Accelerating the Discovery of Rare Materials with Bounded Optimization Techniques

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Abstract

Discovering a rare material within a vast search space exhibits a Needle-in-a-1 Haystack challenge. This challenge of finding a rare material, *i.e.*, the "needle", 2 inside a vast search space, *i.e.*, the "haystack", arises when there is an extreme 3 imbalance of optimum conditions relative to the size of the search space. For ex-4 ample, only 0.82% out of 146k total materials in the open-access Materials Project 5 database have a negative Poisson's ratio, a rare material property. However, current 6 state-of-the-art optimization algorithms are not designed with the capabilities to 7 find solutions to these challenging multidimensional Needle-in-a-Haystack prob-8 lems, resulting in slow convergence to a global optimum or pigeonholing into a 9 local minimum. In this paper, we present a Zooming Memory-Based Initialization 10 algorithm, entitled ZoMBI, that builds on conventional Bayesian optimization 11 principles to quickly and efficiently optimize Needle-in-a-Haystack problems in 12 both less time and fewer experiments by addressing the common convergence 13 and pigeonholing issues. ZoMBI actively extracts knowledge from the previously 14 best performing evaluated experiments to iteratively zoom in the sampling search 15 bounds towards the global optimum "needle" and then prunes the memory of 16 low-performing historical experiments to accelerate compute times by reducing 17 the algorithm time complexity from $O(n^3)$ to O(1), as the number of experiments 18 sampled increases. Additionally, ZoMBI implements two custom acquisition func-19 tions that use active learning to further guide the sampling of new experiments 20 21 towards the global optimum. We validate the algorithm's performance on two real-world 5-dimensional Needle-in-a-Haystack material property optimization 22 datasets: discovery of auxetic Poisson's ratio materials and discovery of high ther-23 moelectric figure of merit materials. The ZoMBI algorithm demonstrates compute 24 time speed-ups of 400x compared to traditional Bayesian optimization as well as 25 efficiently discovering materials in under 100 experiments that are up to 3x more 26 highly optimized than those discovered by current state-of-the-art algorithms. 27

1 Introduction to Rare Material Discovery

Current optimization algorithms perform well on low-dimensional problems that are smooth and have 29 wide basins of attraction. Examples of smooth manifolds with wide basins of attraction within material 30 science include process- and recipe-optimization problems such as tuning perovskite manufacturing 31 variables to achieve higher efficiency [1], optimizing microfluidics flow parameters to achieve ideal 32 droplet formation [2], optimizing silver nanoparticle recipes for optical properties [3], and tuning 33 34 perovskite compositions with physics-based constraints to maximize stability [4]. Optimization techniques like Bayesian optimization (BO) are well-suited to model these simple manifolds using 35 a Gaussian Process (GP) surrogate [5, 6, 7, 8, 9]. However, the performance of this BO with a GP 36 breaks down as the manifold complexity increases. Material property optimization problems that 37

Submitted to 36th Conference on Neural Information Processing Systems (NeurIPS 2022). Do not distribute.

have high technological significance, such as discovering materials with rare properties or materials 38 with a specific combination of properties, have search space manifolds that more closely resemble a 39 *Needle-in-a-Haystack* [10], shown in Figure 1(b), rather than a smooth or convex space. This Needle-40 in-a-Haystack (NiaH) problem arises when only few optimum conditions exist within the entire 41 search space, resulting in an extreme imbalance. Interpolating the parameter space of an imbalanced 42 search space with an estimation function, such as a GP, results in smoothing over the optimum or 43 over-predicting the properties of the materials found near the optimum [11, 12, 13]. Examples of 44 NiaH materials optimization problems include discovering auxetic materials (*i.e.*, materials that 45 have a highly negative Poisson's ratio, ν) for energy absorptive medical devices or protective armor 46 [14, 15, 16] and discovering materials that have a combination of high electrical conductivity and low 47 thermal conductivity (*i.e.*, a highly positive thermoelectric figure of merit, ZT) used from improving 48 sensor technology to enable ubiquitous solid-state cooling [17, 18, 19]. Both of these rare material 49 optimization problems are examples where an extreme data balance exists in the search space because 50 only a fraction of the total number of materials exhibit these rare properties [14, 20, 21, 22, 23]. This 51 NiaH optimization challenge of extremely imbalanced search spaces is largely applicable to many 52 53 fields, not just materials science, including the fields of ecological resource management [24], fraud detection [25, 26], and rare diseases [27, 26]. 54



(a) Process Optimization Manifold

(b) Materials Optimization Manifold

Figure 1: Archetype Manifolds in Materials Science Optimization. (a) Smooth and wide basin of attraction landscape that is common to process optimization problems. This 2D projected manifold is adapted from the 6D perovskite process optimization problem by Liu *et al.* [1]. (b) Rough and narrow basin of attraction landscape that is typical of material property optimization problems. This 2D projected manifold is obtained from the 5D negative Poisson's ratio optimization problem presented in this paper [20, 21].

55 Several challenges exist for the current landscape of computational tools that inhibit effective optimization of these complex NiaH problems. Firstly, the "needle" makes up only a small percentage 56 of the total manifold search space, resulting in a weak correlation between the measured input 57 parameters and the target property of interest, inhibiting discovery of the region containing the needle 58 [28, 29, 11]. This challenge requires the development of an algorithm that can more quickly determine 59 the plausible region of the manifold where the needle exists. The second challenge for algorithms, 60 such as BO, to optimize NiaH manifolds is in the nature of the acquisition function to pigeonhole 61 sampling into local minima because of the narrowness of the needle's basin of attraction [30, 31]. 62 Standard BO acquisition functions, including expected improvement (EI) [32] and lower confidence 63 bound (LCB) [7, 12], are static sampling techniques that only adjust sampling based on the output of 64 the surrogate model, which enacts smoothing of the needle [11, 5, 6]. To overcome this challenge, 65 active learning-based tuning of the acquisition function hyperparameters can be implemented to 66 improve the sampling quality and avoid pigeonholing. We design two active learning acquisition 67 functions, LCB Adaptive and EI Abrupt, further discussed in the Appendix (sections A.1 & A.2). 68

Lastly, there exists a computing challenge for NiaH problems where, typically, several thousands of 69 samples must be observed to find an optimum when using an algorithm that is poorly-suited to tackle 70 NiaH manifolds [10]. The compute time of BO using a GP surrogate scales with the complexity 71 $O(n^3)$, where n is the number of experiments sampled, hence, the compute time of traditional BO 72 blows up as more data is required to find the optimum [33, 34, 35, 5, 6, 36, 37]. To solve this 73 computing challenge, an algorithm must be designed that both efficiently optimizes the space in as 74 few experiments as possible and reduces the effect of compounding compute times over the length of 75 the optimization procedure. 76

77 1.1 Related Literature on Fast and Bounded Optimization

In recent literature, algorithms have been developed to address some of these challenges individually, 78 but not all of them together. The first class of solutions bound the search space using a trust region 79 approach to sample regions with higher probability of containing the optimum. Uber AI develop 80 TuRBO [38] that compiles a set of independent model runs, using separate GP surrogate models to 81 compute a new, smaller search region, narrowed in on the target optimum. Regis develops TRIKE 82 [39] that utilizes maximization of the EI acquisition function to bound a trust region containing the 83 global optimum. Diouane et al. develop TREGO [40], which interleaves sampling between global 84 and local search regions, where the local search regions are defined by the single best historical 85 experiment sampled. Although these methods offer solutions to one of the three challenges presented, 86 each method has its downfalls when optimizing NiaH problems. For example, TuRBO requires the 87 88 computation of several GP model runs, which increases compute time and also does not guarantee that the needle will be resolved due to interpolation effects; TRIKE is inflexible to the use of other 89 acquisition functions as it locks the user in to only using EI, which may pigeonhole into local 90 minima; TREGO uses only the best sampled experiment to define its search regions, which will yield 91 inconsistent or sub-optimal results when the needle consists of a fractional region of the manifold 92 and single point is unlikely to land in its basin of attraction. 93

The second class of solutions to the challenges presented in this paper are designed to decrease the 94 computing time required to run an optimization procedure. A common method for reducing the 95 compute time of BO with a GP surrogate is to introduce a sparse GP [5, 41, 36]. A sparse GP uses a 96 small subset of pseudo data, often denoted as m, to reduce the GP time complexity from $O(n^3)$ to 97 $O(nm^2)$ [42]. However, the process of selecting a useful subset requires minimizing the Kullback-98 Leibler divergence between the sparse GP and true posterior GP, which is often a computationally 99 intensive procedure of using variational inference [43]. In addition to sparse GPs, new algorithms 100 have been developed in literature to improve the compute time of optimization in various ways. 101 Van Stein et al. develop MiP-EGO [44], which parallelizes the function evaluations of efficient 102 103 global optimization (EGO) to discover optima faster and in fewer experiments using derivative-free computation [45]. Joy et al. [46] use directional derivatives to accelerate hyperparameter tuning 104 by 100x and achieve higher accuracy than the FABOLAS baseline by Klein et al. [47]. Zhang et 105 al. develop FLASH [48] to achieve optimization speed-ups of 50% by using a linear parametric 106 model to guide algorithm search within high-dimensional spaces. Snoek et al. [13] design a neural 107 network-based parametric model that reduces the overall time complexity of BO to O(n) compared 108 to the complexity of $O(n^3)$ of standard BO with a GP surrogate model. These existing methods 109 from literature within the class of solutions for accelerating compute time are generally introducing 110 external models necessary to perform optimization, such as neural networks, variational inference, 111 or parameteric models. While these external models do speed-up compute time, they often lack the 112 predictive capabilities to capture the weak correlation between measured input parameters and the 113 target property of interest in NiaH problems. We illustrate this mechanic later in the paper when 114 115 comparing the optimization results on two materials science NiaH problems of a fast algorithm MiP-EGO with that of TuRBO, an algorithm better suited for discovering optima within narrow basins 116 of attraction. 117

Although these methods from existing literature address some of the challenges in optimizing NiaH problems, none of them have been designed specifically to quickly and efficiently discover a needlelike optimum within a haystack of sub-optimal points, resulting in all of them falling short of full solution. Therefore, in this paper, we design an algorithm that addresses all three of the challenges faced when optimizing NiaH problems by (1) zooming in the manifold search bounds iteratively and independently for each dimension based on m number of best memory points to quickly converge to the plausible region containing the global optimum needle, (2) anti-pigeonholing into local minima

by using actively learned acquisition function hyperparameters to tune the exploitation-to-exploration 125 ratio, (3) relieving compute utilization by pruning the low-performing memory points not being 126 used to zoom in the search bounds. The proposed algorithm, entitled [Zo]oming [M]emory-[B]ased 127 [I]nitialization (ZoMBI), combines these three contributions into a method that efficiently optimizes 128 NiaH problems quickly. In essence, this process of scanning broadly and then focusing in on points 129 of interest based on memory was inspired by the way we humans solve similar problems, but stands 130 131 in contrast to the way standard BO methods with static acquisition functions solve problems. We demonstrate the performance of this algorithm on two NiaH materials science datasets: (1) discovery 132 of materials with negative Poisson's ratio and (2) discovery of materials with both high electrical 133 conductivity and low thermal conductivity. The performance of the proposed ZoMBI algorithm is 134 compared against standard BO with static acquisition functions and two state-of-the-art (SoTA) 135 algorithms, one from each of the two classes of partial NiaH solutions: (1) TuRBO (bounded search 136 space) and (2) MiP-EGO (faster compute). 137

138 2 Methodology: Bounded & Memory-Pruning Optimization

The [Zo]oming [M]emory-[B]ased [I]nitialization ZoMBI algorithm has two key features: (1) iterative 139 inward bounding of proceeding search spaces using the m number of best-performing memory 140 points within the prior search space and (2) iterative pruning of low-performing historical search 141 142 space memory. The newly computed search space bounds are unique for each dimension, such 143 that optimum basin of attraction of complex, non-convex NiaH manifolds can be discovered. This algorithm leverages these two key features to guide the acquisition of new data towards more optimal 144 regions while only fitting the surrogate within the suggested optimum region to resolve more detail 145 of the space of interest, as shown in Figure 2. This process subsequently reduces the compute time 146 significantly compared to the compute of a GP in a standard BO procedure, as shown in Figure 3. 147



Figure 2: **Zooming Search Bounds.** For every activation of ZoMBI, the search bounds are zoomed inward based on the prior best-performing memory points. A 4D Ackley function manifold is projected in 2D. The bounding regions of each 2D slice are illustrate by the red and orange boxes. The ϕ number forward experiments sampled are illustrated as black markers. The global optimum is indicated by the red region of the heatmap.

We define m as the number of retained memory points during an activation of ZoMBI. The m memory points are saved to memory while all other data are erased from memory. These are the historical data points that achieve the m lowest (for minimization) target values, y, and they are used to zoom in the search bounds. Using these memory points, the multi-dimensional upper and lower bounds of the zoomed search space are computed for each dimension, d. Let $\mathbf{X} := \{X_1, X_2, \ldots, X_n\}$ be a set of data points, where $X_j \in \mathbb{R}^d$. Let $f : \mathbb{R}^d \to \mathbb{R}$ be the objective function. We first assume that the points in **X** are in general position so that $f(\mathbf{X})$ contains unique elements. Then, for each $m \le n$ define $\mathbf{X}^{(m)} = \{X_{\pi(1)}, \dots, X_{\pi(m)}\}$ where π is a permutation on $\{1, \dots, n\}$ so that $\{f(X_{\pi(j)})\}$ is in ascending order. If $f(\mathbf{X})$ contains repeated elements, we may first remove the points with repeated f values and apply the definition above. Then, for each d, the bounds are defined as:

$$\mathcal{B}_{d}^{l} = \min_{X \in \mathbf{X}^{(m)}} \{ [X]_{d} \}$$

$$\mathcal{B}_{d}^{u} = \max_{X \in \mathbf{X}^{(m)}} \{ [X]_{d} \},$$
(1)

where \mathcal{B}_d^l and \mathcal{B}_d^u computed lower and lower bounds for each dimension, d, respectively. The bounds 158 $[\mathcal{B}_{d}^{l}, \mathcal{B}_{d}^{u}]$ constrain the proceeding acquisition of new data as well as the computation of a GP, such 159 that sampling cannot occur outsides of the bounded region. This constraining process operates 160 independently for each dimension, such that each dimension has a unique lower and upper bound. 161 To initialize the algorithm with data from the constrained space, i data points are sampled from the 162 bounded region using Latin Hypercube Sampling (LHS). LHS splits a d-dimensional space into i * d163 equally spaced strata, where i is the number of points to sample uniformly over d dimensions with 164 low variability, unlike random sampling that has high sampling variability [50]. A GP surrogate 165 model is retrained on these *i* LHS points sampled from the constrained space and then for every 166 proceeding experiment sampled from the space, denoted as a forward experiment, the surrogate model 167 is retrained. Thus, the GP is only being trained on information within the constrained region and as 168 the constrained region iteratively zooms inward and decreases in hypervolume, so does the region 169 computed by the GP. This process allows for more information to be resolve within regions plausibly 170 containing the global optimum basin of attraction. Up to ϕ forward experiments are sampled in serial, 171 where $\{X_i\} \cup \{X_{\phi}\} \subseteq \{X_n\}$. These forward experiments are sampled by maximizing an acquisition 172 value, $a \in [0, 1]$, computed by a user-selected acquisition function from one of the four functions EI, 173 EI Abrupt, LCB, and LCB Adaptive. Once $i + \phi$ number of experiments are sampled, the bounds are 174 175 re-constrained using the *m* best performing experiments, *i* new experiments are sampled from the 176 zoomed-in space using LHS, and then the memory is pruned. The process of collecting ϕ forward



Figure 3: **Wall-clock Compute Time.** The compute time per experiment is illustrate for traditional BO with a GP surrogate (orange) and for ZoMBI with a GP surrogate (blue) with the y-axis in log-scale. Five independent trials of each method were run to optimize a 5D Ackley function with a narrow basin of attraction using an NVIDIA Tesla Volta V100 GPU [49]. The averages of the trials are shown as solid orange and blue lines while the shaded regions indicate the maximum and minimum compute times bounds. The red dashed line indicates the trend of the ZoMBI compute times. The measured compute time includes the time to compute the GP surrogate model and the time to acquire an experiment from the surrogate.

experiments is repeated. A complete constraining-resetting iteration is denoted as an activation, α . This iterative zooming and pruning process over several α significantly speeds up compute time.

179 **3** Rare Material Discovery Results

180 3.1 Compute Time

In this section, we assess the compute time of the developed algorithm in comparison that of standard 181 182 BO methods. As more experiments are amassed and committed to memory to run traditional BO by computing a GP and an acquisition value, the compute time increases polynomially, following 183 the $O(n^3)$ time complexity of GP matrix multiplication [33, 5, 6, 51, 36, 37]. This complexity is 184 unfavorable as it leads to compounding compute times as more experiments are run. Therefore, we 185 implement a memory pruning feature into the ZoMBI algorithm that iteratively selects which prior 186 data points to keep and which ones to prune from the memory during each activation, α . Via memory 187 pruning, the number of experiments used to train the GP surrogate varies between $[i, i+\phi]$ for every α , 188 rather than being proportional to n. This is computationally favorable because $\{X_i\} \cup \{X_{\phi}\} \subseteq \{X_n\}$. 189 Thus, for a single α , the time complexity is $O((i + \phi)^3)$. However, since ϕ resets back to zero after 190 each α , a non-increasing sawtooth pattern in compute time is exhibited, hence, as $\alpha, n \to \infty$, the 191 complexity approaches O(1). Figure 3 illustrates that the sawtooth compute time pattern maps to 192 the resetting interval of ϕ , which trends towards a constant, non-increasing value over many α and 193 n. After collecting 1000 experiments, the compute time of traditional BO trend towards > 400194 195 seconds, whereas after 1000 experiments, the compute time ZoMBI trends towards a constant 1 second. Therefore, the memory pruning feature of ZoMBI accelerates the optimization compute time 196 by over 400x at n = 1000 and achieves further relative acceleration as n increases. The memory 197 pruning mechanic of ZoMBI drives fast compute times without sacrificing the ability to converge on 198 rare materials, demonstrated in the following sections (3.2 & 3.3). 199

200 3.2 Poisson's Ratio

We demonstrate the ability of the ZoMBI algorithm to optimize Needle-in-a-Haystack problems on two real-world datasets. The first dataset consists of 146k materials and the objective is to find the material with the minimum negative Poisson's ratio, ν . The second dataset consists of 1k materials and the objective is to find the material with the maximum thermoelectric merit, ZT, *i.e.*, a material with high electrical conductivity and low thermal conductivity. Both of these datasets are 5-dimensional and are obtained from the open-access Materials Project database [20].

The ν dataset exhibits a Needle-in-a-Haystack problem due to very few materials having negative 207 208 ν values [14, 20, 21, 15]. A positive $\nu > 0$, describes a material that expands when a compressive load is applied to the orthogonal direction [52, 53]. Conversely, a negative $\nu < 0$ describes a material 209 that contracts rather than expands when compressed in the orthogonal direction, denoted as an 210 auxetic material [14, 23] – a rare phenomenon that occurs in only 0.82% of materials within the 211 Materials Project database [20, 21]. Auxetic materials with highly negative Poisson's ratios have 212 energy absorptive properties, which are ideal materials for wearable medical devices and protective 213 armor that must absorb the energy of large impacts to keep bones from shifting or to inhibit the 214 penetration of the protective layer [15, 16]. Thus, for this NiaH problem, the objective is to discover 215 216 the material with the lowest ν value. Figure A.2 illustrates the spread of ν values within the raw dataset as a histrogram as well as a manifold generated by a Random Forest (RF) regression on the 217 raw dataset using 500 trees. The search space generated by the RF is noisy and non-convex with 218 narrow basins of attraction containing each optimum, resulting in a challenging NiaH optimization 219 problem. The ground truth "needle" materials with the lowest ν values are Li₂NbF₆ with $\nu \approx -1.7$ 220 221 and Na₂CO₃ with $\nu \approx -1.2$.

Figure 4 illustrates the performance of ZoMBI in discovering the lowest ν -value material, compared 222 to the SoTA TuRBO and MiP-EGO algorithms. The ZoMBI algorithm is run with each of the four 223 acquisition functions: LCB, LCB Adaptive, EI, and EI Abrupt. In under 100 evaluated experiments, 224 225 LCB and LCB Adaptive discover one of the needles within the dataset (Li_2NbF_6) and, similarly, EI Abrupt discovers the other needle (Na₂CO₃). The distribution of ν values for the final experiment 226 across all ensemble runs is illustrated for each method to highlight the sampling density and general 227 rate of success. LCB Adaptive and EI Abrupt are the first two implementations of ZoMBI to discover 228 a $\nu < 0$ material because of their ability to actively tune their sampling hyperparameters. After 229



Figure 4: **Discovery of Rare Negative Poisson's Ratio Materials.** The optimization objective is to find the material with the minimum Poisson's ratio, ν_{\min} , in 100 experiments. The green, blue, red, and orange lines indicate the median best running evaluated sample of ZoMBI using the LCB, LCB Adaptive, EI, and EI Abrupt acquisition functions, respectively. The pink and black lines indicate the median best running evaluated sample of the SoTA methods, MiP-EGO and TuRBO, respectively. The median for each method is taken over the best 12 independent model runs. The shaded regions indicate the variance between model runs. The crosshatched region indicates the space discovered by standard BO methods, without the use of ZoMBI. The dashed black line indicates the $\nu = 0$ inflection point. The distribution of the final sampled ν value for each method at the 100th experiment is shown as a kernel density estimation with a 0.5 smoothing factor. The materials formulae and unit cells that have the closest evaluated ν value discovered by each ZoMBI method at the end of the 100 experiments are illustrated.

30 experiments, the ZoMBI search bounds have zoomed inward enough for the explorative LCB 230 acquisition function to discover a region of the manifold containing highly negative ν material, 231 eventually leading to the global minimum needle. These three implementations of ZoMBI: LCB, 232 LCB Adaptive, and EI Abrupt, have a steep drop in the discovered ν value, allowing these methods 233 to discover an optimum fast, in fewer experiments than both SoTA methods. Overall, LCB and 234 LCB Adaptive implementations of ZoMBI discover the most optimum minimum $\nu \approx -1.7$, while 235 the SoTA algorithms TuRBO and MiP-EGO only discover $\nu \approx -0.55$ and $\nu \approx -0.20$, respectively. 236 These results demonstrate that with proper selection an acquisition function, ZoMBI achieves better 237 performance and a higher success rate than SoTA on optimizing this real-world materials science 238 NiaH problem. 239

240 3.3 Thermoelectrics

The ZT dataset exhibits a Needle-in-a-Haystack problem, similar to the ν dataset because very few materials have high ZT values [20, 10]. However, rather than ZT being a directly measurable mechanical material property like Poisson's ratio, ZT must be computed using a combination of several thermal and electrical material properties [54]:

$$ZT = \frac{S^2 \sigma}{\kappa} T,$$
(2)

where S is the Seebeck coefficient, σ is electrical conductivity, T is the average temperature, and 245 κ is thermal conductivity. The ZT is computed for each material in the Materials Project database 246 using BoltzTraP [55]. Of the initial 146k materials, 1k of them have the required thermal and 247 electrical properties to compute a ZT value. ZT is a common figure of merit used to describe 248 the thermal-to-electrical or electrical-to-thermal conversion efficiency of thermoelectric materials 249 [56, 57, 58, 59]. A higher ZT indicates that the material is better able to convert a thermal gradient 250 into an electrical current [54]. Materials with large ZT values have a range of applications from 251 usage as solid-state cooling devices to being used as sensors that when heated up, will produce an 252 electrical signal [17, 18, 19]. For this NiaH problem, the objective is to discover the material with 253 the highest ZT value. Figure A.3 illustrates the spread of ZT values within the raw dataset as a 254 hisitogram, as well as a manifold generated by an RF regression on the raw dataset using 500 trees. 255 Similar to the ν manifold, the ZT manifold is noisy and non-convex with narrow basins of attraction 256 [20, 55]. The ground truth "needle" materials with the highest ZT values are $Na_4Al_3Ge_3IO_{12}$ with 257 $ZT \approx 1.4$ and $Sr_4Al_6SO_{12}$ with $ZT \approx 1.9$. 258



Figure 5: **Discovery of Rare Positive Thermoelectric Merit Materials.** The optimization objective is to find the material with the maximum thermoelectric merit, ZT_{max} , in 100 experiments. The green, blue, red, and orange lines indicate the median best running evaluated sample of ZoMBI using the LCB, LCB Adaptive, EI, and EI Abrupt acquisition functions, respectively. The pink and black lines indicate the median best running evaluated sample of the SoTA methods, MiP-EGO and TuRBO, respectively. The median for each method is taken over the best 12 independent model runs. The shaded regions indicate the variance between model runs. The crosshatched region indicates the space discovered by standard BO methods, without the use of ZoMBI. The distribution of the final sampled ZT value for each method at the 100th experiment is shown as a kernel density estimation with a 0.5 smoothing factor. The materials formulae and unit cells that have the closest evaluated ZT value discovered by each ZoMBI method at the end of the 100 experiments are illustrated.

Figure 5 illustrates the performance of ZoMBI in discovering the highest ZT-value material, compared to the SoTA TuRBO and MiP-EGO algorithms. Initially, we see TuRBO outperform all other algorithms, but then it is unable to accelerate its sampling towards the needle basins of attraction. Similarly,

MiP-EGO gets trapped in a local minimum and is unable to escape. Conversely, after 50 evaluated 262 experiments, ZoMBI LCB Adaptive and EI Abrupt supersede TuRBO and quickly discover high ZT263 materials, illustrating the advantage of active learning acquisition functions. Although the active 264 learning acquisition functions prove to be more successful than the SoTA algorithms, none of the 265 tested algorithms are able to discover the maximum global needle, $Sr_4Al_6SO_{12}$, only the second best 266 needle, $Na_4Al_3Ge_3IO_{12}$. This result is likely due to the data imbalance being too extreme that far out 267 on the tail of the ZT dataset, in turn, generating an RF manifold complexity too high, even for ZoMBI. 268 Hence, indicating that there are limitations in the manifold complexity that ZoMBI can optimize, and 269 further illustrating that convergence on the global optimum needle is not guaranteed using this method. 270 However, for the ZT dataset, the LCB Adaptive implementation of ZoMBI discovers the second 271 best needle, Na₄Al₃Ge₃IO₁₂ with $ZT \approx 1.4$, while the SoTA algorithms TuRBO and MiP-EGO only 272 discover $ZT \approx 0.65$ and $ZT \approx 0.45$, respectively. Thus, LCB Adaptive demonstrates the highest 273 performing optimization results across both of the real-world NiaH datasets, discovering the most 274 optimal materials the fastest for both the ν and ZT datasets. 275

276 4 Summary & Conclusion

In this paper, we proposed the [Zo]oming [M]emory-[B]ased [I]nitialization (ZoMBI) algorithm 277 that builds on the principles of Bayesian optimization to accelerate the discovery of rare materials 278 by two-fold, firstly by requiring fewer experiments to achieve a better optimum than state-of-the-279 art, and secondly by pruning the memory of low-performing historical experiments to speed-up 280 compute time. The ZoMBI algorithm exceeds state-of-the-art performance on optimizing Needle-in-281 a-Haystack datasets by (1) using the values of the m best performing previously sampled memory 282 points to iteratively zoom in the search bounds of the manifold uniquely on each dimension and 283 (2) implementing two custom acquisition functions, LCB Adaptive and EI Abrupt, that actively 284 learn information about the manifold during optimization to tune the sampling of new experimental 285 conditions from a surrogate. The main contributions of this algorithm solve three fundamental 286 challenges of optimizing non-convex Needle-in-a-Haystack problems: (1) the challenge of locating 287 the hypervolume region of the manifold containing the narrow global optimum basin of attraction 288 [28, 29, 11] is alleviated by introducing iterative search bounds based on learned knowledge of the 289 manifold; (2) unwanted pigeonholing into local minima [30, 31, 5, 6] is avoided by both the zooming 290 mechanics of ZoMBI as well as using the two acquisition functions developed in his paper, LCB 291 Adaptive and EI Abrupt, that tune their hyperparameters through active learning; (3) the challenge 292 of polynomially increasing compute times of BO using a GP surrogate [33, 34, 35, 5, 6, 36, 37] is 293 addressed by actively pruning the retained memory of the algorithm after each activation, α , in turn, 294 reducing the time complexity from $O(n^3)$ to O(1) as $\alpha, n \to \infty$. By developing the ZoMBI algorithm 295 296 to solve these challenges, it becomes possible to quickly and efficiently find optimal solutions to complex Needle-in-a-Haystack problems in fewer experiments. Hence, this tool can be applied to rare 297 material discovery, a class of data imbalanced Needle-in-a-Haystack problems, to enable widespread 298 discovery of new materials with important technical applications from designing high-performance 299 medical devices to engineering ubiquitous solid-state cooling systems. 300

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