Covariance Matrix Adaptation MAP-Annealing

Matthew C. Fontaine
University of Southern California
Los Angeles, CA, USA
mfontain@usc.edu

Stefanos Nikolaidis
University of Southern California
Los Angeles, CA, USA
nikolaid@usc.edu

ABSTRACT

Single-objective optimization algorithms search for the single highest-quality solution with respect to an objective. Quality diversity (QD) optimization algorithms, such as Covariance Matrix Adaptation MAP-Elites (CMA-ME), search for a collection of solutions that are both high-quality with respect to an objective and diverse with respect to specified measure functions. However, CMA-ME suffers from three major limitations highlighted by the QD community: prematurely abandoning the objective in favor of exploration, struggling to explore flat objectives, and having poor performance for low-resolution archives. We propose a new quality diversity algorithm, Covariance Matrix Adaptation MAP-Annealing (CMA-MAE), that addresses all three limitations. We provide theoretical justifications for the new algorithm with respect to each limitation. Our theory informs our experiments, which support the theory and show that CMA-MAE achieves state-of-the-art performance and robustness.

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1 INTRODUCTION

Consider an example problem of searching for celebrity faces in the latent space of a generative model. As a single-objective optimization problem, we specify an objective \( f \) that targets a celebrity such as Tom Cruise. A single-objective optimizer, such as CMA-ES [35], will converge to a single solution of high objective value, an image that looks like Tom Cruise as much as possible.

However, this objective has ambiguity. How old was Tom Cruise in the photo? Did we want the person in the image to have short or long hair? By instead framing the problem as a quality diversity optimization problem, we additionally specify a measure function \( m_1 \) that quantifies age and a measure function \( m_2 \) that quantifies hair length. A quality diversity algorithm [8, 66], such as CMA-ME [26], can then optimize for a collection of images that are diverse with respect to age and hair length, but all look like Tom Cruise.

While prior work [17, 20, 24, 26] has shown that CMA-ME solves such QD problems efficiently, three important limitations of the algorithm have been discovered. First, on difficult to optimize objectives, variants of CMA-ME will abandon the objective too soon [76], and instead favor exploring the measure space, the vector space defined by the measure function outputs. Second, the CMA-ME algorithm struggles to explore flat objective functions [62]. Third, CMA-ME works well on high-resolution archives, but struggles to explore low-resolution archives [14, 21].

We propose a new algorithm, CMA-MAE, that addresses these three limitations. To address the first limitation, we derive an algorithm that smoothly blends between CMA-ES and CMA-ME. First, consider how CMA-ES and CMA-ME differ. At each step CMA-ES’s objective ranking maximizes the objective function \( f \) by approximating the natural gradient of \( f \) at the current solution point [1]. In contrast, CMA-ME’s improvement ranking moves in the direction of the natural gradient of \( f - f_A \) at the current solution point, where \( f_A \) is a discount function equal to the objective of the best solution so far that has the same measure values as the current solution point. The function \( f - f_A \) quantifies the gap between a candidate solution and the best solution so far at the candidate solution’s position in measure space.

Our key insight is to anneal the function \( f_A \) by a learning rate \( \alpha \). We observe that when \( \alpha = 0 \), then our discount function \( f_A \) never increases and our algorithm behaves like CMA-ES. However, when \( \alpha = 1 \), then our discount function always maintains the best solution for each region in measure space and our algorithm behaves like CMA-ME. For \( 0 < \alpha < 1 \), CMA-MAE smoothly blends between the two algorithms’ behavior, allowing for an algorithm that spends more time on the optimization of \( f \) before transitioning to exploration. Figure 1 is an illustrative example of varying the learning rate \( \alpha \).

Our proposed annealing method naturally addresses the flat objective limitation. Observe that both CMA-ES and CMA-ME struggle on flat objectives \( f \) as the natural gradient becomes 0 in this case and each algorithm will restart. However, we show that, when CMA-MAE optimizes \( f - f_A \) for \( 0 < \alpha < 1 \), the algorithm becomes a descent method on the density histogram defined by the archive.

Finally, CMA-ME’s poor performance on low resolution archives is likely caused by the non-stationary objective \( f - f_A \) changing too quickly for the adaptation mechanism to keep up. Our archive learning rate \( \alpha \) controls how quickly \( f - f_A \) changes. We derive a conversion formula for \( \alpha \) that allows us to derive equivalent \( \alpha \) for different archive resolutions. Our conversion formula guarantees that CMA-MAE is the first QD algorithm invariant to archive resolution.

While our new annealing method benefits CMA-ME, our approach is also compatible with CMA-ME’s differentiable quality diversity (DQD) counterpart, CMA-MEGA [21]. We apply the same modification in the DQD setting to form CMA-MAEGA. To evaluate CMA-MAEGA, we improve the latent space illumination (LSI) [21] domain by introducing a higher-dimensional domain based on StyleGAN2, capable of producing higher quality images. This new domain highlights the advantages of DQD in high-dimensional spaces and demonstrates the performance benefits of our annealing method.

We note that archive resolution affects the performance of all current QD algorithms.
We present several QD algorithms that solve derivative-free QD objective becomes to find a solution problems to provide context for our proposed CMA-MAE algorithm. Appendix D contains information about the DQD algorithm. Overall, our work shows how a simple algorithmic change to CMA-MAE addresses all three major limitations affecting CMA-ME’s performance and robustness. Our theoretical findings justify the aforementioned properties and inform our experiments, which show that CMA-MAE outperforms state-of-the-art QD algorithms and maintains robust performance across different archive resolutions.

2 PROBLEM DEFINITION

Quality Diversity. We adopt the quality diversity (QD) problem definition from prior work [21]. A QD problem consists of an objective function \( f : \mathbb{R}^n \rightarrow \mathbb{R} \) that maps \( n \)-dimensional solution parameters to the scalar value representing the quality of the solution and \( k \) measure functions \( m_i : \mathbb{R}^n \rightarrow \mathbb{R} \) or, as a vector function, \( m : \mathbb{R}^n \rightarrow \mathbb{R}^k \) that quantify the behavior or attributes of each solution.\(^2\) The range of \( m \) forms a measure space \( S = m(\mathbb{R}^n) \). The QD objective is to find a set of solutions \( \theta \in \mathbb{R}^n \), such that \( m(\theta) = s \) for each \( s \) in \( S \) and \( f(\theta) \) is maximized.

The measure space \( S \) is continuous, but solving algorithms need to produce a finite collection of solutions. Therefore, QD algorithms in the MAP-Elites [15, 59] family relax the QD objective by discretizing the space \( S \). Given \( T \) as the tessellation of \( S \) into \( M \) cells, the QD objective becomes to find a solution \( \theta_i \) for each of the \( i \in \{1, \ldots, M\} \) cells, such that each \( \theta_i \) maps to the cell corresponding to \( m(\theta_i) \) in the tessellation \( T \). The QD objective then becomes maximizing the objective value \( f(\theta_i) \) of all cells:

\[
\max \sum_{i=1}^{M} f(\theta_i) \tag{1}
\]

The differentiable quality diversity (DQD) problem [21] is a special case of the QD problem where both the objective \( f \) and measures \( m_i \) are first-order differentiable.

3 PRELIMINARIES

We present several QD algorithms that solve derivative-free QD problems to provide context for our proposed CMA-MEGA algorithm. Appendix D contains information about the DQD algorithm CMA-MEGA, which solves problems where the gradients of the objective and measure functions are available.

MAP-Elites and MAP-Elites (line). The MAP-Elites QD algorithm produces an archive of solutions, where each cell in the archive corresponds to the provided tessellation \( T \) in the QD problem definition. The algorithm initializes the archive by sampling solutions from the solution space \( \mathbb{R}^n \) from a fixed distribution. After initialization, MAP-Elites produces new solutions by selecting occupied cells uniformly at random and perturbing them with isotropic Gaussian noise: \( \theta' = \theta_i + \sigma \mathcal{N}(0, I) \). For each new candidate solution \( \theta' \), the algorithm computes an objective \( f(\theta') \) and measures \( m(\theta') \). MAP-Elites places \( \theta' \) into the archive if the cell corresponding to \( m(\theta') \) is empty or \( \theta' \) obtains a better objective value \( f(\theta') \) than the current occupant. The MAP-Elites algorithm results in an archive of solutions, solutions that are diverse with respect to the measure function \( m \), but also high quality with respect to the objective \( f \). Prior work [79] proposed the MAP-Elites (line) algorithm by augmenting the isotropic Gaussian perturbation with a linear interpolation between two solutions \( \theta_i \) and \( \theta_j \): \( \theta' = \theta_i + \sigma_1 \mathcal{N}(0, I) + \sigma_2 \mathcal{N}(0, 1)(\theta_i - \theta_j) \).

CMA-ME. Covariance Matrix Adaptation MAP-Elites [26] combines the adapting mechanisms of MAP-Elites with the adaptation mechanisms of CMA-ES [35]. Instead of perturbing archive solutions with Gaussian noise, CMA-ME maintains a multivariate Gaussian of search directions \( \mathcal{N}(0, \Sigma) \) and a search point \( \theta \in \mathbb{R}^n \). The algorithm updates the archive by sampling \( \lambda \) solutions around the current search point \( \theta_i \sim \mathcal{N}(\theta, \Sigma) \). After updating the archive, CMA-ME ranks solutions via a two stage ranking. Solutions that discover a new cell are ranked by the objective \( \Delta_i = f(\theta_i) \), and solutions that map to an occupied cell \( \epsilon \) are ranked by the improvement over the incumbent solution \( \theta_\epsilon \) in that cell: \( \Delta_i = f(\theta_i) - f(\theta_\epsilon) \). CMA-ME prioritizes exploration by ranking all solutions that discover a new cell before all solutions that improve upon an existing cell. Finally, CMA-ME moves \( \theta \) towards the largest improvement in the archive, according to the CMA-ES update rules. Prior work [21] showed that the improvement ranking of CMA-ME approximates a natural gradient of a modified QD objective (see Eq. 1).

4 PROPOSED ALGORITHMS

We present the CMA-MAE algorithm. While we focus on CMA-ME, the same augmentations apply to CMA-MEGA to form the novel CMA-MAEGA algorithm (see Appendix D). CMA-MAE, CMA-MAE is an algorithm that adjusts the rate the non-stationary QD objective \( f - f_A \) changes. First, consider a
high level how CMA-ME explores the measure space and discovers high quality solutions. The CMA-ME algorithm maintains a solution point \( \theta \) and an archive \( A \) with previously discovered solutions. When CMA-ME samples a new solution \( \theta' \), the algorithm computes the solution’s objective value \( f(\theta') \) and maps the solution to a cell \( e \) in the archive based on the measure \( m(\theta') \). CMA-ME then computes the improvement of the objective value \( f(\theta') \) of the new solution, over a discount function \( f_A : \mathbb{R}^n \rightarrow \mathbb{R} \). In CMA-ME, we define \( f_A(\theta') \) by computing the cell \( e \) in the archive corresponding to \( m(\theta') \) and letting \( f_A(\theta') = f(\theta_e) \), where \( \theta_e \) is the incumbent solution of cell \( e \). The algorithm ranks candidate solutions by improvement \( \Delta = f(\theta') - f_A(\theta') \) and moves the search in the direction of higher ranked solutions.

Assume that CMA-ME samples a new solution \( \theta' \) with a high objective value of \( f(\theta') = 99 \). If the current occupant \( \theta_e \) of the corresponding cell has a low objective value of \( f(\theta_e) = 0.3 \), then the improvement in the archive \( \Delta = f(\theta') - f(\theta_e) = 98.7 \) is high and the algorithm will move the search point \( \theta \) towards \( \theta' \). Now, assume that in the next iteration the algorithm discovers a new solution \( \theta'' \) with objective value \( f(\theta'') = 100 \) that maps to the same cell as \( \theta' \). The improvement then is \( \Delta = f(\theta'') - f(\theta') = 1 \) as \( \theta'' \) replaced \( \theta_e \) in the archive in the previous iteration. CMA-ME would likely move \( \theta \) away from \( \theta'' \) as the solution resulted in low improvement. In contrast, CMA-ES would move towards \( \theta'' \) as it ranks only by the objective \( f \), ignoring previously discovered solutions with similar measure values.

In the above example, CMA-ME moves away from a high performing region in order to maximize how the archive changes. However, in domains with hard-to-optimize objective functions, it is beneficial to perform more optimization steps towards the objective \( f \) before leaving each high-performing region [76].

Like CMA-ME, CMA-MAE maintains a discount function \( f_A(\theta') \) and ranks solutions by improvement \( f(\theta') - f_A(\theta') \). However, instead of maintaining an elitist archive by setting \( f_A(\theta') \) equal to \( f(\theta_e) \), we maintain a soft archive by setting \( f_A(\theta') \) equal to \( t_e \), where \( t_e \) is an acceptance threshold maintained for each cell in the archive \( A \). When adding a candidate solution to the archive, we control the rate that \( t_e \) changes by the archive learning rate \( \alpha \) as follows:

\[
t_e \leftarrow (1 - \alpha) t_e + \alpha f(\theta').
\]

The archive learning rate \( \alpha \) in CMA-MAE allows us to control how quickly we leave a high-performing region of measure space. For example, consider discovering solutions in the same cell with objective value 100 in 5 consecutive iterations. The improvement values computed by CMA-ME against the elitist archive would be 100, 0, 0, 0, 0, thus CMA-ME would move rapidly away from this cell. The improvement values computed against the soft archive of CMA-MAE with \( \alpha = 0.5 \) would diminish smoothly as follows: 100, 50, 25, 12.5, 6.25, enabling further exploitation of the high-performing region.

Next, we walk through the CMA-MAE algorithm step-by-step. Algorithm 1 shows the pseudo-code for CMA-MAE with the differences from CMA-ME highlighted in yellow. First, on line 3 we initialize the acceptance threshold to \( \min f \). In each iteration we sample \( \lambda \) solutions around the current search point \( \theta \) (line 6). For each candidate solution \( \theta_i \), we evaluate the solution and compute the objective value \( f(\theta_i) \) and measure values \( m(\theta_i) \) (line 7). Next, we compute the cell \( e \) in the archive that corresponds to the measure values and the improvement \( \Delta \) over the current threshold \( t_e \) (lines 8-9). If the objective crosses the acceptance threshold \( t_e \), we replace the incumbent \( \theta_e \) in the archive and increase the acceptance threshold \( t_e \) (lines 10-12). Next, we rank all candidate solutions \( \theta_i \) by their improvement \( \Delta \). Finally, we step our search point \( \theta \) and adapt our covariance matrix \( \Sigma \) towards the direction of largest improvement (lines 15-16) according to CMA-ES’s update rules [35].

**CMA-MAEGA.** We note that our augmentations to the CMA-MAE algorithm only affects how we replace solutions in the archive and how we calculate \( \Delta \), CMA-ME and CMA-MAEGA replace solutions and calculate \( \Delta \) identically, so we apply the same augmentations to CMA-MAEGA to form a new DQD algorithm, CMA-MAEGA, in Appendix D.

## 5 THEORETICAL PROPERTIES OF CMA-MAE

We provide insights about the behavior of CMA-MAE for different \( \alpha \) values. We include all proofs in Appendix E. CMA-MAEGA has similar theoretical properties discussed in Appendix F.

**Theorem 5.1.** The CMA-ES algorithm is equivalent to CMA-MAE when \( \alpha = 0 \), if CMA-ES restarts from an archive solution.
Finally, we wish to address the limitation that CMA-ME performs poorly on flat objectives, where all solutions have the same objective value. Consider how CMA-ME behaves on a flat objective $f(\theta) = C$ for all $\theta \in \mathbb{R}^n$, where $C$ is an arbitrary constant. CMA-ME will only discover each new cell once and will not receive any further feedback from that cell, since any future solution cannot replace the incumbent elite. This hinders the CMA-ME’s movement in measure space, which is based on feedback from changes in the archive. Future candidate solutions will only fall into occupied cells, triggering repeated restarts caused by CMA-ES’s restart rule.

When the objective function plateaus, we still want CMA-ME to perform well and benefit from the CMA-ES adaptation mechanisms. One reasonable approach would be to keep track of the frequency $\omega_t$ that each cell $e$ has been visited in the archive, where $\omega_t$ represents the number of times a solution was generated in that cell. Then, when a flat objective occurs, we rank solutions by descending frequency counts. Conceptually, CMA-ME would descend the density histogram defined by the archive, pushing the search towards regions of the measure space that have been sampled less frequently. Theorem 5.4 shows that we obtain the density descent behavior on flat objectives without additional changes to the CMA-ME algorithm.

**Theorem 5.4.** The CMA-ME algorithm optimizing a constant objective function $f(\theta) = C$ for all $\theta \in \mathbb{R}^n$ is equivalent to density descent, when $0 < \alpha < 1$ and $\min f < C$.

We highlight that the proof of Theorem 5.4 is based on two critical properties. First, the threshold update rule forms a strictly increasing sequence of thresholds for each cell. Second, CMA-ES is invariant to order preserving transformations of its objective $f$. While we have proposed the update rule of line 12 of Algorithm 1, we note that any update rule that satisfies the increasing sequence property retains the density descent property and is thus applicable in CMA-ME.

While Theorem 5.4 assumes a constant objective $f$, we conjecture that the theorem holds true generally when threshold $t_e$ in each cell $e$ approaches the local optimum within the cell boundaries (see Conjecture E.7 in the Appendix).

## 6 EXPERIMENTS

We compare the performance of CMA-ME with the state-of-the-art QD algorithms MAP-Elites, MAP-Elites (line), and CMA-ME, using existing Pyribs [77] QD library implementations. We set $\alpha = 0.01$ for CMA-ME and include additional experiments for varying $\alpha$ in section 7. Because annealing methods replace solutions based on the threshold, we retain the best solution in each cell for comparison purposes. We include additional comparisons between CMA-MEGA and CMA-MAEGA – the DQD counterpart to CMA-ME.

We select the benchmark domains from prior work [21]: linear projection [26], arm repertoire [16], and latent space illumination [24]. To evaluate the good exploration properties of CMA-ME on flat objectives, we introduce a variant of the linear projection domain to include a “plateau” objective function that is constant everywhere for solutions within a fixed range and has a quadratic penalty for solutions outside the range. We describe the domains in detail in Appendix B.

We additionally introduce a second LSI experiment on StyleGAN2 [47], configured by insights from the generative art community [12, 27] that improve the quality of single-objective latent
space optimization. To improve control over image synthesis, the LSI (StyleGAN) domain optimizes the full 9216-dimensional latent w-space, rather than a compressed 512-dimensional latent space in the LSI (StyleGAN) experiments. We exclude CMA-ME and CMA-MAE from this domain due to the prohibitive size of the covariance matrix. The LSI (StyleGAN2) domain allows us to evaluate the performance of DQD algorithms on a much more challenging DQD domain than prior work. We describe the domain in Appendix I.

### 6.1 Experiment Design

**Independent Variables.** We follow a between-groups design with two independent variables: the algorithm and the domain.

**Dependent Variables.** We use the sum of f values of all cells in the archive, defined as the QD-score [66], as a metric for the quality and diversity of solutions. Following prior work [21], we normalize the QD-score metric by the archive size (the total number of cells from the tessellation of measure space) to make the metric invariant to archive resolution. We additionally compute the coverage, defined as the number of occupied cells in the archive divided by the total number of cells.

### 6.2 Analysis

**Derivative-free QD Algorithms.** Table 1 shows the QD-score and coverage values for each algorithm and domain, averaged over 20 trials for the linear projection (LP) and arm repertoire domains and over 5 trials for the LSI domain. Table 2 shows the QD-score and coverage values for each algorithm and domain, averaged over 20 trials for the linear projection (LP) and arm repertoire domains and over 5 trials for the LSI domains. We conducted a two-way ANOVA to examine the effect of the algorithm and domain on the QD-score. There was a significant interaction between the search algorithm and the domain (F(5, 168) = 165.7, p < 0.001). Simple main effects analysis with Bonferroni corrections showed that CMA-MAE outperformed CMA-MEGA in the LP (sphere), arm repertoire, and LSI (StyleGAN) domains. There was no statistically significant difference between the two algorithms in the LP (Rastrigin), LP (plateau), and LSI (StyleGAN) domains.

We observe the trends that CMA-MAE initially explores regions of the measure space that have high-objective values. Once the archive becomes saturated, CMA-MAE reduces to approximate density descent, as we prove in Theorem 5.4 for flat objectives. On the other hand, CMA-ME does not receive any exploration signal when the objective landscape becomes flat, resulting in poor performance.

**DQD Algorithms.** We additionally compare CMA-MEGA and CMA-MAE in the five benchmark domains. The LSI (StyleGAN) domain optimizes the full 9216-dimensional latent space, rather than a compressed 512-dimensional latent space in the LSI (StyleGAN) experiments. We exclude CMA-ME and CMA-MAE from this domain due to the prohibitive size of the covariance matrix. The LSI (StyleGAN2) domain allows us to evaluate the performance of DQD algorithms on a much more challenging DQD domain than prior work. We describe the domain in Appendix I.

We attribute the absence of a statistical difference in the QD-score between the two algorithms on the LP (Rastrigin) and LP (plateau) domains on the perfect coverage obtained by both algorithms. Thus, any differences in QD-score are based on the objective values of the solutions returned by each algorithm.

In LP (plateau), the optimal objective for each cell is easily obtainable for both methods. The LP (Rastrigin) domain contains many local optima, because of the form of the objective function (Eq. 5). CMA-MEGA will converge to these optima before restarting, behaving as a single-objective optimizer within each local optimum. Because of the large number of local optima in the domain, CMA-MEGA obtains a higher QD-score.

In the LSI (StyleGAN) domain, we attribute similar performance between CMA-MEGA and CMA-MAE to the restart rules used to keep each search within the training distribution of StyleGAN. The ill-conditioned latent space of StyleGAN also explains why CMA-MAE outperforms both DQD algorithms on this domain. Being a natural gradient optimizer, CMA-MAE is an approximate second-order method, and second-order methods are better suited for optimizing spaces with ill-conditioned curvature.

On the other hand, in the LSI (StyleGAN2) domain, we regularize the search space by an L2 penalty in latent space, allowing for a...
Figure 3: QD-score plot with 95% confidence intervals and heatmaps of generated archives by CMA-MAE and CMA-ME for the linear projection sphere (top), plateau (middle), and arm repertoire (bottom) domains. Each heatmap visualizes an archive of solutions across a 2D measure space.

larger learning rate and a basic restart rule for both algorithms, while still preventing drift out of the training distribution of StyleGAN2. Because of the fewer restarts, CMA-MAEGA can take advantage of the density descent property, which was shown to improve exploration in CMA-MAE, and outperform CMA-MEGA. Fig. 4 shows an example collage generated from the final archive of CMA-MAEGA on the LSI (StyleGAN2) domain. We note that because StyleGAN2 has a better conditioning on the latent space [47], the model is better suited for first-order optimization of its latent space, which helps distinguish between the two DQD algorithms.
We include runs for MAP-Elites and MAP-Elites (line) on the LSI (StyleGAN2) domain in Table 1 for comparison purposes. In the LSI (StyleGAN2) domain, the two algorithms drift out of distribution and suffer a regularization penalty that results in negative objective values. We observe that the gap in performance between derivative-free QD algorithms and DQD algorithms is higher in the LSI (StyleGAN2) domain than in LSI (StyleGAN) domain. This highlights the benefits of leveraging gradients of the objective and measure functions in high-dimensional search spaces.

7 ON THE ROBUSTNESS OF CMA-MAE

Next, we present two studies that evaluate CMA-MAE robustness across two hyperparameters that may affect algorithm performance: the archive learning rate \( \alpha \) and the archive resolution.

**Archive Learning Rate.** We examine the effect of different archive learning rates on the performance of CMA-MAE in the linear projection and arm repertoire domains. We vary the learning rate from 0 to 1 on an exponential scale, while keeping the resolution constant.

Table 3 shows that running CMA-MAE with the different \( 0 < \alpha < 1 \) values results in similar performance, showing that CMA-MAE is fairly robust to the exact choice of \( \alpha \) value. On the other hand, if \( \alpha = 0 \) or \( \alpha = 1 \) the performance drops drastically. Setting \( \alpha = 1 \) results in very similar performance with CMA-ME, which supports our insight from Theorem 5.2.

**Archive Resolution.** As noted in prior work [14, 21], quality diversity algorithms in the MAP-Elites family sometimes perform differently when run with different archive resolutions. For example, in the linear projection domain proposed in prior work [26], CMA-ME outperformed MAP-Elites and MAP-Elites (line) for archives of resolution \( 500 \times 500 \), while in this paper we observe that it performs worse for resolution \( 100 \times 100 \). In this study, we investigate how CMA-MAE performs at different archive resolutions.

First, we note that the optimal archive learning rate \( \alpha \) is dependent on the resolution of the archive. Consider as an example a sequence of solution additions to two archives \( A_1 \) and \( A_2 \) of resolution \( 100 \times 100 \) and \( 200 \times 200 \) respectively. \( A_2 \) subdivides each cell in \( A_1 \) into four cells, thus archive \( A_2 \)'s thresholds \( t_\alpha \) should increase at a four times faster rate than \( A_1 \). To account for this difference, we compute \( \alpha_2 \) for \( A_2 \) via a conversion formula \( \alpha_2 = 1 - (1 - \alpha_1)^r \) (see derivation in Appendix G), where \( r \) is the ratio of cell counts between archives \( A_1 \) and \( A_2 \). We initialize \( \alpha_1 = 0.01 \) for \( A_1 \). In the above example, \( \alpha_2 = 1 - (1 - 0.01)^4 = 0.0394 \).

Fig. 5 shows the QD-score of CMA-MAE with the resolution-dependent archive learning rate and the baselines for each benchmark domain. CMA-ME performs worse as the resolution decreases because the archive changes quickly at small resolutions, affecting CMA-ME’s adaptation mechanism. On the contrary, MAP-Elites and MAP-Elites (line) perform worse as the resolution increases.
due to having more elites to perturb. CMA-MAE’s performance is invariant to the resolution of the archive.

8 RELATED WORK

Quality Diversity Optimization. The predecessor to quality diversity optimization, simply called diversity optimization, originated with the Novelty Search algorithm [54], which searches for a collection of solutions that are diverse in measure space. Later work introduced the Novelty Search with Local Competition (NSLC) [55] and MAP-Elites [15, 59] algorithms, which combined single-objective optimization with diversity optimization and were the first QD algorithms. Since then, several QD algorithms have been proposed, based on a variety of single-objective optimization methods, such as Bayesian optimization [49], evolution strategies [10, 11, 26], differential evolution [9], and gradient ascent [21]. Several works have improved selection mechanisms [16, 71], archives [23, 72, 78], perturbation operators [61, 79], and resolution scaling [13, 23, 32].

QD with Gradient Information. Several works combine gradient information with QD optimization without leveraging the objective and measure gradients directly. For example, in model-based QD optimization [7, 28, 29, 34, 48, 57, 80], prior work [68] trains an autoencoder on the archive of solutions and leverages the Jacobian of the decoder network to compute the covariance of the Gaussian perturbation. Works in quality diversity reinforcement learning (QD-RL) [60, 63, 65, 76] approximate a reward gradient or diversity gradient via a critic network, action space noise, or evolution strategies and incorporate those gradients into a QD-RL algorithm.

Acceptance Thresholds. Our proposed archive learning rate $\alpha$ was loosely inspired by simulated annealing methods [2] that maintain an acceptance threshold that gradually becomes more selective as the algorithm progresses. The notion of an acceptance threshold is also closely related to minimal criterion methods in evolutionary computation [5, 6, 53, 73]. Our work differs by both 1) maintaining an acceptance threshold per archive cell rather than a global threshold and 2) annealing the threshold.

9 LIMITATIONS AND FUTURE WORK

Our approach introduced two hyperparameters, $\alpha$ and $\min_f$ (see Appendix J for $\min_f$ analysis), to control the rate that $f = f_\alpha$ changes. We observed that an $\alpha$ set strictly between 0 and 1 yields theoretical exploration improvements and that CMA-MAE is robust with respect to the exact choice of $\alpha$. We additionally derived a conversion formula that converts an $\alpha_1$ for a specific archive resolution to an equivalent $\alpha_2$ for a different resolution. However, the conversion formula still requires practitioners to specify a good initial value of $\alpha_1$. Future work will explore ways to automatically initialize $\alpha$, similar to how CMA-ES automatically assigns internal parameters [35].

CMA-MAE’s DQD counterpart CMA-MAEGA sets a new state-of-the-art in DQD optimization. However, observing its performance benefits required the more challenging LSI (StyleGAN2) domain. This highlights the need for more challenging DQD problems to advance research in DQD algorithms, since many of the current benchmark domains can be solved optimally by existing algorithms.

Quality diversity optimization is a rapidly growing branch of stochastic optimization with applications in generative design [28, 29, 33], automatic scenario generation in robotics [20, 22, 25], reinforcement learning [60, 63, 65, 76], damage recovery in robotics [15], and procedural content generation [4, 17, 24, 31, 50, 69, 70, 75, 80]. Our paper introduces a new quality diversity algorithm, CMA-MAE. Our theoretical findings inform our experiments, which show that CMA-MAE addresses three major limitations affecting CMA-ME, leading to state-of-the-art performance and robustness.

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