8, 23, 29]. Often in these settings, access to constant order moments suffice (e.g. 3rd or 4th moments), although we note that [8] is a notable exception. In contrast, we do not make any non-degeneracy assumptions, and our methods need to access much higher moments.

We also mention a line of work studying the theoretical behavior of the popular expectation-maximization (EM) algorithm for learning mixtures of Gaussians [15, 17, 46]. However, while the above works can prove that the dynamics of EM converge in limited settings, it is known that EM fails to converge in general, even for mixtures of 3 Gaussians [17, 45].

Finally, we note that our work bears some vague resemblance to the general line of work that uses spectral-based methods to speed up Sum-of-Squares (SoS) algorithms. Spectral techniques have been used to demonstrate to speed up SoS-based algorithms in various settings such as tensor decomposition [27, 28, 35, 41] and refuting random CSPs [39]. Similarly, it has been observed that in some settings, SoS-based algorithms can be sped up, when the SoS proofs are much smaller than the overall size of the program [24, 42].
algorithm shares some qualitative similarities with some of these approaches—for instance, it is based on a (fairly complicated) spectral algorithm. However, we do not know of a concrete connection between our algorithm and this line of work. It is possible, for instance, that our algorithm could be interpreted as extracting a specific randomized polynomially-sized SoS proof of identifiability, but we leave further investigations of this to future work.

2 FORMAL PROBLEM SETUP AND OUR RESULTS

In this section, we formally define the problems we consider, and state our formal results. For the remainder of this paper, we will always let \( \| \cdot \| \) denote the \( \ell_2 \) norm.

2.1 Clustering Mixtures of Poincare Distributions

The general problem that we study involves clustering mixtures of Poincare distributions. We begin with a few definitions.

Definition 2.1 (Poincare Distribution). For a parameter \( \sigma \), we say a distribution \( \mathcal{D} \) on \( \mathbb{R}^d \) is \( \sigma \)-Poincare if for all differentiable functions \( f : \mathbb{R}^d \to \mathbb{R} \),

\[
\text{Var} \left( f(z) \right) \leq \sigma^2 \mathbb{E}_{z \sim \mathcal{D}} \left[ \| \nabla f(z) \|^2 \right].
\]

Definition 2.2. Let \( \mathcal{D} \) be a distribution on \( \mathbb{R}^d \). We use \( \mathcal{D}(\mu_1) \) for \( \mu_1 \in \mathbb{R}^d \) to denote the distribution obtained by shifting \( \mathcal{D} \) by the vector \( \mu_1 \).

We assume that there is some \( \sigma \)-Poincare distribution \( \mathcal{D} \) on \( \mathbb{R}^d \) that we have sample access to. Since everything will be scale invariant, it will suffice to focus on the case \( \sigma = 1 \). For simplicity we assume that \( \mathcal{D} \) has mean 0 (it is easy to reduce to this case when we can simply estimate the mean of \( \mathcal{D} \) and subtract it out). We also assume that we have sample access to a mixture

\[
\mathcal{M} = w_1 \mathcal{D}(\mu_1) + \cdots + w_d \mathcal{D}(\mu_d)
\]

where the mixing weights \( w_1, \ldots, w_d \) and means \( \mu_1, \ldots, \mu_d \) are unknown. We will assume that we are given a lower bound on the mixing weights \( w_{\text{min}} \). We consider the setting where there is some minimum separation between all pairs of means \( \mu_i, \mu_j \) so that the mixture is clusterable. In the proceeding sections, when we say an event happens with high probability, we mean that the failure probability is smaller than any inverse polynomial in \( \log(1/\delta) \).

Our main theorem is stated below.

Theorem 2.3. Let \( \mathcal{D} \) be a 1-Poincare distribution on \( \mathbb{R}^d \). Let

\[
\mathcal{M} = w_1 \mathcal{D}(\mu_1) + \cdots + w_d \mathcal{D}(\mu_d)
\]

be a mixture of translated copies of \( \mathcal{D} \). Let \( w_{\text{min}}, s \) be parameters such that \( w_i \geq w_{\text{min}} \) for all \( i \) and \( \| \mu_i - \mu_j \| \geq s \) for all \( i \neq j \). Let \( \alpha = \left( w_{\text{min}} / k \right)^{O(1)} \) be some desired accuracy (that is inverse polynomial)\(^3\). Assume that

\[
s \geq (\log(k/w_{\text{min}}))^{1+c}
\]

for some \( 0 < c < 1 \). Also assume that

\[
\max \| \mu_i - \mu_j \| \leq s^C
\]

for some \( C \). There is an algorithm that takes

\[
n = \text{poly}(kd/(w_{\text{min}}\alpha)^{(C/c)})
\]

samples from \( \mathcal{M} \) and \( \mathcal{D} \) and runs in \( \text{poly}(n) \) time and with high probability, outputs estimates 

\[
\tilde{\mu}_1, \ldots, \tilde{\mu}_k, \tilde{\mu}_1, \ldots, \tilde{\mu}_k
\]

such that for some permutation \( \pi \) on \( [k] \),

\[
|\mu_i - \tilde{\mu}_{\pi(i)}| \leq \alpha, \| \mu_i - \tilde{\mu}_{\pi(i)} \| \leq \alpha
\]

for all \( i \).

Remark. If we could remove the assumption \( \| \mu_i - \mu_j \| \leq s^C \), then we would get a complete polynomial time learning result. Still, our learning algorithm works in polynomial time for mixtures where the maximum separation is polynomially bounded in terms of the minimum separation.

An immediate consequence of Theorem 2.3 is that we can cluster samples from the mixture with accuracy better than any inverse polynomial.

Corollary 2.4. Under the same assumptions as Theorem 2.3, we can recover the ground-truth clustering of the samples with high probability i.e. we output \( k \) clusters \( S_1, \ldots, S_k \) such that for some permutation \( \pi \) on \( [k] \), the set \( S_{\pi(i)} \) consists precisely of the samples from the component \( \mathcal{D}(\mu_i) \) for all \( i \).

2.2 Clustering Mixtures of Gaussians

In the case where the distribution \( \mathcal{D} \) in the setup in Section 2.1 is a Gaussian, we can obtain a stronger result that works in full generality, without any assumption about the maximum separation. The result for Gaussians also works with a smaller separation of \( (\log(k/w_{\text{min}}))^{1+c} \) which, as mentioned before, is essentially optimal for Gaussians.

Theorem 2.5. Let \( \mathcal{M} = w_1 N(\mu_1, I) + \cdots + w_k N(\mu_k, I) \) be an unknown mixture of Gaussians in \( \mathbb{R}^d \) such that \( w_i \geq w_{\text{min}} \) for all \( i \) and \( \| \mu_i - \mu_j \| \geq (\log(k/w_{\text{min}}))^{1+c} \) for some constant \( c > 0 \). Then for any desired (inverse polynomial) accuracy \( \alpha \geq \left( w_{\text{min}} / k \right)^{O(1)} \) given \( n = \text{poly}(dk/(w_{\text{min}}\alpha)^{(C/c)}) \) samples and \( \text{poly}(n) \) runtime, there is an algorithm that with high probability outputs estimates \( \tilde{\mu}_1, \ldots, \tilde{\mu}_k \) and \( \tilde{\mu}_1, \ldots, \tilde{\mu}_k \) such that for some permutation \( \pi \) on \( [k] \), we have

\[
|\mu_i - \tilde{\mu}_{\pi(i)}| \leq \alpha, \| \mu_i - \tilde{\mu}_{\pi(i)} \| \leq \alpha
\]

for all \( i \in [k] \).

Again, once we have estimated the parameters of \( \mathcal{M} \), it is easy to cluster samples from \( \mathcal{M} \) into each of the components with accuracy better than any inverse polynomial.

Corollary 2.6. Under the same assumptions as Theorem 2.5, with high probability, we can recover the ground-truth clustering of the samples i.e. we output \( k \) clusters \( S_1, \ldots, S_k \) such that for some permutation \( \pi \) on \( [k] \), the set \( S_{\pi(i)} \) consists precisely of the samples from the component \( N(\mu_i, I) \) for all \( i \).
Note that throughout this paper, we do not actually need to know the true number of components \( k \). All of the algorithms that we write will work if instead of the number of components being \( k \), the number of components is upper bounded by \( k \) i.e. we only need to be told an upper bound on the number of components. In fact, we can simply use \( 1/w_{\text{min}} \) as the upper bound on the number of components.

2.3 Organization

In Section 3, we introduce basic notation and prove a few basic facts that will be used later on.

Clustering Test: In Sections 4 - 7, we develop our key clustering test i.e. we show how to test if two samples are from the same component or not. Note that throughout these sections, when we work with a mixture

\[
M = w_1 \mathcal{D}(\mu_1) + \cdots + w_k \mathcal{D}(\mu_k),
\]

this will correspond to the ‘difference mixture’ of the mixture that we are trying to learn and thus our goal will be to test whether a sample came from a component with \( \mu_j = 0 \) or \( \mu_j \) far from 0.

In Section 4, we discuss how to construct estimators for the moments of a Poincare distribution that can be manipulated implicitly. In the end, we construct a random polynomial \( R_t \) such that

\[
\mathbb{E}_{x \sim \mathcal{D}(\mu)} [R_t(x)] = \mu^{\otimes t}
\]

and such that \( R_t \) can be written as the sum of polynomially many rank-1 tensors (see Corollary 4.9). In Section 5, we describe our iterative projection technique and explain how it can be used to implicitly store and apply a projection map \( \Gamma : \mathbb{R}^{d^k} \rightarrow \mathbb{R}^k \) in polynomial time to rank-1 tensors. In Section 6, we combine the techniques in Section 4 and Section 5 to achieve the following: given samples from

\[
M = w_1 \mathcal{D}(\mu_1) + \cdots + w_k \mathcal{D}(\mu_k)
\]

we can find a \( k \times d^t \) projection matrix \( \Gamma_t \) (where \( t \sim \log k / \log \log k \)) whose row span essentially contains all of \( \mu_1^{\otimes t}, \ldots, \mu_k^{\otimes t} \) (see Lemma 6.5). Finally, in Section 7, we use the projection map computed in the previous step and argue that if \( x \sim \mathcal{D}(0) \) then \( \|\Gamma_t R_t(x)\| \) is small and if \( x \sim \mathcal{D}(\mu) \) for \( \mu \) far from 0, then \( \|\Gamma_t R_t(x)\| \) is large with high probability. Thus, to test a sample \( x \), it suffices to measure the length of \( \|\Gamma_t R_t(x)\| \).

Main Result for Mixtures of Poincare Distributions: In Section 8, we prove Theorem 2.3, our main result for mixtures of Poincare distributions. It will follow fairly easily from the guarantees of the clustering test in Section 7.

Main Result for Mixtures of Gaussians: Proving our main result for mixtures of Gaussians requires some additional work although all of the machinery from Sections 4 - 7 can still be used. Roughly, the two additional ingredients are

1. Quantitatively stronger versions of the bounds in Sections 4 - 7 that exploit special properties of Gaussians in order to get down from \( \log^k k \) to \( \log^{1/2+\epsilon} k \) separation
2. A recursive clustering procedure that allows us to eliminate the assumption about the maximum separation.

Due to space constraints these parts are omitted here and deferred to the full version. While the additional pieces require working through many technical details, the core of our learning algorithm is still built from the machinery in Sections 4 - 7.

3 PRELIMINARIES

We now introduce some notation that will be used throughout the paper. We use \( I_n \) to denote the \( n \times n \) identity matrix. For matrices \( A, B \) we define \( A \otimes B \) to be their Kronecker product. This is to avoid confusion with our notation for tensor products. For a tensor \( T \), we use flatten \((T)\) to denote the flattening of \( T \) into a vector. We assume that this is done in a canonical way throughout this paper.

3.1 Manipulating Tensors

We will need to do many manipulations with tensors later on so we first introduce some notation for working with tensors.

Definition 3.1 (Tensor Notation). For an order-\( t \) tensor, we index its dimensions \((1, 2, \ldots, t)\). For a partition of \( [t] \) into subsets \( S_1, \ldots, S_d \) and tensors \( T_1, \ldots, T_d \) of orders \(|S_1|, \ldots, |S_d|\) respectively we write

\[
T_1^{(S_1)} \otimes \cdots \otimes T_d^{(S_d)}
\]

to denote the tensor obtained by taking the tensor product of \( T_1 \) in the dimensions indexed by \( S_1 \), \( T_2 \) in the dimensions indexed by \( S_2 \), and so on for \( T_3, \ldots, T_d \).

Definition 3.2. For a vector \( x \) (viewed as an order-1 tensor), we will use the shorthand \( x^{\otimes S} \) to denote \((x^{\otimes |S|})^{(S)}\) (i.e. the product of copies \( x \) in dimensions indexed by elements of \( S \)). For example,

\[
x^{\otimes \{1,3\}} \otimes y^{\otimes \{2,4\}} = x \otimes y \otimes x \otimes y.
\]

Definition 3.3 (Tensor Slicing). For an order-\( t \) tensor \( T \), we can imagine indexing its entries with indices \((\eta_1, \ldots, \eta_t) \in [d_1] \times \cdots \times [d_t]\) where \( d_1, \ldots, d_t \) are the dimensions of \( T \). We use the notation

\[
T_{\eta_1 = b_1, \ldots, \eta_t = b_t}
\]

to denote the slice of \( T \) entries whose indices satisfy the constraints \( \eta_1 = b_1, \ldots, \eta_t = b_t \).

Definition 3.4 (Unordered Partitions). We use \( Z_t(S) \) to denote all partitions of \( S \) into unordered, possibly empty, parts.

Remark. Note the partitions are not ordered so \( \{1\} \neq \{2\} \) is the same as \( \{2\} \neq \{1\} \).

Definition 3.5. For a collection of sets \( S_1, \ldots, S_t \), we define

\[
C(\{S_1, \ldots, S_t\})
\]

to be the number of sets among \( S_1, \ldots, S_t \) that are nonempty.

Definition 3.6 (Symmetrization). Let \( A_1, \ldots, A_n \) be tensors such that \( A_i \) is an order \( a_i \) tensor having dimensions \( d \times \cdots \times d \) for some \( a_i \). We define

\[
\sum_{\text{sym}} (A_1 \otimes \cdots \otimes A_n) = \sum_{S \subseteq [n]} A_1^{(S_1)} \otimes \cdots \otimes A_n^{(S_n)}.
\]

In other words, we sum over all ways to tensor \( A_1, \ldots, A_n \) together to form a tensor of order \( a_1 + \cdots + a_n \).
3.2 Properties of Poincare Distributions

Here we state a few standard facts about Poincare distributions that will be used later on.

Fact 3.7. Poincare distributions satisfy the following properties:

- Direct Products: If \( D_1 \) and \( D_2 \) are \( \sigma \)-Poincare distributions then their product \( D_1 \times D_2 \) is \( \sigma \)-Poincare.
- Linear Mappings: If \( D \) is \( \sigma \)-Poincare and \( A \) is a linear mapping then the distribution \( Ax \) for \( x \sim D \) is \( \sigma \|A\|_{op} \)-Poincare.
- Concentration: If \( D \) is \( \sigma \)-Poincare then for any \( L \)-Lipschitz function \( f \) and any parameter \( t \geq 0 \), we have

\[
\Pr_{z \sim D} [ |f(z) - E[f(z)]| \geq t ] \leq 6e^{-t^2/(\sigma L)}.
\]

The following concentration inequality for samples from a Poincare distribution is also standard.

Claim 3.8. Let \( D \) be a distribution in \( \mathbb{R}^d \) that is \( 1 \)-Poincare. Let \( 0 < \epsilon < 0.1 \) be some parameter. Given \( n \geq (d/\epsilon)^8 \) independent samples \( z_1, \ldots, z_n \sim D \), with probability at least \( 1 - 2^{-d/\epsilon} \), we have

\[
\frac{1}{n} \sum_{i=1}^{n} z_i - E[z_i] \leq \epsilon.
\]

3.3 Basic Observations

Before we begin with the main proofs of Theorem 3.2 and Theorem 2.5, it will be useful to make a few simple reductions that allow us to make the following simplifying assumptions:

- Means Polynomially Bounded: For all \( i \), we have \( |\mu_i - \mu_j| \leq O((k/w_{\min})^2) \), and
- Dimension Not Too High: We have \( d \leq k \).

Reducing to the case when the above assumptions hold is straightforward and omitted due to space constraints. The reductions work in both settings (general Poincare distributions and Gaussians) so in future sections, we will be able to work assuming that the above properties hold.

4 MOMENT ESTIMATION

We will now work towards proving our result for Poincare distributions. A key ingredient in our algorithm will be estimating the moment tensor of a mixture \( M \) i.e. for an unknown mixture

\[
M = w_1 D(\mu_1) + \cdots + w_k D(\mu_k)
\]

we would like to estimate the tensor

\[
w_1 \mu_1^t + \cdots + w_k \mu_k^t
\]

for various values of \( t \) using samples from \( M \). Naturally, it suffices to consider the case where we are given samples from \( D(\mu) \) for some unknown \( \mu \) and our goal is to estimate the tensor \( \mu_{\mu_1}^t \). For our full algorithm, we will need to go up to \( t \sim \log k / \log \log k \) but of course for such \( t \), our estimate has to be implicit because we cannot write down the full tensor in polynomial time. In this section, we address this task by constructing an unbiased estimator with bounded variance that can be easily manipulated implicitly.

We make the following definition to simplify notation later on.

Definition 4.1. For integers \( t \) and a distribution \( D \), we define the tensor

\[
D_{t, D} = \mathbb{E}_{z \sim D} [ z^{\otimes t} ].
\]

We will drop the subscript \( D \) when it is clear from context.

4.1 Adjusted Polynomials

First, we just construct an unbiased estimator for \( \mu_{\mu}^t \) (without worrying about making it implicit). This estimator is given in the definition below.

Definition 4.2. Let \( D \) be a distribution on \( \mathbb{R}^d \). For \( x \in \mathbb{R}^d \), define the polynomials \( P_{t, D}(x) \) for positive integers \( t \) as follows. \( P_{0, D}(x) = 1 \) and for \( t \geq 1 \),

\[
P_{t, D}(x) = x^{\otimes t} - \sum_{s = 0}^{t-1} D_{s, D} \otimes P_{t-s, D}(x).
\]

We call \( P_{t, D} \) the \( D \)-adjusted polynomials and will sometimes drop the subscript \( D \) when it is clear from context.

Through direct computation, we can verify that the \( D \)-adjusted polynomials indeed give an unbiased estimator for \( \mu_{\mu}^t \) when given samples from \( D(\mu) \).

Claim 4.3. For any \( \mu \in \mathbb{R}^d \),

\[
\mathbb{E}_{z \sim D(\mu)} [ P_{t, D}(z) ] = \mu_{\mu}^t.
\]

4.2 Variance Bounds for Poincare Distributions

In the previous section, we showed that the \( D \)-adjusted polynomials give an unbiased estimator for \( \mu_{\mu}^t \). We now show that they also have bounded variance when \( D \) is Poincare. This will rely on the following claim which shows that the \( D \)-adjusted polynomials recur under differentiation.

Claim 4.4. Let \( D \) be a distribution on \( \mathbb{R}^d \). Then

\[
\frac{\partial P_{t, D}(x)}{\partial x_i} = \sum e_i \otimes P_{t-1, D}(x)
\]

where we imagine \( x = (x_1, \ldots, x_d) \) so \( x_i \) is the \( i \)-th coordinate of \( x \) and

\[
e_i = (0, \ldots, 0, 1, \ldots, 0)
\]

denotes the \( i \)-th coordinate basis vector.

Now, by using the Poincare property, we can prove a bound on the variance of the estimator \( P_{t, D}(x) \).

Claim 4.5. Let \( D \) be a distribution on \( \mathbb{R}^d \) that is \( 1 \)-Poincare. Let \( v \in \mathbb{R}^d \) be a vector. Then

\[
\mathbb{E}_{z \sim D(\mu)} [ |v \cdot flatten(P_{t, D}(z))|^2 ] \leq (\|v\|^2 + t^2)^t \|v\|^2.
\]
4.3 Efficient Implicit Representation

In the previous section, we showed that for $x \sim D(\mu)$ for unknown $\mu$, $P_{\mu}D(x)$ gives us an unbiased estimator of $\mu^{ot}$ with bounded variance. Still, it is not feasible to actually compute $P_{\mu}D(x)$ in polynomial time because we cannot write down all of its entries and there is no nice way to implicitly work with terms such as $D_{\mu}x$ that appear in $P_{\mu}D(x)$. In this section, we construct a modified estimator that is closely related to $P_{\mu}D(x)$ but is also easy to work with implicitly because all of the terms will be rank-1 i.e. of the form $v_1 \otimes \cdots \otimes v_k$ for some vectors $v_1, \ldots, v_k \in \mathbb{R}^d$. Throughout this section, we will assume that the distribution $D$ that we are working with is fixed and we will drop it from all subscripts as there will be no ambiguity.

Roughly, the way that we construct this modified estimator is that we take multiple variables $x_1, \ldots, x_k \in \mathbb{R}^d$. We start with $P_t(x_1)$. We then add various products $P_{a_1}(x_1) \otimes \cdots \otimes P_{a_t}(x_t)$ to it in a way that when expanded as monomials, only the leading terms, which are rank-1 since they are a direct product of the form $x_1^{a_1} \otimes \cdots \otimes x_t^{a_t}$, remain. If we then take $x_1 \sim D(\mu)$ and $x_2, \ldots, x_t \sim D$, then Claim 4.3 will immediately imply that the expectation is $\mu^{ot}$. The key properties are stated formally in Corollary 4.10 and Corollary 4.11.

The following polynomial and its properties will be the key in our construction.

**Definition 4.6.** For $x_1, \ldots, x_t \in \mathbb{R}^d$, define the polynomial $Q_t(x_1, \ldots, x_t)$ as

$$Q_t(x_1, \ldots, x_t) = \sum_{S_1, \ldots, S_t = \emptyset} (-1)^{|S_1| + \cdots + |S_t|} \left( \prod_{i=1}^t (P_{S_i}(x_i)) \right) \otimes \cdots \otimes \left( \prod_{i=1}^t (P_{S_i}(x_i)) \right).$$

**Lemma 4.7.** All nonzero monomials in $Q_t$ either have total degree $t$ in the variables $x_1, \ldots, x_t$ or are constant.

Now, we can easily eliminate the constant term by subtracting off $Q_t(x_{t+1}, \ldots, x_k)$ for some additional variables $x_{t+1}, \ldots, x_k$ and we will be left with only degree-$t$ terms. It will be immediate that the degree-$t$ terms are all rank-1 and this will give us an estimator that can be efficiently manipulated implicitly.

**Definition 4.8.** For $x_1, \ldots, x_{2t} \in \mathbb{R}^d$, define the polynomial $R_t(x_1, \ldots, x_{2t}) = -Q_t(x_1, \ldots, x_t) + Q_t(x_{t+1}, \ldots, x_{2t})$.

**Corollary 4.9.** We have the identity $R_t(x_1, \ldots, x_{2t}) = \sum_{S_1, \ldots, S_t = \emptyset} (-1)^{|S_1| + \cdots + |S_t|} \left( \prod_{i=1}^t (P_{S_i}(x_i)) \right) \otimes \cdots \otimes \left( \prod_{i=1}^t (P_{S_i}(x_i)) \right)$.

Corollary 4.9 gives us a convenient representation for working implicitly with $R_t(x_1, \ldots, x_{2t})$. We now show why this polynomial is actually useful. In particular, we show that for $x_1 \sim D(\mu)$ and $x_2, \ldots, x_{2t} \sim D$, $R_t(x_1, \ldots, x_{2t})$ is an unbiased estimator of $\mu^{ot}$ and furthermore that its variance is bounded. These properties will follow directly from the definitions of $R_t, Q_t$ combined with Claim 4.3 and Claim 4.5.

**Corollary 4.10.** We have

$$E \left[ z_1 \sim D(\mu), \ldots, z_{2t} \sim D \mid R_t(z_1, \ldots, z_{2t}) \right] = \mu^{ot}$$

and for fixed $z_1$, we have

$$E \left[ z_2, \ldots, z_{2t} \sim D \mid R_t(z_1, \ldots, z_{2t}) \right] = P_t(z_1).$$

**Corollary 4.11.** Let $D$ be a distribution that is 1-Poincare. We have

$$E \left[ \text{flatten}(R_t(z_1, \ldots, z_{2t}))^{ot} \right] \leq (20t)^{2t} (\|\mu\|^{2t} + 1) I_d.$$
Note that in our paper, $\Pi_1, \ldots, \Pi_t$ will always have orthonormal rows so $\Pi_{1,\ldots,\Pi_t}$ always does as well. This fact will often be used without explicitly stating it. The key point about the construction of $\Pi_{1,\ldots,\Pi_t}$ is that instead of storing a full $c_t \times d^t$-sized matrix, it suffices to store the individual matrices $\Pi_1, \ldots, \Pi_t$ which are all polynomially sized. The next important observation is that for certain vectors $v \in \mathbb{R}^d$, the expression $\Pi_{1,\ldots,\Pi_t}v$ can be computed efficiently. This is shown in the following claim.

**Claim 5.3.** Let $c_0 = 1$ and $c_1, \ldots, c_t$ be positive integers. Let $\Pi_1 \in \mathbb{R}^{c_1 \times d_0}, \Pi_2 \in \mathbb{R}^{c_2 \times d_1}, \ldots, \Pi_t \in \mathbb{R}^{c_t \times d_{t-1}}$ be matrices. Let $v \in \mathbb{R}^d$ satisfy $v = \text{flatten}(v_1 \otimes \cdots \otimes v_1)$ for some $v_1, \ldots, v_t \in \mathbb{R}^d$. Then in $\text{poly}(d, t, \max(c_i))$ time, we can compute $\Gamma_{\Pi_{1,\ldots,\Pi_t}}v$.

**Proof.** We will prove the claim by induction on $t$. For each $t' = 1, 2, \ldots, t$, we compute

$$
\Pi_{1,\ldots,\Pi_t}v = \text{flatten}(v_1 \otimes \cdots \otimes v_1)
$$

by implicitly estimating the moment tensor. For the base case of the induction, it is clear from context. For the induction step, note that

$$
\Pi_{1,\ldots,\Pi_t}v = \Pi_{t',\ldots,\Pi_t}v = \text{flatten}\left(\Pi_{t',\ldots,\Pi_t}v \otimes \cdots \otimes \Pi_{t',\ldots,\Pi_t}v\right).
$$

It is clear that this computation can be done in $\text{poly}(d, t, \max(c_i))$ time so iterating this operation completes the proof. \hfill \blacksquare

Throughout our paper, we will only compute nested projections of the above form so it will be easy to verify that all steps can be implemented in polynomial time.

## 6 IMPLICITLY ESTIMATING THE MOMENT TENSOR

In this section, we combine the iterative projection techniques from Section 5 with the estimators from Section 4 to show how to implicitly estimate the moment tensor given sensor access to the distribution $\mathcal{D}$ and the mixture

$$
M = w_1 \mathcal{D}(\mu_1) + \cdots + w_k \mathcal{D}(\mu_k).
$$

By implicitly estimate, we mean that we will compute projection matrices $\Pi_1, \ldots, \Pi_t$ such that the row-span of $\Pi_{1,\ldots,\Pi_t}$ essentially contains all of the flattenings of $\mu_1^{\otimes t}, \ldots, \mu_k^{\otimes t}$.

It will be convenient to make the following definitions.

**Definition 6.1.** For a mixture $M = w_1 \mathcal{D}(\mu_1) + \cdots + w_k \mathcal{D}(\mu_k)$, we use $\Pi_{1,M}$ to denote the tensor $w_1 \mu_1^{\otimes t} + \cdots + w_k \mu_k^{\otimes t}$. We may drop the subscript $M$ and just write $\Pi_t$ when it is clear from context.

**Definition 6.2.** For a mixture $M = w_1 \mathcal{D}(\mu_1) + \cdots + w_k \mathcal{D}(\mu_k)$, we define $\Pi_{2k,M}$ to be the tensor $w_1 \mu_1^{\otimes t} + \cdots + w_k \mu_k^{\otimes t}$. We may drop the subscript $M$ when it is clear from context.

We define $\mu_{\max} = \max(1, \|\mu_1\|, \ldots, \|\mu_k\|)$. We do not assume that we know $\mu_{\max}$ in advance. However, the reduction in Section 3.3 means that it suffices to consider when $\mu_{\max}$ is polynomially bounded. Also we can assume $d = k$ i.e. the dimension of the underlying space is equal to the number of components. This is because we can use the reduction in Section 3.3 and if $d < k$, then we can simply add independent standard Gaussian entries in the remaining $k - d$ dimensions.

We now describe our algorithm for implicitly estimating the moment tensor. For the remainder of this section, we will only work with a fixed mixture $M$ so we will drop it from all subscripts e.g. in Definitions 6.1 and 6.2. At a high level, we will recursively compute a sequence of projection matrices $\Pi_1 \in \mathbb{R}^{k \times d}, \Pi_2, \ldots, \Pi_t \in \mathbb{R}^{k \times d}$.

Our goal will be to ensure that $\Pi_{1,\ldots,\Pi_t}$ (which is a $k \times d$ matrix) essentially contains the flattenings of $\mu_1^{\otimes t}, \ldots, \mu_k^{\otimes t}$ in its row span.

To see how to do this, assume that we have computed $\Pi_{0,\ldots,\Pi_t}$ so far. By the inductive hypothesis, $\Pi_{1,\ldots,\Pi_t}$ tells us a $k$-dimensional subspace that essentially contains the flattenings of $\mu_1^{\otimes t-1}, \ldots, \mu_k^{\otimes t-1}$. Thus, we trivially have a $dk$ dimensional subspace, given by the rows of $(\Pi_{0,\ldots,\Pi_t})^{\otimes t}$. It remains to reduce from this $dk$ dimensional space back to a $k$-dimensional space. However, we can now write everything out in this $dk$-dimensional space and simply run PCA and take the top-$k$ singular subspace. Our algorithm is described in full below.

### Algorithm 1 Iterative Projection Step

**Input:** Samples $z_1, \ldots, z_n$ from unknown mixture $M = w_1 \mathcal{D}(\mu_1) + \cdots + w_k \mathcal{D}(\mu_k)$

**Output:** Integer $t > 0$

Split samples into $t$ sets $S_1, \ldots, S_t$ of equal size

Let $\Pi_1 = I_d$ (recall $k = d$)

for $s = 2, \ldots, t$ do

Run Estimate Moment Tensor using samples $S_s$ to get approximation $\hat{A}_{2s} \in \mathbb{R}^{d \times k}$

$A_{2s} = (I_d \otimes_k \Pi_{1,\ldots,\Pi_t}) M_{2s} (I_d \otimes_k \Pi_{1,\ldots,\Pi_t})^T$

Let $\Pi_k \in \mathbb{R}^{k \times d}$ have rows forming an orthonormal basis of the top $k$ singular subspace of $A_{2s}$

**Output:** $(\Pi_k, \ldots, \Pi_t)$

### Algorithm 2 Estimate Moment Tensor

**Input:** Samples $z_1, \ldots, z_n$ from unknown mixture $M = w_1 \mathcal{D}(\mu_1) + \cdots + w_k \mathcal{D}(\mu_k)$

**Input:** Integer $s > 0$

**Input:** Matrices $\Pi_{s-1} \in \mathbb{R}^{k \times d}, \ldots, \Pi_2 \in \mathbb{R}^{k \times d}, \Pi_1 \in \mathbb{R}^{k \times d}$

for $i = 1, 2, \ldots, n$ do

Independently draw samples $x_{1i}, \ldots, x_{4k-1}$ from $\mathcal{D}$

Compute the $(kd)^2$-dimensional vector (recall Definition 4.8)

$X_i = ((I_d \otimes_k \Pi_{1,\ldots,\Pi_t}) \otimes_k (I_d \otimes_k \Pi_{1,\ldots,\Pi_t}))$

flatten ($R_{2s}(x_i, x_{1i}, \ldots, x_{4k-1})$).

Let $K_i$ be the rearrangement of $X_i$ into a square $dk \times dk$-dimensional matrix

**Output:** $A = (K_1 + \cdots + K_n)/n$
The main algorithm, Algorithm 1, computes the projection matrices $\Pi_1, \ldots, \Pi_s$ following the outline above. As a subroutine, it needs to estimate the matrix $A_{2s}$. This is done in Algorithm 2 which relies on the results in Section 4, namely Corollary 4.10 and Corollary 4.11.

### 6.1 Efficient Implementation

A naive implementation of Algorithm 1 requires $d^t$ time, which is too large. However, in this section, we show that we can implement all of the steps more efficiently using only $\text{poly}(ndk, t^4)$ time.

**Remark.** We will later show that it suffices to consider

$$t \sim O\left(\frac{\log(k/w_{\text{min}})}{\log \log(k/w_{\text{min}})}\right)$$

so this runtime is actually polynomial in all parameters that we need.

**Claim 6.3.** Algorithm 1 can be implemented to run in $\text{poly}(n, d, k, t^4)$ time.

**Proof.** Follows easily from Claim 5.3.

### 6.2 Accuracy Analysis

Now, we analyze the correctness of Algorithm 1 namely that the span of the rows of the matrix $\Pi_{k_1, \ldots, k_t}$ essentially contain all of $\mu_1^{(s)}, \ldots, \mu_k^{(s)}$. To simplify notation, we make the following definition.

**Definition 6.4.** For all $s$, we define the matrix

$$A_{2s} = (I_d \otimes_k \Pi_{k_1, \ldots, k_t}^1) M_{2s} (I_d \otimes_k \Pi_{k_1, \ldots, k_t})^T.$$

**Remark.** Note that in the execution of Algorithm 1, $\tilde{A}_{2s}$ is intended to be an estimate of $A_{2s}$.

The main result that we will prove is stated below. Note that this lemma does not require any assumptions about minimum mixing weights or means in the mixture. Instead, it simply says that the subspace spanned by the rows of $\Pi_{k_1, \ldots, k_t}$ essentially contains $\text{flattened}(\mu_i)$ for all components $D(\mu_i)$ with mean and mixing weight bounded away from 0.

**Lemma 6.5.** Let $D$ be a distribution on $\mathbb{R}^d$ that is 1-Poincare and let $M = W_1 D(\mu_1) + \cdots + W_k D(\mu_k)$ be a mixture of translations of $D$. Let $w^*, \epsilon > 0$ be parameters. Assume that the number of samples satisfies

$$n \geq \left(\frac{t^d \mu_{\text{max}}^2 k d}{w^* \epsilon}\right)^C$$

for some sufficiently large universal constant $C$. Then with probability at least $1 - n^{-0.2}$ in the execution of Algorithm 1, the following condition holds: for all $i \in [k]$ such that $\|\mu_i\| \geq 1$ and $w_i \geq w^*$, we have

$$\|\Pi_{k_1, \ldots, k_t} \text{flattened}(\mu_i^{(s)})\| \geq (1 - \epsilon) \|\mu_i\|^s$$

for all $s = 1, 2, \ldots, t$.

**Remark.** The parameter $\epsilon$ represents the desired accuracy and the parameter $w^*$ is a weight cutoff threshold where we guarantee to recover “significant” components whose mixing weight is at least $w^*$.

Roughly, the proof of Lemma 6.5 will involve following the outline at the beginning of this section but quantitatively tracking the errors more precisely. Before we prove Lemma 6.5, we first prove a preliminary claim that our estimation error $\|A_{2s} - \tilde{A}_{2s}\|_F$ is small.

**Claim 6.6.** Assume that for a fixed integer $s$, Algorithm 2 is run with a number of samples

$$n \geq \left(\frac{s^d \mu_{\text{max}}^2 k d}{w^* \epsilon}\right)^C$$

for some sufficiently large universal constant $C$. Then with probability at least $1 - n^{-0.4}$, its output $\tilde{A}_{2s}$ satisfies

$$\|A_{2s} - \tilde{A}_{2s}\|_F \leq 0.5 w^* \epsilon^2.$$
On the other hand, note that $A_{2\delta}$ has rank at most $k$. Assuming that
the hypothesis of Claim 6.6 holds, the $k + 1$st singular value of $A_{2\delta}$
has size at most $w^k \epsilon^2$. Thus,
$$u_i^T A_{2\delta} u_i \leq 0.5w^k \epsilon^2 \|u_i\|^2.$$ 
Finally, using the hypothesis of Claim 6.6 again, we must have
$$u_i^T (A_{2\delta} - A_{2\epsilon}) u_i \leq 0.5w^k \epsilon^2 \|u_i\|^2.$$ 
Putting the previous three inequalities together, we deduce that we must have
$$w_i \|u_i\|^2 \leq w^k \epsilon^2 \|u_i\|^2$$
which implies \( \|u_i\| \leq \epsilon \). Also, the induction hypothesis implies that
$$\|\Pi_{i-1} \ldots \Pi_{i} \text{flatten}(\mu_\delta^{i-1})\| \geq (1 - (s - 1)\epsilon) \|\mu_i\|^{s-1}$$
and thus
$$\|u_i\| \geq (1 - (s - 1)\epsilon) \|\mu_i\|^s.$$ 
Finally, note that
$$\|\Pi_{i-1} \ldots \Pi_{i} \text{flatten}(\mu_i^*)\| = \|u_i - u_i\| \geq (1 - (s - 1)\epsilon) \|\mu_i\|^s - \epsilon \geq (1 - s\epsilon) \|\mu_i\|^s.$$ 
This completes the inductive step. Finally, it remains to note that the overall failure probability can be bounded by union bounding over all applications of Claim 6.6 and is clearly at most $n^{-0.2}$ as long as we choose $n$ sufficiently large. This completes the proof. \( \blacksquare \)

7 Testing Samples Using Implicit Moments

Now we show how to use the projection maps $\Pi_1, \ldots, \Pi_2$ computed by Algorithm 1 to test whether a sample came from a component with mean close to 0 or mean far away from 0. Roughly, given a sample $z$, the test simply works by computing $R_i(z, z_1, \ldots, z_{2t-1})$ for $z_1, \ldots, z_{2t-1} \sim D$ and computing
$$\|\Pi_{i-1} \ldots \Pi_{i} \text{flatten}(R_i(z, z_1, \ldots, z_{2t-1}))\|.$$ 
We output $\text{FA}$ if the above is larger than some threshold and otherwise we output $\text{CLOSE}$. For technical reasons, we will actually average over multiple independent draws for $z_1, \ldots, z_{2t-1} \sim D$.

Roughly, the intuition for why this test works is as follows. Note that if $z \sim D$ then by Corollary 4.10,
$$E[\Pi_{i-1} \ldots \Pi_{i} \text{flatten}(R_i(z, z_1, \ldots, z_{2t-1}))] = 0$$
and if we control the variance using Corollary 4.11, then we can upper bound the length with reasonable probability. On the other hand if $z \sim D(\mu)$ for some $\mu$ with large norm, then
$$E[\Pi_{i-1} \ldots \Pi_{i} \text{flatten}(R_i(z, z_1, \ldots, z_{2t-1}))] = \Pi_{i-1} \ldots \Pi_{i} \text{flatten}(\mu_i^*)$$
and since the algorithm in the previous section can ensure that $\mu_i^*$ is essentially contained in the row span of $\Pi_{i-1} \ldots \Pi_{i}$, the RHS above has large norm. The details of our algorithm for testing samples are described below.

Algorithm 3 Test Samples

| Input: Projection matrices $\Pi_1, \ldots, \Pi_2 \in \mathbb{R}^{k \times d}$, $\Pi_1 \in \mathbb{R}^{k \times d}$ |
| Input: Sample $z \in \mathbb{R}^d$ to test |
| Input: Threshold $\tau$, desired accuracy $\delta$ |
| for $i = 1, 2, \ldots, n$ do |
| Draw samples $z_1, \ldots, z_{2t-1} \sim D$ |
| Let $A_i = \Pi_{i-1} \ldots \Pi_{i} \text{flatten}(R_i(z, z_1, \ldots, z_{2t-1}))$ |
| if $\|A_i\| \geq \tau$ then |
| Output: $\text{FA}$ |
| else |
| Output: $\text{CLOSE}$ |

7.1 Analysis of Test Samples

Now we analyze the behavior of Algorithm 3. The key properties that the test satisfies are summarized in the following two lemmas. Lemma 7.1 says that with $1 - \delta$ probability the test will successfully output $\text{FA}$ for samples from a component with mean far from 0 and Lemma 7.1 says that with $1 - \delta$ probability, the test will successfully output $\text{CLOSE}$ for samples from a component with mean 0.

Note that Lemma 7.1 requires that the row span of $\Pi_{i-1} \ldots \Pi_{i}$ essentially contains $\text{flatten}(\mu_i^*)$ (which can be guaranteed by Algorithm 1 and Lemma 6.5). Lemma 7.1 actually does not require anything about $\Pi_1, \ldots, \Pi_2$ (other than the fact that they are actually projections).

Lemma 7.1. Let $D$ be a distribution that is 1-Poincaré. Let $t \in \mathbb{N}$ and $0 < \delta < 0.01$ be some parameters. Let $\Pi_1, \ldots, \Pi_2 \in \mathbb{R}^{k \times d}$, $\Pi_1 \in \mathbb{R}^{k \times d}$ be any matrices whose rows are orthonormal. Let $\tau$ be some parameter satisfying $\tau \geq (20t)^k k / \delta$. Let $z \sim D$. Then with probability at least $1 - \delta$, Algorithm 3 run with these parameters outputs $\text{CLOSE}$ where the randomness is over $z$ and the random choices within Algorithm 3.

Lemma 7.2. Let $D$ be a distribution that is 1-Poincaré. Let $t \in \mathbb{N}$ and $0 < \delta < 0.01$ be some parameters. Let $z \sim D(\mu_i)$ where $\|\mu_i\| \geq 10^t (\log(1 / \delta) + t)$. Let $\tau$ be some parameter satisfying $\tau \leq (0.4 \|\mu_i\|)^t$. Assume that the matrices $\Pi_1, \ldots, \Pi_2 \in \mathbb{R}^{k \times d}$, $\Pi_1 \in \mathbb{R}^{k \times d}$ satisfy that
$$\|\Pi_{i-1} \ldots \Pi_{i} \text{flatten}(\mu_i^*)\| \geq (1 - t\epsilon) \|\mu_i\|^t,$$
where
$$\epsilon < \frac{\delta}{(10t \|\mu_i\|)^{2t}}.$$ 
Then with probability at least $1 - \delta$, Algorithm 3 run with these parameters outputs $\text{FA}$ (where the randomness is over $z$ and the random choices within Algorithm 3).

Remark. Note that we will want to run the test with some inverse polynomial failure probability i.e. $\delta = \text{poly}(k / w^k)$ for some weight $w^k$. In order to be able to combine Lemma 7.1 and Lemma 7.2 meaningfully, we need
$$(0.4 \|\mu_i\|)^t \geq (20t)^k k / \delta.$$ 
If $\|\mu_i\| \geq (\log(k / w^k))^{1 + c}$ for some constant $c > 0$ then setting $t \sim O(c^2 \log(k / w^k) / \log(\log(k / w^k)))$ ensures that the above inequality is true. Note that for this setting, $t^t = \text{poly}(k / w^k)$ (where we treat
2.3 and Algorithm 3.7 will follow easily from it.

We are now ready to prove Theorem 2.3, our full result for mixtures of Poincare distributions. Let \( D' = (D - D)/\sqrt{2} \) i.e. \( D' \) is the distribution of the difference between two independent samples from \( D \) scaled down by \( \sqrt{2} \). The \( \sqrt{2} \) scaling ensures that \( D' \) is 1-Poincare by Fact 3.7. Note that we can take pairwise differences between samples to simulate access to the mixture \[
M' = (w_1^2 + \cdots + w_k^2)D' + \sum_{i \neq j} w_i w_j D'((\mu_i - \mu_j)/\sqrt{2}).
\]

We will run Algorithm 1 and Algorithm 3 for the distribution \( D' \) and mixture \( M' \). Note that the test in Algorithm 3 will test for a pair of samples say \( z, z' \), from components \( D(\mu_i), D(\mu_j) \) of the original mixture, whether \( |\mu_i - \mu_j| \) is large or zero. Running this test between all pairs of samples will let us form clusters of samples that correspond to each of the components. We begin with a lemma that more precisely specifies the guarantees of this test and then Theorem 2.3 will follow easily from it.

**Lemma 8.1.** Let \( D \) be a 1-Poincare distribution on \( \mathbb{R}^d \). Let \[
M = w_1 D(\mu_1) + \cdots + w_k D(\mu_k)
\]
be a mixture of translated copies of \( D \). Let \( w^*, s, \delta \) be parameters and assume that \( s \geq (\log k/(w^* \delta))^4c/3 \) for some \( 0 < c < 1 \). Also assume that \[
\max_i \|w_i - \mu_i\| \leq \min_{\alpha}(k/(w^* \delta))^2, s^C
\]
for some \( C \). There is an algorithm that takes \( n = \text{poly}(kd/(w^* \delta)^C) \) samples from \( M \) and \( D \) and runs in \( \text{poly}(n) \) time and achieves the following testing guarantees: for a pair of (independent) samples \( z \sim D(\mu_i), z' \sim D(\mu_j) \):

- If \( i = j \) and \( w_i \geq w^* \) then with probability \( 1 - \delta \) the output is accept
- If \( w_i, w_j \geq w^* \) and \( |\mu_i - \mu_j| \geq s \) then with probability \( 1 - \delta \) the output is reject

where the randomness is over the samples \( z, z' \) and the random choices in the algorithm.

Using Lemma 8.1, it is not difficult to prove Theorem 2.3.

**Proof of Theorem 2.3.** Recall by the reduction in Section 3.3 that we may assume \( d \leq k \) and that \( |\mu_i - \mu_j| \leq O(k/w_{\min}^2) \) for all \( i, j \). Now we will do the following process to estimate the means of the components.

1. Draw a sample \( z \sim M \)
2. Take \( m = (k/(w_{\min} \alpha))^{10^9} \) samples \( z_1, \ldots, z_m \sim M \)
3. Use Lemma 8.1 with parameters \( w^* = w_{\min}, \delta = (w_{\min} \alpha/k)^{10^9} \)

To test the pair of samples \( z, z_j \) for all \( i \in [m] \)
4. Let \( S \subset [m] \) be the set of all \( i \) that are ACCEPTED and compute \( \mu_i = \frac{1}{|S_i|} \sum_{i \in S} z_i \)

We first argue that if \( z \) is a sample from \( D(\mu_i) \) for some \( i \), then with \( 1 - (w_{\min} \alpha/k)^{10^9} \) probability, the procedure returns \( \mu_i \) such that \( |\mu_i - \mu_i| \leq 0.1\alpha \). This is because by the guarantees of Lemma 8.1, with probability at least \( 1 - (w_{\min} \alpha/k)^{10^9} \) the test will accept all samples from among \( z_1, \ldots, z_m \) that are from the component \( D(\mu_i) \) and reject all of the others. Also, with high probability, there will be at least \( (k/(w_{\min} \alpha))^{99} \) samples from the component \( D(\mu_i) \) so by Claim 3.8 and union bounding all of the failure probabilities, we have that if \( z \) is a sample from \( D(\mu_i) \) then \( |\mu_i - \mu_i| \leq 0.1\alpha \) with probability at least \( 1 - (w_{\min} \alpha/k)^{10^9} \).

Now it suffices to repeat the procedure in steps 1 - 4 for \( l = (k/(w_{\min} \alpha))^{10^9} \) independent samples \( z \sim M \). This gives us a list of means say \( S = \{\tilde{\mu}_1, \ldots, \tilde{\mu}_k\} \). The previous argument implies that most of these estimates will be close to one of the true means and that all of the true means will be represented. To ensure that with high probability we output exactly one estimate corresponding to each true mean and no extraneous estimates, we do a sort of majority voting.

We will inspect the estimates in \( S \) one at a time and decide whether to output them or not. Let \( T \) be the set of estimates that we will output. Note that \( T \) is initially empty. Now for each \( i \), let \( S_i \) be the subset of \( \{\tilde{\mu}_1, \ldots, \tilde{\mu}_k\} \) consisting of all means with \( |\tilde{\mu}_j - \tilde{\mu}_i| \leq 0.2\alpha \). If \( |S_i| \geq 0.9w_{\min}l \) and \( \tilde{\mu}_i \) is not within \( \alpha \) of any element of \( T \) then add \( \tilde{\mu}_i \) to \( T \). Otherwise do nothing.

We claim that with high probability, this procedure returns one estimate corresponding to each true mean and nothing extraneous. First, with high probability there will be no extraneous outputs because if say \( \tilde{\mu}_i \) is at least \( 0.5\alpha \) away from all of the true means, then with high probability we will have \( |S_i| < 0.9w_{\min}l \). Now it is also clear that with high probability we output exactly one estimate corresponding to each true mean (since \( l \) is sufficiently large that with high probability we get enough samples from each component). Thus, with high probability, the final output will be a set of means that are within \( \alpha \) of the true means up to some permutation. Once we have learned the means, we can learn the mixing weights by simply taking fresh samples from \( M \) and clustering since with high probability, we can uniquely identify which component a sample came from. This completes the proof.

The clustering guarantee in Corollary 2.4 follows as an immediate consequence of Theorem 2.3.

**Proof of Corollary 2.4.** Let the estimated means computed by Theorem 2.3 for \( \alpha = (w_{\min}/k)^{10} \) be \( \tilde{\mu}_1, \ldots, \tilde{\mu}_k \). Now for all \( j_1, j_2 \in [k] \) with \( j_1 \neq j_2 \), let

\[
\delta_{j_1, j_2} = \frac{|\tilde{\mu}_j - \tilde{\mu}_n|}{|\tilde{\mu}_j - \tilde{\mu}_n|}. \]

Now given a sample \( z \) from \( M \), we compute the index \( j \) such that for all \( j_1, j_2 \), we have

\[
|\delta_{j_1, j_2} \cdot (\tilde{\mu}_j - z)| \leq (\log(k/w_{\min}))^{1.15}.0.5c.\]

Note that by the guarantees of Theorem 2.3, there is a permutation \( \pi \) such that \( |\tilde{\mu}_{\pi(i)} - \mu_i| \leq \alpha \) for all \( i \). If \( z \) is a sample from \( D(\mu_i) \), then by the tail bound in Fact 3.7, with high probability the unique index \( j \) that satisfies the above is exactly \( j = \pi(i) \) and thus, we recover the ground truth clustering with high probability.
9 LEARNING MIXTURES OF GAUSSIANS

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