



Evolutionary Algorithms

- Population-based, stochastic search algorithms
- Exploitation: selection
- Exploration: mutation & crossover

Model-Based Evolutionary Algorithms

- Population-based, stochastic search algorithms
- Exploitation: selection
- Exploration:
 - 1. Learn a model from selected solutions
 - 2. Generate new solutions from the model (& population)

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What ?

Model-Based Evolutionary Algorithms (MBEA)

- a.k.a. Estimation of Distribution Algorithms (EDAs)
- ▶ a.k.a. Probabilistic Model-Building Genetic Algorithms
- > a.k.a. Iterated Density Estimation Evolutionary Algorithms

MBEA = Evolutionary Computing + Machine Learning

Note: model not necessarily probabilistic

Why ? Goal: Black Box Optimization Little known about the structure of the problem Clean separation optimizer from problem definition Easy and generally applicable Approach * Classical EAs: need suitable representation & variation operators * Model-Based EAs: learn structure from good solutions

Discrete Representation Typically binary representation Higher order cardinality: similar approach Current Selected New population population population 11001 01111 11001 **Probabilistic** 11001 11101 10101 01011 Model 11011 11000 00111



Univariate PMBGA

Model

- * Model: probability vector $[p_1, ..., p_\ell]$ (ℓ : string length)
- * p_i : probability of value 1 at string position i
- * $p(X) = \prod_{i=1}^{\ell} p(x_i) (p(x_i): \text{ univariate marginal distribution})$
- Learn model: count proportions of 1 in selected population
- Sample model: generate new solutions with specified probabilities





A hard problem for the univariate FOS

:	000	000	-	Marginal Product (MP) FOS				_
	111	111	_	Ŷ	$(X_0X_1\lambda$	(₂)	$(X_3 X_4 X_5)$)
	010	101	_	000	0.3		0.3	_
	1010	010		001	0.0		0.0	
	000	010		010	0.2		0.2	
	111	000		011	0.0		0.0	
	010	111		100	0.0		0.0	
	111	000		101	0.1		0.1	
	000	111		110	0.0		0.0	
	111	111	_	111	0.4		0.4	
			ι	Univariate FOS				
		$\hat{P}(X_0)$	$\hat{P}(X_1)$	$\hat{P}(X_2)$	$\hat{P}(X_3)$	$\hat{P}(X_4)$	$\hat{P}(X_5)$	
	0	0.5	0.4	0.5	0.5	0.4	0.5	
	1	0.5	0.6	0.5	0.5	0.6	0.5	
What is	the	proba	ability	of ger	eratin	g 111	111?	
Univariat	te F	OS: 0).5 · 0.	6 · 0.5	· 0.5 ·	0.6 • 0	0.5 = 0	.02
							N	

Learning problem structure on the fly Without a "good" decomposition of the problem, important partial solutions (building blocks) are likely to get disrupted in variation. Disruption leads to inefficiency. Can we automatically configure the model structure favorably? Selection increases proportion of good building blocks and thus "correlations" between variables of these building blocks. So, learn which variables are "correlated". See the population (or selection) as a data set. Apply statistics / probability theory / probabilistic modeling.

Bivariate PMBGA Model • Need more than just probabilities of bit values • Model pairwise interactions: conditional probabilities • MIMIC (de Bonet, Isbell, Viola; 1996) • Dependency Chain • COMIT (Baluja, Davies; 1997) • Dependency Tree • BMDA (Pelikan , Mühlenbein; 1998) • Independent trees (forest)

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Bivariate PMBGA

MIMIC

- Model: chain of pairwise dependencies.
- $p(X) = \prod_{i=1}^{\ell-1} p(x_{i+1}|x_i) p(x_1).$
- MIMIC greedily searches for the optimal permutation of variables that minimizes Kullack-Leibler divergence.

Bivariate PMBGA

COMIT

- Optimal dependency tree instead of linear chain.
- Compute fully connected weighted graph between problem variables.
- ► Weights are the mutual information *I*(*X*, *Y*) between the variables.
- ► $I(X, Y) = \sum_{y \in Y} \sum_{x \in X} p(x, y) \log \frac{p(x, y)}{p(x)p(y)}$.
- COMIT computes the maximum spanning tree of the weighted graph.

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Bivariate PMBGA DSMGA Dependency Structure Matrix Genetic Algorithm (Yu, Goldberg, Sastry, Lima, Pelikan; 2009) Dependency Structure Matrix (DSM) contains the information of pairwise interactions. DSMGA constructs the DSM by using mutual information metric. DSM clustering aims to transfer the pair-wise interaction information into higher-order interaction information. DSM Clustering Metric based on the minimum description length principle (MDL).

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Bivariate PMBGA

DSMGA-II

- Extended version \Rightarrow DSMGA-II (Hsu, Yu; 2015).
- DSMGA-II consists of four major components:
 - 1. pair-wise linkage detection
 - 2. model building
 - 3. restricted mixing
 - 4. back mixing
- Clustering the DSM leads to the Incremental Linkage Set: starting from one gene, incrementally add the next most dependent gene one-by-one.
- Restricted mixing: focus on building-block supply.
- Back mixing: when no improvement occurs, switch to the equal-acceptance criterion to reduce unnecessary evaluations on plateaus.

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Multivariate PMBGA

Marginal Product Model

- Extended Compact GA (ECGA) (Harik; 1999) was first EDA going beyond pairwise dependencies.
- Greedily searches for the Marginal Product Model that minimizes the minimum description length (MDL).
- ► $p(X) = \prod_{g=1}^{G} p(X_g)$
- Choose the probability distribution with the lowest MDL score.
- ► Start from simplest model: the univariate factorization.
- Join two groups that result in the largest improvement in the used scoring measure.
- ► Stop when no joining of two groups improves the score further.

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Multivariate PMBGA

Learning MP model

Start from univariate FOS:

 {{0}, {1}, {2}, ..., {*l* - 2}, {*l* - 1}}

 All possible pairs of partitions are temporarily merged:

 {{0, 1}, {2}, ..., {*l* - 2}, {*l* - 1}}
 {{0, 2}, {1}, ..., {*l* - 2}, {*l* - 1}}
 {{0, 2}, {1}, ..., {*l* - 2}, {*l* - 1}}
 {{0}, {1, 2}, ..., {*l* - 2}, {*l* - 1}}
 {{0}, {1, 2}, ..., {*l* - 2, {*l* - 1}}
 {{0}, {1}, {2}, ..., {*l* - 2, *l* - 1}}

 Compute MDL score of each factorization.
 Choose the best scoring factorization if better than current.
 Repeat until no better scoring factorization is found.

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Multivariate PMBGA Bayesian Network Probability vector, dependency tree, and marginal product model are limited probability models. Bayesian network much more powerful model. Acyclic directed graph. Nodes are problem variables. Edges represent conditional dependencies.









Multivariate PMBGA

Markov Network

- Markov Netwok EDA (MN-EDA: Santana, 2005) (DEUM: Shakya & McCall, 2007).
- Probability model is undirected graph.
- Factorise the joint probability distribution in cliques of the undirected graph and sample it.
- Most recent version: Markovian Optimisation Algorithm (MOA) (Shakya & Santana, 2008).
- MOA does not explicitly factorise the distribution but uses the local Markov property and Gibbs sampling to generate new solutions.

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Linkage Tree Learning

- This agglomerative hierarchical clustering algorithm is computationally efficient.
- ► Only the mutual information between pairs of variables needs to be computed once, which is a O(ℓ²) operation.
- The bottom-up hierarchical clustering can also be done in $O(\ell^2)$ computation by using the *reciprocal nearest neighbor chain* algorithm.

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Benchmark problems

- Hierarchical If-and-only-iFF (HIFF) (Watson, Hornby and Pollack, 1998)
- Computed over multiple layers
- Nodes are combined as a perfectly balanced binary tree
- Problem lengths are powers of two (i.e., $\ell = 2, 4, 8, 16, 32, ...$)
- Each variable is considered to be a leaf
- Leaf contributes 1
- ► Internal node contributes 2^{height} if children both 0 or both 1
- ▶ Internal node is 0 if children both 0; 1 if both 1; NIL else

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Benchmark problems

- Weighted MAX-CUT (Karp, 1972)
- Given a weighted graph (V, E), divide nodes into two sets so that total weight of edges between sets is maximized
- Identify binary variable x_i with each node v_i

$$f_{\text{Weighted MAX-CUT}}(\mathbf{x}) = \sum_{(v_i, v_j) \in E} \begin{cases} w_{ij} & \text{if } x_i \neq x_j \\ 0 & otherwise \end{cases}$$

- For now, considered as a black-box problem (no partial evaluations allowed)
- ► 5 instance types:
 - Fully connected graphs (β -distributed, $\alpha = 100$, $\beta = 1$)
 - > 2D Square-grid graphs (β -distributed, $\alpha = 100$, $\beta = 1$)
 - ▶ 3D Square-torus graphs (β -distributed, $\alpha = 100$, $\beta = 1$)
 - Uniformly distributed in a box, fully connected
 - Uniformly distributed in a box, $\lfloor \sqrt{(\ell)} \rfloor$ nearest neighbors

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Experiments: conclusion

- LTGA (= GOMEA with LT FOS) very efficient on Deceptive Trap function, Nearest-Neighbor NK landscape, and Hierarchical Trap function.
- Other FOS models possible: Linkage Neighborhood OM (Bosman & Thierens, 2012).
- Linkage Tree seems to be good compromise between FOS model complexity and search efficiency.

Parameter-less Population Pyramid



- Similar to Harik-Lobo scheme: eliminates population-size parameter
- Each level of a pyramid-like structure is a population of solutions.
- Solutions are always hill-climbed.
- ► All solutions encountered are stored in the pyramid structure.
- At each level a Linkage Tree GA is run.
- Solutions climb the pyramid ladder with increasing fitness.
- ▶ Whenever a solution enters a level the linkage tree is relearned.

Predetermined vs. Learned FOS

- Problem structure unknown: learn a FOS model.
- Problem structure Information available: predetermined FOS model.
- What is a good predetermined FOS model ?
- Direct mapping of dependency structure of problem definition to a predetermined FOS model ?
- Predetermined linkage models mirroring the static structure of the problem not sufficient (Thierens & Bosman, 2012).
- Dynamically learned tree model superior to mirror structured models and to static tree model.
- Question: is there an optimal, predetermined linkage model that outperforms the learned (tree) model ?

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Conclusions

- "Blind" Evolutionary Algorithms are limited in their capability to detect and mix/exploit/re-use partial solutions (building blocks).
- One requires luck or analyzing and designing ways of structure exploitation directly into problem representation and search operators.
- ► Having a configurable model can help overcome this.
- Algorithm then must learn to configure the model and thereby exploit structure online during optimization (e.g. EDAs, OMEAs).



















EDAs based on the Normal Distribution

► Make decisions based on better fit and increased complexity (e.g. P̂(X₀, X₁) vs. P̂(X₀)P̂(X₁))

















Illustration on the 1-D sphere function

 $\mathfrak{F}(\mathbf{x}) = x_0^2$







$$\lim_{t\to\infty} \left\{ \hat{\sigma}(t) \right\} = \lim_{t\to\infty} \left\{ \hat{\sigma}(0) c(\tau)^t \right\} = 0$$

This limits mean shift to a fixed factor times initial spread!

$$\lim_{t o\infty}\left\{\hat{\mu}(t)
ight\}=\hat{\mu}(0)+rac{d(au)}{1-\sqrt{c(au)}}\hat{\sigma}(0) \;,$$

- $c(\tau)$ and $d(\tau)$ functions of
 - $\phi()$ (standard normal distribution) and
 - Φ() (inverse cumulative normal distribution)

















AMaLGaM, CMA-ES, NES, and RP

- Differences are only in how the distribution is obtained. Where AMaLGaM uses maximum-likelihood estimates from the current generation, CMA-ES and NES base estimates on differences between subsequent generations as well as many elaborate enhancements (see tutorial on CMA-ES) and RP uses ensembles of random projections to lower dimensions to estimate covariance matrices more efficiently.
- On typical unimodal benchmark problems (sphere, (rotated) ellipsoid, cigar, etc) these algorithms exhibit polynomial scalability in both minimally required population size and required number of function evaluations
- CMA-ES, NES scale better than AMaLGaM on such problems





Running length / dimension







Permutation Model-Based Evolutionary Algorithms

- Building permutation models directly not straightforward
- Potential aid in the form of random keys (Bean (1997))
- Random keys encode permutations in real-valued space (via sorting)



- Real-valued approaches can thus be used directly
 - Bosman and Thierens (2001) (normal EDA)
 - Larrañaga et al (2001) (normal EDA)
- Inefficient scale-up behavior on deceptive additively decomposable relative ordering problems
- Highly redundant encoding that is hard to model with a normal distribution

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Permutation Model-Based Evolutionary Algorithms

- Binary/Integer representations are discrete, but also Cartesian
- Other discrete search spaces exist that are non-Cartesian
- Most notably: permutation-based problems
- Important real-world relevance, e.g. routing and scheduling
- Brings different challenges than Cartesian spaces however
 - Relative ordering problems
 - Absolute ordering problems
 - Neighbor ordering problems
 - Combinations of these
- Different types of models are more suited for specific types of ordering problem

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Permutation Model-Based Evolutionary Algorithms

- Generate instance for each subset of variables independently
- Then map to the real-valued domain using random keys and then translate the entire string into a valid permutation
- Preserves relative ordering of variables in subsets
- Can sample directly instead of using crossover (crossover still more robust however)
- Scales polynomially and much better than normal-pdf induced crossover

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Permutation Model-Based Evolutionary Algorithms Edge-histogram based sampling Tsutsui, Pelikan, and Goldberg (2003) Maps well to problems with neighboring variable relations Model is a matrix with probabilities of edges Matrix needs to be adjusted while sampling For problems with neighboring relations works better than random keys

Permutation Model-Based Evolutionary Algorithms

- Gaussian "equivalent" in permutation space: Mallows model (GM-EDA)
 - ► Ceberio, Mendiburu, and Lozano (2011)
- Requires a distance measure between permutations and a central permutation
- Also requires a spread parameter (not estimated from data)
- Most commonly used distance: Kendall- τ , allows factorization
- Finding central permutation is NP-hard however
- ▶ Fast heuristics are possible (linear in *l* and *n*)
- Final parameter estimation and sampling are not trivial and require dedicated algorithms
- First results are promising (permutation flow shop), outperforming Tsutsui
 - Ceberio, Irurozki, Mendiburu, and Lozano (2014)

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- Alternative approach: grammar-based
- Start with basic production rules
- Learning: assign probabilities to rules and increase complexity and specificity of rules using heuristics
- Sampling: select probabilistically from appropriate production rules
- Results are promising in that less function evaluations are often needed than standard GP, but time-complexity is (much) larger
 - Shan, McKay, Baxter, Abbass, and Essam (2003)
 - Bosman and de Jong (2004)
 - Shan, McKay, Baxter, Abbass, Essam, and Hoai (2004)
 - Hasegawa and Iba (2007)

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Model-Based Genetic Programming

Impossible to cover everything in this tutorial, see literature

▶ Kim, Shan, Nguyen, and McKay (2014)



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Multi-objective Model-Based Evolutionary Algorithms

- Each distribution explores own region
- Learning may however by incremental (CMA-ES, iAMaLGaM, iBOA, etc)
- Assign each distribution own adaptive incremental mechanisms
- Cannot combine directly with clustering each generation
- Need correspondence over generations
- Number of clusters fixed beforehand (k)

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Conclusions

- "Blind" metaheuristics are limited in their capability to detect and mix/exploit/re-use structural features of an optimization problem (e.g. partial solutions, building blocks, promising search directions, etc).
- One requires luck or analyzing and designing ways of structure exploitation directly into problem representation and search operators.
- Having a configurable model can help "overcome" this / help to do this automatically.
- Algorithm then must learn to configure the model and thereby exploit structure online during optimization.
- Having an explicitly tunable model can really help

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Conclusions Efficient model-based evolutionary algorithms (EDAs/IDEAs/PMBGAs/OMEAs) exist Binary/Integer/Permutation/Real-valued/GP & multi-objective Research is ongoing Especially useful when optimizing from a black-box perspective (e.g. complex simulations) Also useful from a white-box perspective Can learn more about the problem through learnt models Models configurable by hand (remove "expensive" learning overhead)

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Books Larrañaga and Lozano (eds) (2001). Estimation of Distribution Algorithms: A New Tool for Evolutionary Computation. Kluwer. Lozano, Larrañaga, Inza, Bengoetxea (2006). Towards a New Evolutionary Computation: Advances on Estimation of Distribution Algorithms, Springer. Pelikan, Sastry, Cantú-Paz (eds) (2006). Scalable Optimization via Probabilistic Modeling: From Algorithms to Applications, Springer.

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