

Tutorial: CMA-ES — Evolution Strategies and Covariance Matrix Adaptation

Anne Auger & Nikolaus Hansen

INRIA Research Centre Saclay – Île-de-France
Project team TAO
University Paris-Sud, LRI (UMR 8623), Bat. 490
91405 ORSAY Cedex, France

Copyright is held by the author/owner(s).
GECCO'11, July 12–16, 2011, Dublin, Ireland.
ACM 978-1-4503-0690-4/11/07.

Content

- 1 Problem Statement
 - Black Box Optimization and Its Difficulties
 - Non-Separable Problems
 - Ill-Conditioned Problems
- 2 Evolution Strategies
 - A Search Template
 - The Normal Distribution
 - Invariance
- 3 Step-Size Control
 - Why Step-Size Control
 - One-Fifth Success Rule
 - Path Length Control (CSA)
- 4 Covariance Matrix Adaptation
 - Covariance Matrix Rank-One Update
 - Cumulation—the Evolution Path
 - Covariance Matrix Rank- μ Update
- 5 Theoretical Foundations
- 6 Experiments
- 7 Summary and Final Remarks

Problem Statement

Continuous Domain Search/Optimization

- Task: **minimize** an **objective function** (*fitness function, loss function*) in continuous domain

$$f : \mathcal{X} \subseteq \mathbb{R}^n \rightarrow \mathbb{R}, \quad x \mapsto f(x)$$

- **Black Box** scenario (direct search scenario)



- gradients are not available or not useful
- problem domain specific knowledge is used only within the black box, e.g. within an appropriate encoding
- Search **costs**: number of function evaluations

Problem Statement

Continuous Domain Search/Optimization

- Goal
 - fast convergence to the global optimum
 - solution x with **small function value** $f(x)$ *... or to a robust solution x with **least search cost***
there are two conflicting objectives
- Typical Examples
 - shape optimization (e.g. using CFD)
 - model calibration
 - parameter calibration*curve fitting, airfoils, biological, physical controller, plants, images*
- Problems
 - exhaustive search is infeasible
 - naive random search takes too long
 - deterministic search is not successful / takes too long

Approach: stochastic search, Evolutionary Algorithms

Objective Function Properties

We assume $f : \mathcal{X} \subset \mathbb{R}^n \rightarrow \mathbb{R}$ to be *non-linear, non-separable* and to have at least moderate dimensionality, say $n \not\ll 10$.

Additionally, f can be

- non-convex
- multimodal
- non-smooth
- discontinuous
- ill-conditioned
- noisy
- ...

there are possibly many local optima

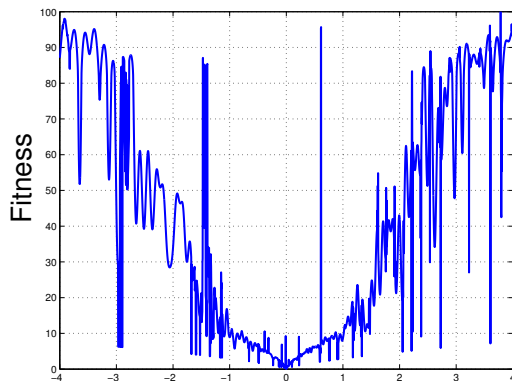
derivatives do not exist

Goal : cope with any of these function properties

they are related to real-world problems

Ruggedness

non-smooth, discontinuous, multimodal, and/or noisy

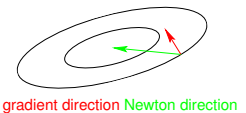
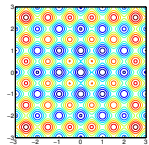
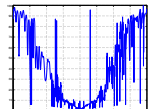
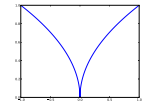


cut from a 5-D example, (easily) solvable with evolution strategies

What Makes a Function Difficult to Solve?

Why stochastic search?

- non-linear, non-quadratic, non-convex
on linear and quadratic functions much better search policies are available
- ruggedness
non-smooth, discontinuous, multimodal, and/or noisy function
- dimensionality (size of search space)
(considerably) larger than three
- non-separability
dependencies between the objective variables
- ill-conditioning



Curse of Dimensionality

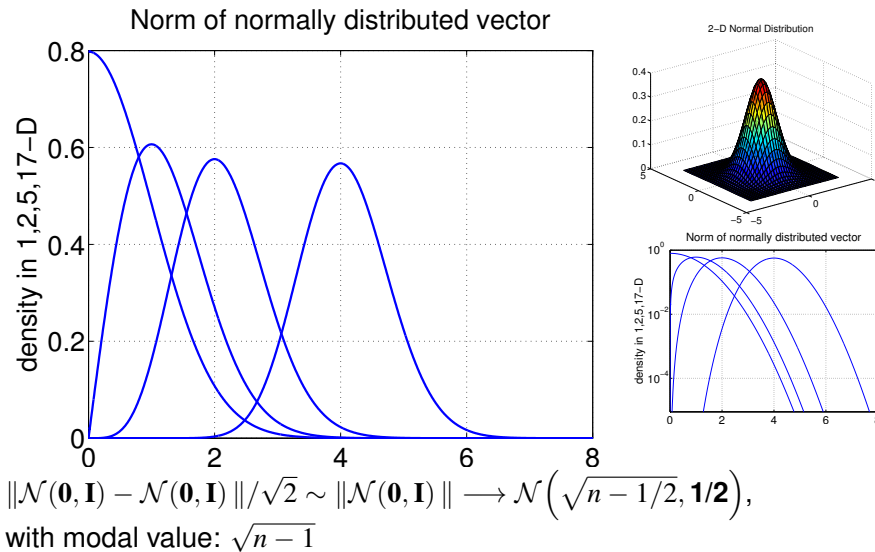
The term *Curse of dimensionality* (Richard Bellman) refers to problems caused by the **rapid increase in volume** associated with adding extra dimensions to a (mathematical) space.

Example: Consider placing 100 points onto a real interval, say $[0, 1]$. To get **similar coverage**, in terms of distance between adjacent points, of the 10-dimensional space $[0, 1]^{10}$ would require $100^{10} = 10^{20}$ points. A 100 points appear now as isolated points in a vast empty space.

Remark: **distance measures** break down in higher dimensionalities (the central limit theorem kicks in)

Consequently, a **search policy** (e.g. exhaustive search) that is valuable in small dimensions **might be useless** in moderate or large dimensional search spaces.

Curse of Dimensionality: Example



Separable Problems

Definition (Separable Problem)

A function f is separable if

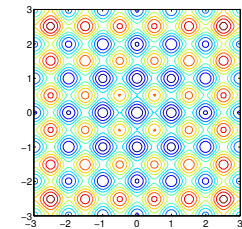
$$\arg \min_{(x_1, \dots, x_n)} f(x_1, \dots, x_n) = \left(\arg \min_{x_1} f(x_1, \dots), \dots, \arg \min_{x_n} f(\dots, x_n) \right)$$

$\Rightarrow$ it follows that f can be optimized in a sequence of n independent 1-D optimization processes

Example: Additively decomposable functions

$$f(x_1, \dots, x_n) = \sum_{i=1}^n f_i(x_i)$$

Rastrigin function



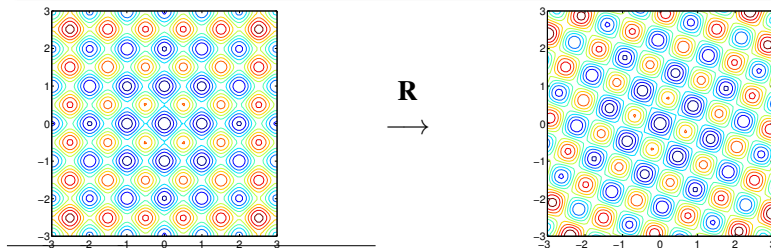
Non-Separable Problems

Building a non-separable problem from a separable one ^(1,2)

Rotating the coordinate system

- $f : x \mapsto f(x)$ separable
- $f : x \mapsto f(\mathbf{R}x)$ non-separable

$\mathbf{R}$ rotation matrix



¹ Hansen, Ostermeier, Gawelczyk (1995). On the adaptation of arbitrary normal mutation distributions in evolution strategies: The generating set adaptation. Sixth ICGA, pp. 57-64, Morgan Kaufmann

² Salomon (1996). "Reevaluating Genetic Algorithm Performance under Coordinate Rotation of Benchmark Functions: A survey of some theoretical and practical aspects of genetic algorithms." BioSystems, 39(3):263-278

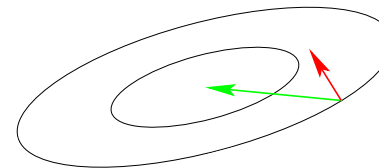
Ill-Conditioned Problems

Curvature of level sets

Consider the convex-quadratic function

$$f(\mathbf{x}) = \frac{1}{2}(\mathbf{x} - \mathbf{x}^*)^T \mathbf{H}(\mathbf{x} - \mathbf{x}^*) = \frac{1}{2} \sum_i h_{i,i} x_i^2 + \frac{1}{2} \sum_{i \neq j} h_{i,j} x_i x_j$$

$\mathbf{H}$ is Hessian matrix of f and symmetric positive definite



gradient direction $-f'(\mathbf{x})^T$

Newton direction $-\mathbf{H}^{-1}f'(\mathbf{x})^T$

Ill-conditioning means **squeezed level sets** (high curvature). Condition number equals nine here. Condition numbers up to 10^{10} are not unusual in real world problems.

If $\mathbf{H} \approx \mathbf{I}$ (small condition number of $\mathbf{H}$) first order information (e.g. the gradient) is sufficient. Otherwise **second order information** (estimation of $\mathbf{H}^{-1}$) is necessary.

What Makes a Function Difficult to Solve?

... and what can be done

The Problem	The Approach in ESs and continuous EDAs
Dimensionality, Non-Separability	exploiting the problem structure locality, neighborhood, encoding
Ill-conditioning	second order approach changes the neighborhood metric
Ruggedness	non-local policy, large sampling width (step-size) as large as possible while preserving a reasonable convergence speed stochastic, non-elitistic, population-based method recombination operator serves as repair mechanism restarts

... metaphors

Evolution Strategies

- 1 Problem Statement
 - Black Box Optimization and Its Difficulties
 - Non-Separable Problems
 - Ill-Conditioned Problems
- 2 Evolution Strategies
 - A Search Template
 - The Normal Distribution
 - Invariance
- 3 Step-Size Control
 - Why Step-Size Control
 - One-Fifth Success Rule
 - Path Length Control (CSA)
- 4 Covariance Matrix Adaptation
 - Covariance Matrix Rank-One Update
 - Cumulation—the Evolution Path
 - Covariance Matrix Rank- μ Update
- 5 Theoretical Foundations
- 6 Experiments
- 7 Summary and Final Remarks

Metaphors

Evolutionary Computation

individual, offspring, parent $\longleftrightarrow$

population
fitness function

generation

Optimization

candidate solution
decision variables
design variables
object variables

set of candidate solutions
objective function
loss function
cost function
iteration

... methods: ESs

Evolution Strategies

A Search Template

Stochastic Search

A black box search template to minimize $f : \mathbb{R}^n \rightarrow \mathbb{R}$

Initialize distribution parameters θ , **set** population size $\lambda \in \mathbb{N}$

While not terminate

- 1 **Sample** distribution $P(x|\theta) \rightarrow x_1, \dots, x_\lambda \in \mathbb{R}^n$
- 2 **Evaluate** $x_1, \dots, x_\lambda$ on f
- 3 **Update** parameters $\theta \leftarrow F_\theta(\theta, x_1, \dots, x_\lambda, f(x_1), \dots, f(x_\lambda))$

Everything depends on the definition of P and F_θ

deterministic algorithms are covered as well

In Evolutionary Algorithms the distribution P is often implicitly defined via **operators on a population**, in particular, selection, recombination and mutation

Natural template for *Estimation of Distribution Algorithms*

Evolution Strategies

New search points are sampled normally distributed

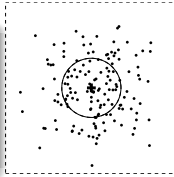
$$\mathbf{x}_i \sim \mathbf{m} + \sigma \mathcal{N}_i(\mathbf{0}, \mathbf{C}) \quad \text{for } i = 1, \dots, \lambda$$

as perturbations of $\mathbf{m}$, where $\mathbf{x}_i, \mathbf{m} \in \mathbb{R}^n$, $\sigma \in \mathbb{R}_+$, $\mathbf{C} \in \mathbb{R}^{n \times n}$
where

- the **mean** vector $\mathbf{m} \in \mathbb{R}^n$ represents the favorite solution
- the so-called **step-size** $\sigma \in \mathbb{R}_+$ controls the **step length**
- the **covariance matrix** $\mathbf{C} \in \mathbb{R}^{n \times n}$ determines the **shape** of the distribution ellipsoid

here, all new points are sampled with the same parameters

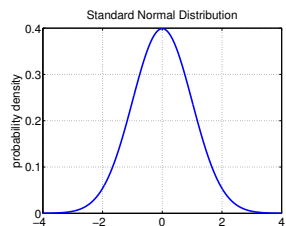
The question remains how to update $\mathbf{m}$, $\mathbf{C}$, and σ .



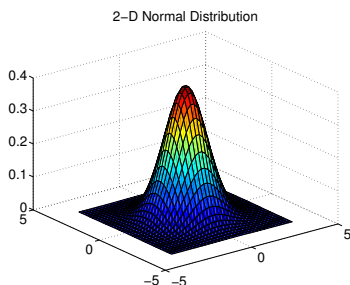
Why Normal Distributions?

- 1 widely observed in nature, for example as phenotypic traits
- 2 only stable distribution with finite variance
stable means the sum of normal variates is again normal, helpful in **design and analysis** of algorithms
- 3 most convenient way to generate **isotropic** search points
the isotropic distribution does **not favor any direction** (unfoundedly), supports rotational invariance
- 4 maximum entropy distribution with finite variance
the least possible assumptions on f in the distribution shape

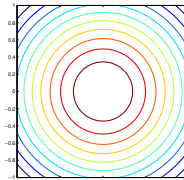
Normal Distribution



probability density of the 1-D standard normal distribution



probability density of a 2-D normal distribution



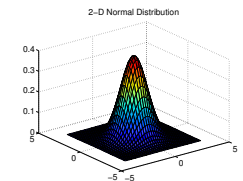
Why Normal Distributions?

The Multi-Variate (n -Dimensional) Normal Distribution

Any multi-variate normal distribution $\mathcal{N}(\mathbf{m}, \mathbf{C})$ is uniquely determined by its mean value $\mathbf{m} \in \mathbb{R}^n$ and its symmetric positive definite $n \times n$ covariance matrix $\mathbf{C}$.

The **mean** value $\mathbf{m}$

- determines the displacement (translation)
- value with the largest density (modal value)
- the distribution is symmetric about the distribution mean

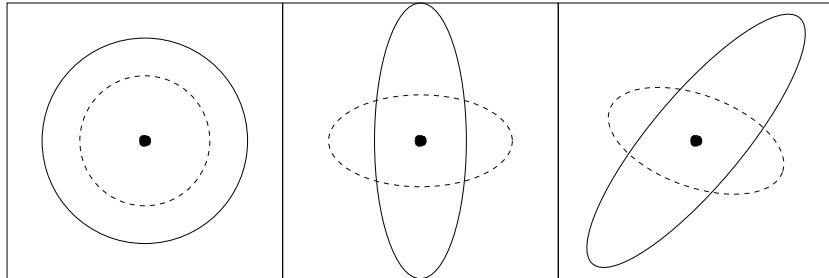


The **covariance matrix** $\mathbf{C}$

- determines the shape
- geometrical interpretation:** any covariance matrix can be uniquely identified with the iso-density ellipsoid $\{\mathbf{x} \in \mathbb{R}^n \mid (\mathbf{x} - \mathbf{m})^T \mathbf{C}^{-1} (\mathbf{x} - \mathbf{m}) = 1\}$

... any **covariance matrix** can be uniquely identified with the iso-density ellipsoid $\{x \in \mathbb{R}^n \mid x^T C^{-1} x = 1\}$

Lines of Equal Density



$\mathcal{N}(m, \sigma^2 \mathbf{I}) \sim m + \sigma \mathcal{N}(\mathbf{0}, \mathbf{I})$ **one degree of freedom** σ components are independent standard normally distributed
 $\mathcal{N}(m, \mathbf{D}^2) \sim m + \mathbf{D} \mathcal{N}(\mathbf{0}, \mathbf{I})$ **n degrees of freedom** components are independent, scaled
 $\mathcal{N}(m, \mathbf{C}) \sim m + \mathbf{C}^{\frac{1}{2}} \mathcal{N}(\mathbf{0}, \mathbf{I})$ **$(n^2 + n)/2$ degrees of freedom** components are correlated

where $\mathbf{I}$ is the identity matrix (isotropic case) and $\mathbf{D}$ is a diagonal matrix (reasonable for separable problems) and $\mathbf{A} \times \mathcal{N}(\mathbf{0}, \mathbf{I}) \sim \mathcal{N}(\mathbf{0}, \mathbf{A} \mathbf{A}^T)$ holds for all $\mathbf{A}$.

...ESs

Evolution Strategies

Terminology

Let μ : # parents, λ : # offspring

Plus (elitist) and comma (non-elitist) selection

$(\mu + \lambda)$ -ES: selection in $\{\text{parents}\} \cup \{\text{offspring}\}$

(μ, λ) -ES: selection in $\{\text{offspring}\}$

$(1 + 1)$ -ES

Sample one offspring from parent m

$$x = m + \sigma \mathcal{N}(\mathbf{0}, \mathbf{C})$$

If x better than m select

$$m \leftarrow x$$

...why?

The $(\mu/\mu, \lambda)$ -ES

Non-elitist selection and intermediate (weighted) recombination

Given the i -th solution point $x_i = m + \sigma \underbrace{\mathcal{N}_i(\mathbf{0}, \mathbf{C})}_{=: y_i} = m + \sigma y_i$

Let $x_{i:\lambda}$ the i -th **ranked** solution point, such that $f(x_{1:\lambda}) \leq \dots \leq f(x_{\lambda:\lambda})$.
The new mean reads

$$m \leftarrow \sum_{i=1}^{\mu} w_i x_{i:\lambda} = m + \sigma \underbrace{\sum_{i=1}^{\mu} w_i y_{i:\lambda}}_{=: y_w}$$

where

$$w_1 \geq \dots \geq w_{\mu} > 0, \quad \sum_{i=1}^{\mu} w_i = 1, \quad \frac{1}{\sum_{i=1}^{\mu} w_i^2} =: \mu_w \approx \frac{\lambda}{4}$$

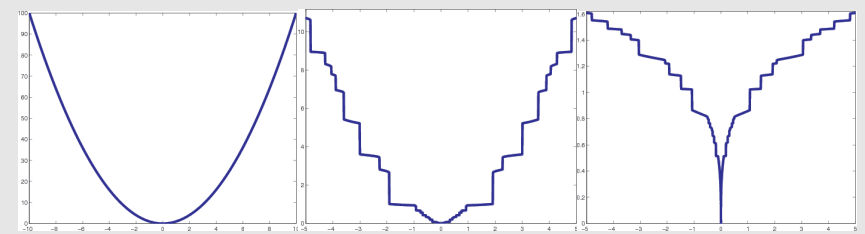
The best μ points are selected from the new solutions (non-elitistic) and **weighted intermediate recombination** is applied.

Invariance Under Monotonically Increasing Functions

Rank-based algorithms

Update of all parameters uses only the ranks

$$f(x_{1:\lambda}) \leq f(x_{2:\lambda}) \leq \dots \leq f(x_{\lambda:\lambda})$$



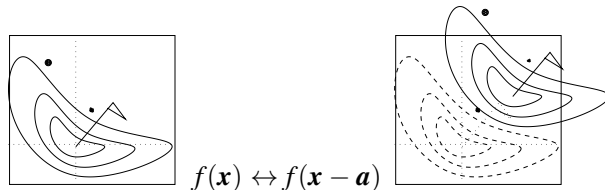
$$g(f(x_{1:\lambda})) \leq g(f(x_{2:\lambda})) \leq \dots \leq g(f(x_{\lambda:\lambda})) \quad \forall g$$

g is strictly monotonically increasing
 g preserves ranks

Basic Invariance in Search Space

- translation invariance

is true for most optimization algorithms



Identical behavior on f and f_a

$$\begin{aligned} f &: \mathbf{x} \mapsto f(\mathbf{x}), & \mathbf{x}^{(t=0)} &= \mathbf{x}_0 \\ f_a &: \mathbf{x} \mapsto f(\mathbf{x} - \mathbf{a}), & \mathbf{x}^{(t=0)} &= \mathbf{x}_0 + \mathbf{a} \end{aligned}$$

No difference can be observed w.r.t. the argument of f

Invariance

Impact

The grand aim of all science is to cover the greatest number of empirical facts by logical deduction from the smallest number of hypotheses or axioms.

— Albert Einstein

- empirical performance results, for example

- from benchmark functions
- from solved real world problems

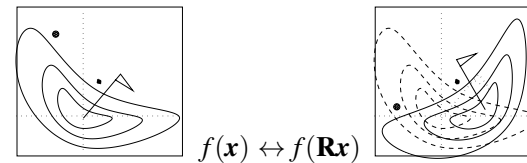
are only useful if they do **generalize** to other problems

- Invariance** is a strong **non-empirical** statement about the feasibility of generalization
generalizing (identical) performance from a single function to a whole class of functions

consequently, invariance is important for the evaluation of search algorithms

Rotational Invariance in Search Space

- invariance to an orthogonal transformation $\mathbf{R}$, where $\mathbf{R}\mathbf{R}^T = \mathbf{I}$
e.g. true for simple evolution strategies
recombination operators might jeopardize rotational invariance



Identical behavior on f and $f_{\mathbf{R}}$

$$\begin{aligned} f &: \mathbf{x} \mapsto f(\mathbf{x}), & \mathbf{x}^{(t=0)} &= \mathbf{x}_0 \\ f_{\mathbf{R}} &: \mathbf{x} \mapsto f(\mathbf{R}\mathbf{x}), & \mathbf{x}^{(t=0)} &= \mathbf{R}^{-1}(\mathbf{x}_0) \end{aligned}$$

No difference can be observed w.r.t. the argument of f

34

³Salomon (1996). "Reevaluating Genetic Algorithm Performance under Coordinate Rotation of Benchmark

- 1 Problem Statement
 - Black Box Optimization and Its Difficulties
 - Non-Separable Problems
 - Ill-Conditioned Problems
- 2 Evolution Strategies
 - A Search Template
 - The Normal Distribution
 - Invariance
- 3 Step-Size Control
 - Why Step-Size Control
 - One-Fifth Success Rule
 - Path Length Control (CSA)
- 4 Covariance Matrix Adaptation
 - Covariance Matrix Rank-One Update
 - Cumulation—the Evolution Path
 - Covariance Matrix Rank- μ Update
- 5 Theoretical Foundations
- 6 Experiments
- 7 Summary and Final Remarks

Evolution Strategies

Recalling

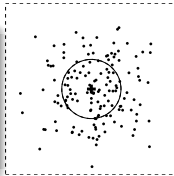
New search points are sampled normally distributed

$$\mathbf{x}_i \sim \mathbf{m} + \sigma \mathcal{N}(\mathbf{0}, \mathbf{C}) \quad \text{for } i = 1, \dots, \lambda$$

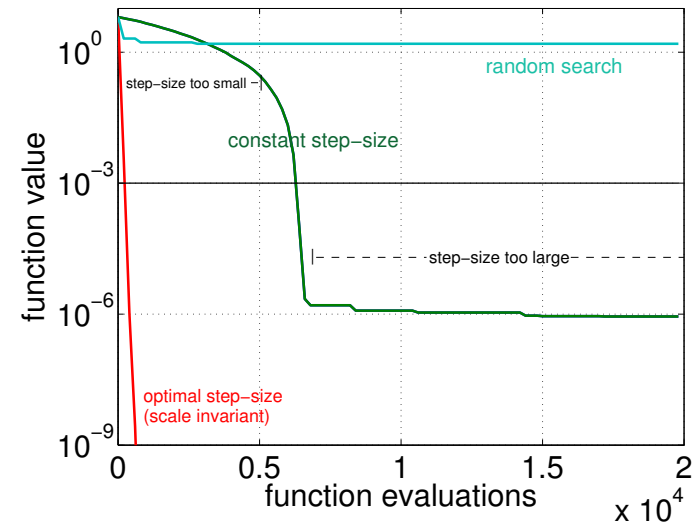
as perturbations of $\mathbf{m}$, where $\mathbf{x}_i, \mathbf{m} \in \mathbb{R}^n$, $\sigma \in \mathbb{R}_+$, $\mathbf{C} \in \mathbb{R}^{n \times n}$ where

- the **mean** vector $\mathbf{m} \in \mathbb{R}^n$ represents the favorite solution
- the so-called **step-size** $\sigma \in \mathbb{R}_+$ controls the *step length*
- the **covariance matrix** $\mathbf{C} \in \mathbb{R}^{n \times n}$ determines the **shape** of the distribution ellipsoid

The remaining question is how to update σ and $\mathbf{C}$.



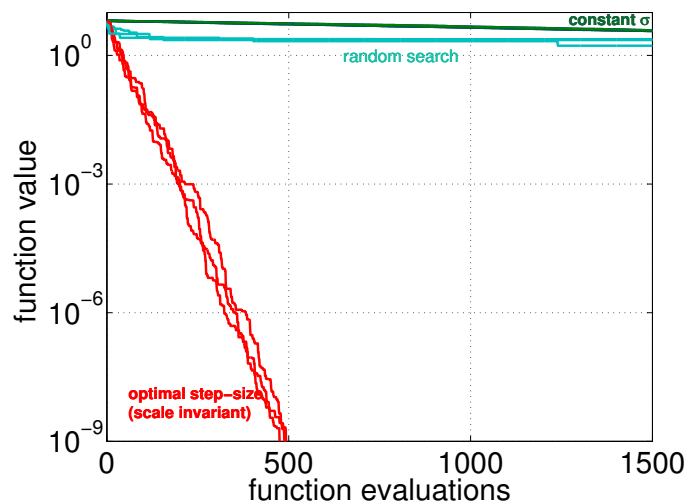
Why Step-Size Control?



$$f(\mathbf{x}) = \sum_{i=1}^n x_i^2$$

in $[-0.2, 0.8]^n$
for $n = 10$

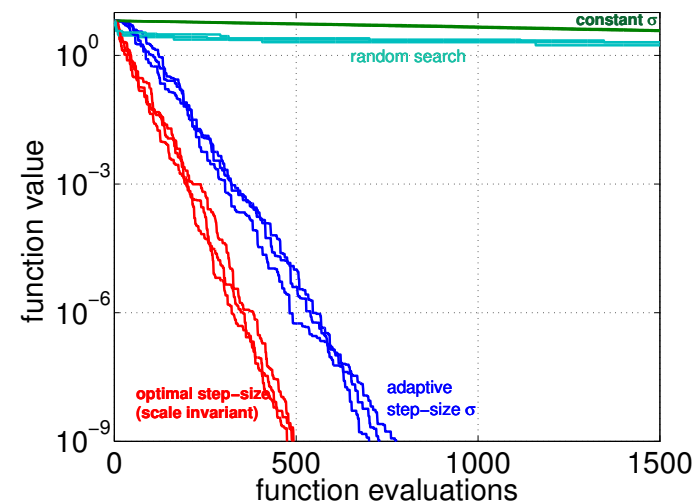
Why Step-Size Control?



$$f(\mathbf{x}) = \sum_{i=1}^n x_i^2$$

in $[-0.2, 0.8]^n$
for $n = 10$

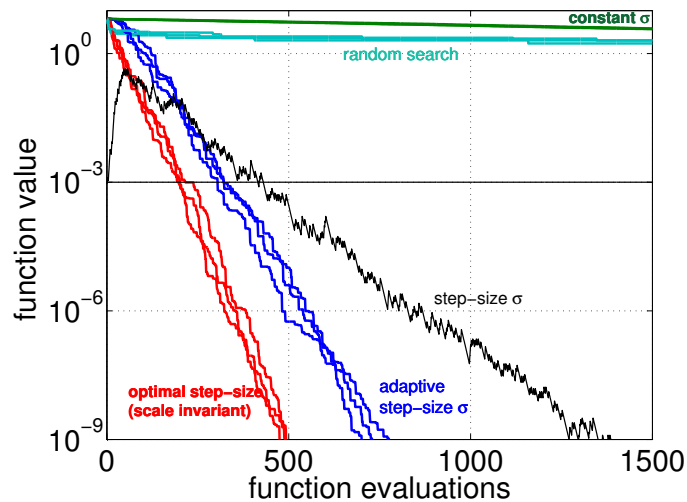
Why Step-Size Control?



$$f(\mathbf{x}) = \sum_{i=1}^n x_i^2$$

in $[-0.2, 0.8]^n$
for $n = 10$

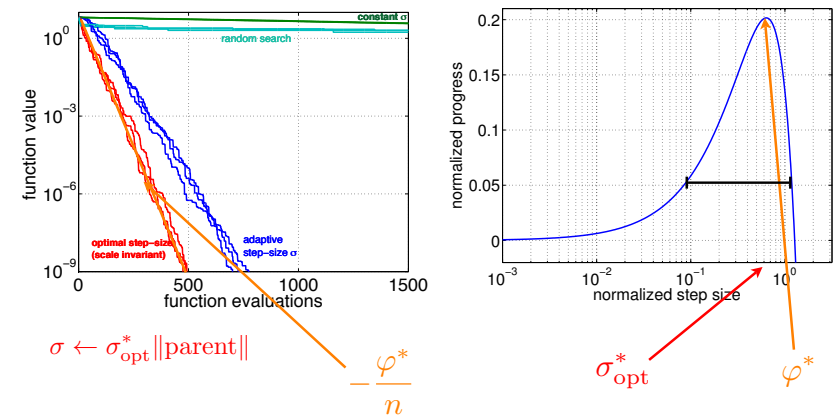
Why Step-Size Control?



$$f(\mathbf{x}) = \sum_{i=1}^n x_i^2$$

in $[-0.2, 0.8]^n$
for $n = 10$

Why Step-Size Control?



evolution window refers to the step-size interval (—) where reasonable performance is observed

Methods for Step-Size Control

- 1/5-th success rule^{ab}, often applied with “+”-selection
 - increase step-size if more than 20% of the new solutions are successful, decrease otherwise
- σ -self-adaptation^c, applied with “,”-selection
 - mutation is applied to the step-size and the better, according to the objective function value, is selected
 - simplified “global” self-adaptation
- path length control^d (Cumulative Step-size Adaptation, CSA)^e
 - self-adaptation derandomized and non-localized

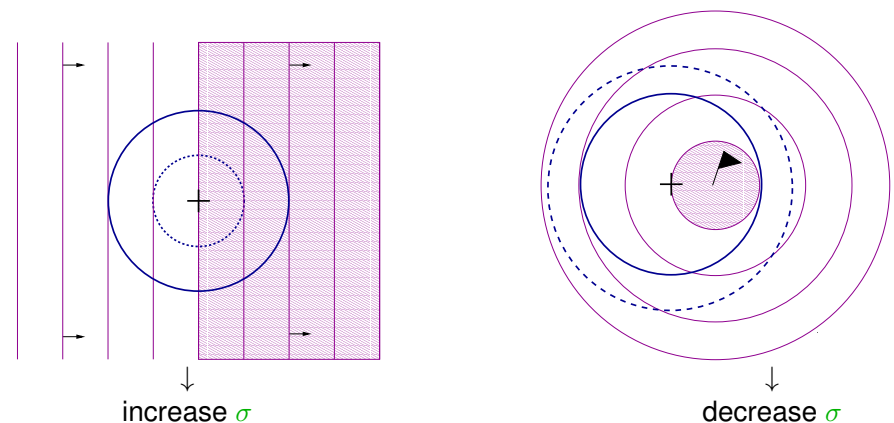
^aRechenberg 1973, *Evolutionstrategie, Optimierung technischer Systeme nach Prinzipien der biologischen Evolution*, Frommann-Holzboog

^bSchumer and Steiglitz 1968, Adaptive step size random search. *IEEE TAC*

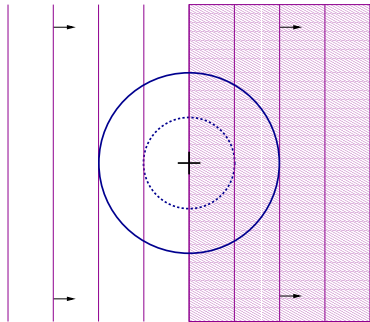
^cSchweifel 1981, *Numerical Optimization of Computer Models*, Wiley

^dHansen & Ostermeier 2001, Completely Derandomized Self-Adaptation in Evolution Strategies, *Evol. Comput.*

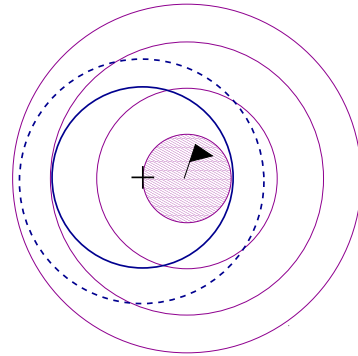
One-fifth success rule



One-fifth success rule

Probability of success (p_s)

1/2

Probability of success (p_s)

1/5

"too small"

One-fifth success rule

 p_s : # of successful offspring / # offspring (per generation)

$$\sigma \leftarrow \sigma \times \exp\left(\frac{1}{3} \times \frac{p_s - p_{\text{target}}}{1 - p_{\text{target}}}\right)$$

Increase σ if $p_s > p_{\text{target}}$
 Decrease σ if $p_s < p_{\text{target}}$

(1 + 1)-ES

$$p_{\text{target}} = 1/5$$

IF offspring better parent

$$p_s = 1, \sigma \leftarrow \sigma \times \exp(1/3)$$

ELSE

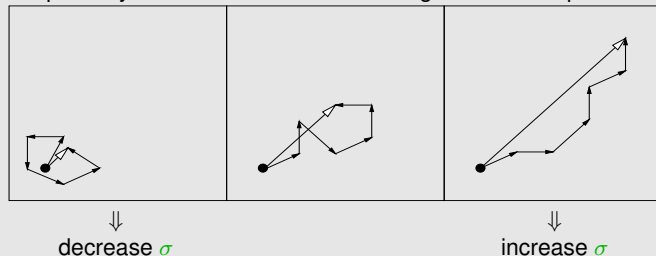
$$p_s = 0, \sigma \leftarrow \sigma / \exp(1/3)^{1/4}$$

Path Length Control (CSA)

The Concept of Cumulative Step-Size Adaptation

$$\begin{aligned} \mathbf{x}_i &= \mathbf{m} + \sigma \mathbf{y}_i \\ \mathbf{m} &\leftarrow \mathbf{m} + \sigma \mathbf{y}_w \end{aligned}$$

Measure the length of the evolution path

the pathway of the mean vector $\mathbf{m}$ in the generation sequence

loosely speaking steps are

- perpendicular under random selection (in expectation)
- perpendicular in the desired situation (to be most efficient)

Path Length Control (CSA)

The Equations

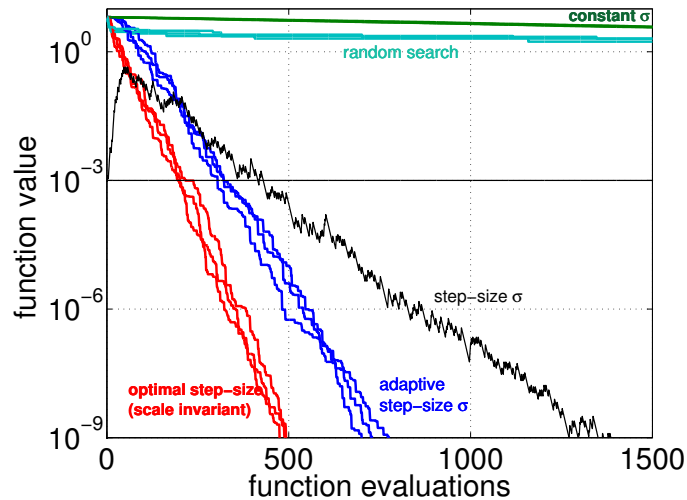
Initialize $\mathbf{m} \in \mathbb{R}^n$, $\sigma \in \mathbb{R}_+$, evolution path $\mathbf{p}_\sigma = \mathbf{0}$,
 set $c_\sigma \approx 4/n$, $d_\sigma \approx 1$.

$$\mathbf{m} \leftarrow \mathbf{m} + \sigma \mathbf{y}_w \quad \text{where } \mathbf{y}_w = \sum_{i=1}^{\mu} w_i \mathbf{y}_{i:\lambda} \quad \text{update mean}$$

$$\mathbf{p}_\sigma \leftarrow (1 - c_\sigma) \mathbf{p}_\sigma + \underbrace{\sqrt{1 - (1 - c_\sigma)^2}}_{\text{accounts for } 1 - c_\sigma} \underbrace{\sqrt{\mu_w}}_{\text{accounts for } w_i} \mathbf{y}_w$$

$$\sigma \leftarrow \sigma \times \exp\left(\frac{c_\sigma}{d_\sigma} \left(\frac{\|\mathbf{p}_\sigma\|}{\mathbb{E}\|\mathcal{N}(\mathbf{0}, \mathbf{I})\|} - 1\right)\right) \quad \text{update step-size}$$

$> 1 \iff \|\mathbf{p}_\sigma\|$ is greater than its expectation



$$f(\mathbf{x}) = \sum_{i=1}^n x_i^2$$

in $[-0.2, 0.8]^n$
for $n = 10$

- 1 Problem Statement
- 2 Evolution Strategies
- 3 Step-Size Control
- 4 Covariance Matrix Adaptation
 - Covariance Matrix Rank-One Update
 - Cumulation—the Evolution Path
 - Covariance Matrix Rank- μ Update
- 5 Theoretical Foundations
- 6 Experiments
- 7 Summary and Final Remarks

Evolution Strategies

Recalling

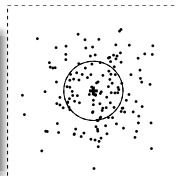
New search points are sampled normally distributed

$$\mathbf{x}_i \sim \mathbf{m} + \sigma \mathcal{N}_i(\mathbf{0}, \mathbf{C}) \quad \text{for } i = 1, \dots, \lambda$$

as perturbations of $\mathbf{m}$, where $\mathbf{x}_i, \mathbf{m} \in \mathbb{R}^n$, $\sigma \in \mathbb{R}_+$, $\mathbf{C} \in \mathbb{R}^{n \times n}$
where

- the **mean** vector $\mathbf{m} \in \mathbb{R}^n$ represents the favorite solution
- the so-called **step-size** $\sigma \in \mathbb{R}_+$ controls the *step length*
- the **covariance matrix** $\mathbf{C} \in \mathbb{R}^{n \times n}$ determines the **shape** of the distribution ellipsoid

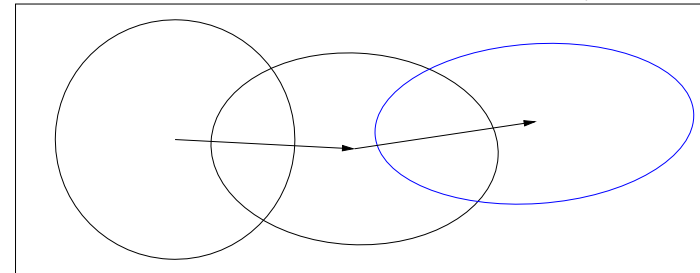
The remaining question is how to update $\mathbf{C}$.



Covariance Matrix Adaptation

Rank-One Update

$$\mathbf{m} \leftarrow \mathbf{m} + \sigma \mathbf{y}_w, \quad \mathbf{y}_w = \sum_{i=1}^{\mu} w_i \mathbf{y}_i, \quad \mathbf{y}_i \sim \mathcal{N}_i(\mathbf{0}, \mathbf{C})$$



new distribution,

$$\mathbf{C} \leftarrow 0.8 \times \mathbf{C} + 0.2 \times \mathbf{y}_w \mathbf{y}_w^T$$

the ruling principle: the adaptation **increases the likelihood of successful steps**, $\mathbf{y}_w$, to appear again
another viewpoint: the adaptation **follows a natural gradient**
approximation of the expected fitness

... equations

Covariance Matrix Adaptation

Rank-One Update

Initialize $\mathbf{m} \in \mathbb{R}^n$, and $\mathbf{C} = \mathbf{I}$, set $\sigma = 1$, learning rate $c_{\text{cov}} \approx 2/n^2$
While not terminate

$$\mathbf{x}_i = \mathbf{m} + \sigma \mathbf{y}_i, \quad \mathbf{y}_i \sim \mathcal{N}(\mathbf{0}, \mathbf{C}),$$

$$\mathbf{m} \leftarrow \mathbf{m} + \sigma \mathbf{y}_w \quad \text{where } \mathbf{y}_w = \sum_{i=1}^{\mu} w_i \mathbf{y}_{i:\lambda}$$

$$\mathbf{C} \leftarrow (1 - c_{\text{cov}})\mathbf{C} + c_{\text{cov}} \underbrace{\mu_w \mathbf{y}_w \mathbf{y}_w^T}_{\text{rank-one}} \quad \text{where } \mu_w = \frac{1}{\sum_{i=1}^{\mu} w_i^2} \geq 1$$

- 1 Problem Statement
- 2 Evolution Strategies
- 3 Step-Size Control
- 4 Covariance Matrix Adaptation
 - Covariance Matrix Rank-One Update
 - Cumulation—the Evolution Path
 - Covariance Matrix Rank- μ Update

5 Theoretical Foundations

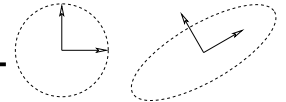
6 Experiments

7 Summary and Final Remarks

$$\mathbf{C} \leftarrow (1 - c_{\text{cov}})\mathbf{C} + c_{\text{cov}} \mu_w \mathbf{y}_w \mathbf{y}_w^T$$

covariance matrix adaptation

- learns all **pairwise dependencies** between variables
off-diagonal entries in the covariance matrix reflect the dependencies
- conducts a **principle component analysis** (PCA) of steps $\mathbf{y}_w$, sequentially in time and space
eigenvectors of the covariance matrix $\mathbf{C}$ are the principle components / the principle axes of the mutation ellipsoid, rotational invariant
- learns a new, **rotated problem representation** and a **new metric** (Mahalanobis)
components are independent (only) in the new representation rotational invariant
- approximates the **inverse Hessian** on quadratic functions
overwhelming empirical evidence, proof is in progress
- is **entirely independent** of the given coordinate system
algebraic formulation possible



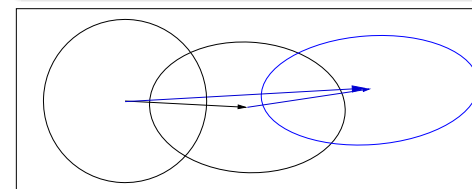
... cumulation, rank- μ

Cumulation

The Evolution Path

Evolution Path

Conceptually, the evolution path is the **search path** the strategy takes **over a number of generation steps**. It can be expressed as a sum of consecutive **steps** of the mean $\mathbf{m}$.



An exponentially weighted sum of steps $\mathbf{y}_w$ is used

$$\mathbf{p}_c \propto \sum_{i=0}^g \underbrace{(1 - c_c)^{g-i}}_{\text{exponentially fading weights}} \mathbf{y}_w^{(i)}$$

The recursive construction of the evolution path (cumulation):

$$\mathbf{p}_c \leftarrow \underbrace{(1 - c_c)}_{\text{decay factor}} \mathbf{p}_c + \underbrace{\sqrt{1 - (1 - c_c)^2}}_{\text{normalization factor}} \underbrace{\mu_w}_{\text{input} = \frac{\mathbf{m} - \mathbf{m}_{\text{old}}}{\sigma}} \mathbf{y}_w$$

where $\mu_w = \frac{1}{\sum w_i^2}$, $c_c \ll 1$. **History information** is accumulated in the evolution path.

“Cumulation” is a widely used technique and also known as

- *exponential smoothing* in time series, forecasting
- exponentially weighted *moving average*
- *iterate averaging* in stochastic approximation
- *momentum* in the back-propagation algorithm for ANNs
- ...

... why?

Using an **evolution path** for the **rank-one update** of the covariance matrix reduces the number of function evaluations to adapt to a straight ridge **from $\mathcal{O}(n^2)$ to $\mathcal{O}(n)$** .^(a)

^aHansen, Müller and Koumoutsakos 2003. Reducing the Time Complexity of the Derandomized Evolution Strategy with Covariance Matrix Adaptation (CMA-ES). *Evolutionary Computation*, 11(1), pp. 1-18

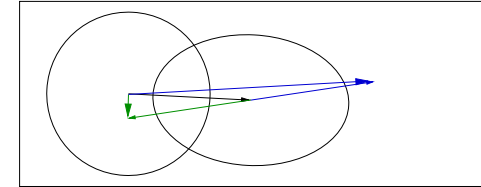
The overall model complexity is n^2 but important parts of the model can be learned in time of order n

... rank- μ update

Cumulation

Utilizing the Evolution Path

We used $\mathbf{y}_w \mathbf{y}_w^T$ for updating $\mathbf{C}$. Because $\mathbf{y}_w \mathbf{y}_w^T = -\mathbf{y}_w (-\mathbf{y}_w)^T$ the sign of $\mathbf{y}_w$ is lost.



The sign information is (re-)introduced by using the *evolution path*.

$$\begin{aligned} \mathbf{p}_c &\leftarrow \underbrace{(1 - c_c)}_{\text{decay factor}} \mathbf{p}_c + \underbrace{\sqrt{1 - (1 - c_c)^2} \sqrt{\mu_w}}_{\text{normalization factor}} \mathbf{y}_w \\ \mathbf{C} &\leftarrow (1 - c_{\text{cov}}) \mathbf{C} + c_{\text{cov}} \underbrace{\mathbf{p}_c \mathbf{p}_c^T}_{\text{rank-one}} \end{aligned}$$

where $\mu_w = \frac{1}{\sum w_i^2}$, $c_c \ll 1$.

Rank- μ Update

$$\begin{aligned} \mathbf{x}_i &= \mathbf{m} + \sigma \mathbf{y}_i, & \mathbf{y}_i &\sim \mathcal{N}_i(\mathbf{0}, \mathbf{C}), \\ \mathbf{m} &\leftarrow \mathbf{m} + \sigma \mathbf{y}_w, & \mathbf{y}_w &= \sum_{i=1}^{\mu} w_i \mathbf{y}_{i:\lambda} \end{aligned}$$

The rank- μ update extends the update rule for **large population sizes λ** using $\mu > 1$ vectors to update $\mathbf{C}$ at each generation step.

The matrix

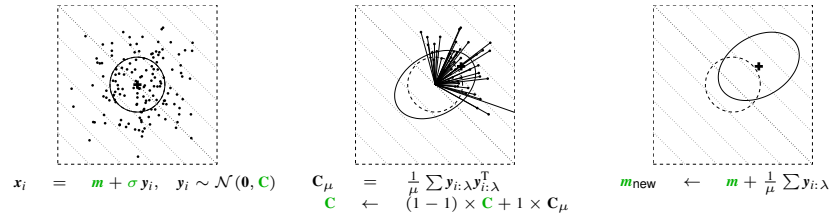
$$\mathbf{C}_\mu = \sum_{i=1}^{\mu} w_i \mathbf{y}_{i:\lambda} \mathbf{y}_{i:\lambda}^T$$

computes a weighted mean of the outer products of the best μ steps and has rank $\min(\mu, n)$ with probability one.

The rank- μ update then reads

$$\mathbf{C} \leftarrow (1 - c_{\text{cov}}) \mathbf{C} + c_{\text{cov}} \mathbf{C}_\mu$$

where $c_{\text{cov}} \approx \mu_w / n^2$ and $c_{\text{cov}} \leq 1$.

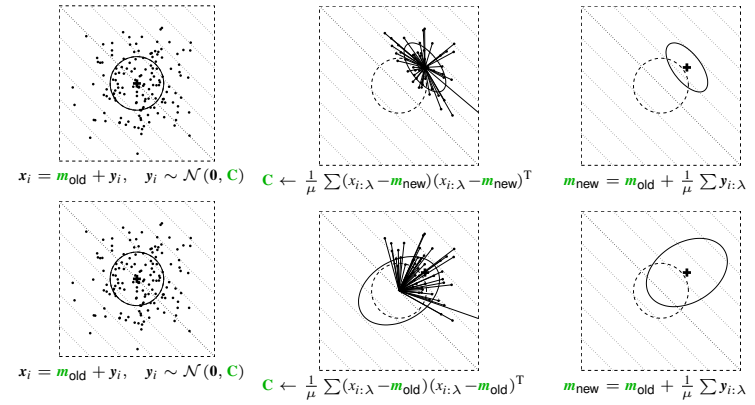


sampling of $\lambda = 150$
solutions where
 $C = I$ and $\sigma = 1$

calculating C where
 $\mu = 50$,
 $w_1 = \dots = w_\mu = \frac{1}{\mu}$,
and $c_{\text{cov}} = 1$

new distribution

Estimation of Multivariate Normal Algorithm EMNA_{global} versus rank- μ CMA⁵



EMNA_{global}
conducts a
PCA of
points

rank- μ CMA
conducts a
PCA of
steps

sampling of $\lambda = 150$
solutions (dots)

calculating C from $\mu = 50$
solutions

new distribution

The CMA-update yields a larger variance in particular in gradient direction, because m_{new} is the minimizer for the variances when calculating C

⁵ Hansen, N. (2006). The CMA Evolution Strategy: A Comparing Review. In J.A. Lozano, P. Larranga, I. Inza and E. Bengoetxea (Eds.). Towards a new evolutionary computation. Advances in estimation of distribution algorithms. pp. 75-102

The rank- μ update

- increases the possible learning rate in large populations
roughly from $2/n^2$ to μ_w/n^2
- can reduce the number of necessary **generations** roughly from $\mathcal{O}(n^2)$ to $\mathcal{O}(n)$ ⁽⁶⁾
given $\mu_w \propto \lambda \propto n$

Therefore the rank- μ update is the primary mechanism whenever a large population size is used

say $\lambda \geq 3n + 10$

The rank-one update

- uses the evolution path and reduces the number of necessary **function evaluations** to learn straight ridges from $\mathcal{O}(n^2)$ to $\mathcal{O}(n)$.

Rank-one update and rank- μ update can be combined

... all equations

⁶ Hansen, Müller, and Koumoutsakos 2003. Reducing the Time Complexity of the Derandomized Evolution Strategy with Covariance Matrix Adaptation (CMA-ES). *Evolutionary Computation*, 11(1), pp. 1-18

Summary of Equations

The Covariance Matrix Adaptation Evolution Strategy

Input: $m \in \mathbb{R}^n, \sigma \in \mathbb{R}_+, \lambda$

Initialize: $C = I$, and $p_c = 0, p_\sigma = 0$,

Set: $c_c \approx 4/n, c_\sigma \approx 4/n, c_1 \approx 2/n^2, c_\mu \approx \mu_w/n^2, c_1 + c_\mu \leq 1, d_\sigma \approx 1 + \sqrt{\frac{\mu_w}{n}}$,
and $w_{i=1 \dots \lambda}$ such that $\mu_w = \frac{1}{\sum_{i=1}^\lambda w_i^2} \approx 0.3 \lambda$

While not terminate

$x_i = m + \sigma y_i, \quad y_i \sim \mathcal{N}(0, C), \quad \text{for } i = 1, \dots, \lambda$ sampling

$m \leftarrow \sum_{i=1}^\mu w_i x_{i:\lambda} = m + \sigma y_w$ where $y_w = \sum_{i=1}^\mu w_i y_{i:\lambda}$ update mean

$p_c \leftarrow (1 - c_c) p_c + \mathbb{1}_{\{\|p_c\| < 1.5\sqrt{n}\}} \sqrt{1 - (1 - c_c)^2} \sqrt{\mu_w} y_w$ cumulation for C

$p_\sigma \leftarrow (1 - c_\sigma) p_\sigma + \sqrt{1 - (1 - c_\sigma)^2} \sqrt{\mu_w} C^{-\frac{1}{2}} y_w$ cumulation for σ

$C \leftarrow (1 - c_1 - c_\mu) C + c_1 p_c p_c^T + c_\mu \sum_{i=1}^\mu w_i y_{i:\lambda} y_{i:\lambda}^T$ update C

$\sigma \leftarrow \sigma \times \exp\left(\frac{c_\sigma}{d_\sigma} \left(\frac{\|p_\sigma\|}{\mathbb{E}\|\mathcal{N}(0, I)\|} - 1\right)\right)$ update of σ

Not covered on this slide: termination, restarts, useful output, boundaries and encoding

Source Code Snippet

```

counteval = 0; % the next 40 lines contain the 20 lines of interesting code
while counteval < stopeval

    % Generate and evaluate lambda offspring
    for k=1:lambda,
        arx(:,k) = xmean + sigma * B * (D .* randn(N,1)); % m + sig * Normal(0,C)
        arfitness(k) = feval(strfitnessfcn, arx(:,k)); % objective function call
        counteval = counteval+1;
    end

    % Sort by fitness and compute weighted mean into xmean
    [arfitness, arindex] = sort(arfitness); % minimization
    xold = xmean;
    xmean = arx(:,arindex(1:mu))*weights; % recombination, new mean value

    % Cumulation: Update evolution paths
    ps = (1-cs)*ps ...
        + sqrt(cs*(2-cs)*mueff) * invsqrtC * (xmean-xold) / sigma;
    hsig = norm(ps)/sqrt(1-(1-cs)^(2*counteval/lambda))/chiN < 1.4 + 2/(N+1);
    pc = (1-cs)*pc ...
        + hsig * sqrt(cc*(2-cc)*mueff) * (xmean-xold) / sigma;

    % Adapt covariance matrix C
    artmp = (1/sigma) * (arx(:,arindex(1:mu))- repmat(xold,1,mu));
    C = (1-cl-cmu) * C ... % regard old matrix
        + cl * (pc*pc') ... % plus rank one update
        + (1-hsig) * cc*(2-cc) * C ... % minor correction if hsig==0
        + cmu * artmp * diag(weights) * artmp'; % plus rank mu update

    % Adapt step size sigma
    sigma = sigma * exp((cs/damps)*(norm(ps)/chiN - 1));

    % Decomposition of C into B*diag(D.^2)*B' (diagonalization)
    if counteval - eigeneval > lambda*(cl*cmu)/N/10 % to achieve O(N^2)
        eigeneval = counteval;
        C = triu(C) + triu(C,1)'; % enforce symmetry
        [B,D] = eig(C); % eigen decomposition, B=normalized eigenvectors
        D = sqrt(diag(D)); % D is a vector of standard deviations now
        invsqrtC = B * diag(D.^-1) * B';
    end
end

```

Anne Auger & Nikolaus Hansen ()

CMA-ES

July, 2011 57 / 80

Theoretical Foundations

- 1 Problem Statement
- 2 Evolution Strategies
- 3 Step-Size Control
- 4 Covariance Matrix Adaptation
- 5 Theoretical Foundations
- 6 Experiments
- 7 Summary and Final Remarks

Anne Auger & Nikolaus Hansen ()

CMA-ES

July, 2011 59 / 80

Evolution Strategies in a Nutshell

- 1 **Sampling** from a multi-variate normal distribution
with maximum entropy
- 2 **Rank-based selection**: same performance on $g(f(x))$ for any g
 $g: \mathbb{R} \rightarrow \mathbb{R}$ strictly monotonic (order preserving)
- 3 **Step-size control**: converge log-linearly on the sphere function and many others
- 4 **Covariance matrix adaptation**: reduce any g^{-1} -convex quadratic function

$$f(x) = g(x^T H x)$$

to the sphere function

$$f(x) = x^T x$$

lines of equal density align with lines of equal fitness $C \propto H^{-1}$ without use of derivatives

... Theory

Anne Auger & Nikolaus Hansen ()

CMA-ES

July, 2011 58 / 80

Theoretical Foundations

Maximum Likelihood Update

The new distribution mean m maximizes the log-likelihood

$$m_{\text{new}} = \arg \max_m \sum_{i=1}^{\mu} w_i \log p_{\mathcal{N}}(x_{i:\lambda} | m)$$

independently of the given covariance matrix

The rank- μ update matrix C_μ maximizes the log-likelihood

$$C_\mu = \arg \max_C \sum_{i=1}^{\mu} w_i \log p_{\mathcal{N}} \left(\frac{x_{i:\lambda} - m_{\text{old}}}{\sigma} \middle| m_{\text{old}}, C \right)$$

$\log p_{\mathcal{N}}(x | m, C) = -\frac{1}{2} \log \det(2\pi C) - \frac{1}{2} (x - m)^T C^{-1} (x - m)$
 $p_{\mathcal{N}}$ is the density of the multi-variate normal distribution

Anne Auger & Nikolaus Hansen ()

CMA-ES

July, 2011 60 / 80

Natural Gradient Descent

- Consider $\arg \min_{\theta} E(f(\mathbf{x})|\theta)$ under the sampling distribution $p(\cdot|\theta)$
we could improve $E(f(\mathbf{x})|\theta)$ by following the gradient $\nabla_{\theta} E(f(\mathbf{x})|\theta)$:

$$\theta \leftarrow \theta - \eta \nabla_{\theta} E(f(\mathbf{x})|\theta), \quad \eta > 0$$

∇_{θ} depends on the parameterization of the distribution, specifically

- Consider the **natural gradient** of the expected weighted fitness

$$\begin{aligned} \tilde{\nabla} E(w(f(\mathbf{x})|\theta)) &= F_{\theta}^{-1} \nabla_{\theta} E(w(f(\mathbf{x})|\theta)) \\ &= E(w(f(\mathbf{x})) F_{\theta}^{-1} \nabla_{\theta} \ln p(\mathbf{x}|\theta)) \end{aligned}$$

using the Fisher information matrix $F_{\theta} = \left(E \left(\frac{\partial^2 \log p(\mathbf{x}|\theta)}{\partial \theta_i \partial \theta_j} \right) \right)_{ij}$ of the density p .

The natural gradient is **invariant under re-parameterization** of the distribution.

- A **Monte-Carlo approximation** reads

$$\tilde{\nabla} \hat{E}(\hat{w}(f(\mathbf{x})|\theta)) = \sum_{i=1}^{\lambda} w_i F_{\theta}^{-1} \nabla_{\theta} \ln p(\mathbf{x}_{i:\lambda}|\theta), \quad w_i = \hat{w}(f(\mathbf{x}_{i:\lambda})|\theta)$$

Variable Metric

On the function class

$$f(\mathbf{x}) = g \left(\frac{1}{2} (\mathbf{x} - \mathbf{x}^*)^T \mathbf{H} (\mathbf{x} - \mathbf{x}^*) \right)$$

the covariance matrix approximates the inverse Hessian up to a constant factor, that is:

$$\mathbf{C} \propto \mathbf{H}^{-1} \quad (\text{approximately})$$

In effect, ellipsoidal level-sets are transformed into spherical level-sets.

$g : \mathbb{R} \rightarrow \mathbb{R}$ is strictly increasing

CMA-ES = Natural Evolution Strategy + Cumulation

Natural gradient descent using the MC approximation and the normal distribution

- Rewriting the update of the distribution mean

$$\mathbf{m}_{\text{new}} \leftarrow \sum_{i=1}^{\mu} w_i \mathbf{x}_{i:\lambda} = \mathbf{m} - \underbrace{\sum_{i=1}^{\mu} w_i (\mathbf{m} - \mathbf{x}_{i:\lambda})}_{\text{natural gradient for mean } \frac{\partial}{\partial \mathbf{m}} \hat{E}(f(\mathbf{x})|\mathbf{m}, \mathbf{C})}$$

- Rewriting the update of the covariance matrix⁷

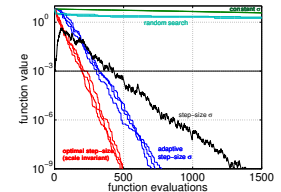
$$\begin{aligned} \mathbf{C}_{\text{new}} &\leftarrow \mathbf{C} + c_1 \underbrace{(\mathbf{p}_c \mathbf{p}_c^T)}_{\text{rank one}} - \mathbf{C} \\ &\quad - \frac{c_{\mu}}{\sigma^2} \sum_{i=1}^{\mu} w_i \left(\sigma^2 \mathbf{C} - \underbrace{(\mathbf{x}_{i:\lambda} - \mathbf{m})(\mathbf{x}_{i:\lambda} - \mathbf{m})^T}_{\text{rank-}\mu} \right) \\ &\quad \text{natural gradient for covariance matrix } \frac{\partial}{\partial \mathbf{C}} \hat{E}(f(\mathbf{x})|\mathbf{m}, \mathbf{C}) \end{aligned}$$

⁷ Akimoto et.al. (2010): Bidirectional Relation between CMA Evolution Strategies and Natural Evolution

On Convergence

Evolution Strategies converge with probability one on,
e.g., $g \left(\frac{1}{2} \mathbf{x}^T \mathbf{H} \mathbf{x} \right)$ like

$$\|\mathbf{m}_k - \mathbf{x}^*\| \propto e^{-ck}, \quad c \leq \frac{0.25}{n}$$



Monte Carlo pure random search converges like

$$\|\mathbf{m}_k - \mathbf{x}^*\| \propto k^{-c} = e^{-c \log k}, \quad c = \frac{1}{n}$$

- 1 Problem Statement
- 2 Evolution Strategies
- 3 Step-Size Control
- 4 Covariance Matrix Adaptation
- 5 Theoretical Foundations
- 6 Experiments
- 7 Summary and Final Remarks

Experimentum Crucis (0)

What did we want to achieve?

- reduce any convex-quadratic function

$$f(\mathbf{x}) = \mathbf{x}^T \mathbf{H} \mathbf{x}$$

$$\text{e.g. } f(\mathbf{x}) = \sum_{i=1}^n 10^{\frac{i-1}{n-1}} x_i^2$$

to the sphere model

$$f(\mathbf{x}) = \mathbf{x}^T \mathbf{x}$$

without use of derivatives

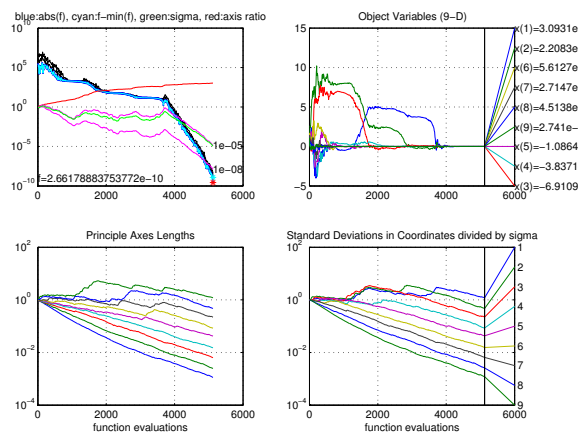
- lines of equal density align with lines of equal fitness

$$\mathbf{C} \propto \mathbf{H}^{-1}$$

in a stochastic sense

Experimentum Crucis (1)

f convex quadratic, separable

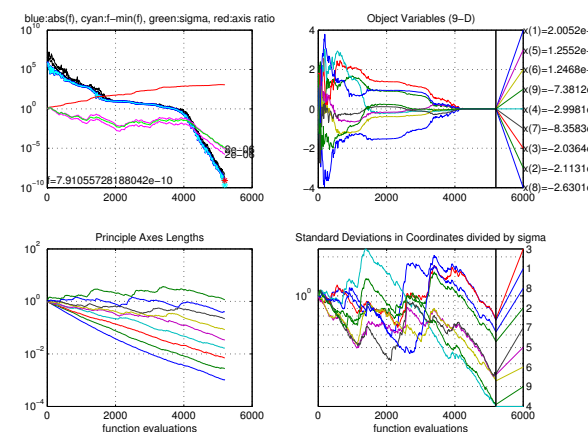


$$f(\mathbf{x}) = \sum_{i=1}^n 10^{\alpha \frac{i-1}{n-1}} x_i^2, \alpha = 6$$

... non-separable

Experimentum Crucis (2)

f convex quadratic, as before but non-separable (rotated)



$$f(\mathbf{x}) = g(\mathbf{x}^T \mathbf{H} \mathbf{x}), g: \mathbb{R} \rightarrow \mathbb{R} \text{ strictly increasing}$$

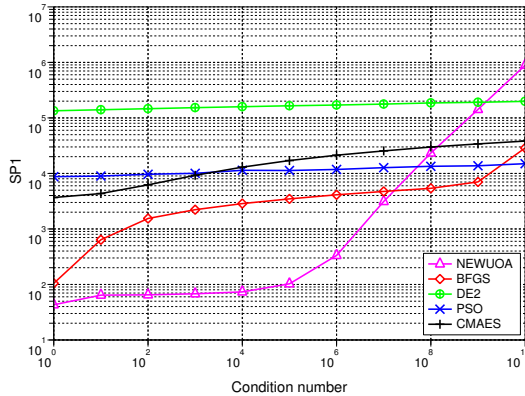
$$\mathbf{C} \propto \mathbf{H}^{-1} \text{ for all } g, \mathbf{H}$$

... internal parameters

Comparison to BFGS, NEWUOA, PSO and DE

f convex quadratic, separable with varying condition number α

Ellipsoid dimension 20, 21 trials, tolerance $1e-09$, eval max $1e+07$



BFGS (Broyden et al 1970)
NEWUOA (Powell 2004)
DE (Storn & Price 1996)
PSO (Kennedy & Eberhart 1995)
CMA-ES (Hansen & Ostermeier 2001)

$f(x) = g(x^T H x)$ with
 H diagonal
 g identity (for **BFGS** and **NEWUOA**)
 g any order-preserving = strictly increasing function (for all other)

SP1 = average number of objective function evaluations⁸ to reach the target function value of $g^{-1}(10^{-9})$

⁸ Auger et.al. (2009): Experimental comparisons of derivative free optimization algorithms, SEA

Anne Auger & Nikolaus Hansen ()

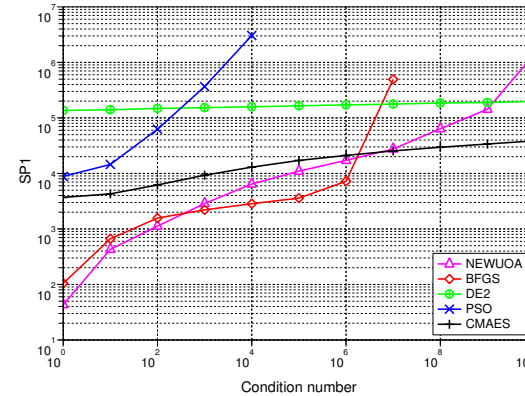
CMA-ES

July, 2011 69 / 80

Comparison to BFGS, NEWUOA, PSO and DE

f convex quadratic, non-separable (rotated) with varying condition number α

Rotated Ellipsoid dimension 20, 21 trials, tolerance $1e-09$, eval max $1e+07$



BFGS (Broyden et al 1970)
NEWUOA (Powell 2004)
DE (Storn & Price 1996)
PSO (Kennedy & Eberhart 1995)
CMA-ES (Hansen & Ostermeier 2001)

$f(x) = g(x^T H x)$ with
 H full
 g identity (for **BFGS** and **NEWUOA**)
 g any order-preserving = strictly increasing function (for all other)

SP1 = average number of objective function evaluations⁹ to reach the target function value of $g^{-1}(10^{-9})$

⁹ Auger et.al. (2009): Experimental comparisons of derivative free optimization algorithms, SEA

Anne Auger & Nikolaus Hansen ()

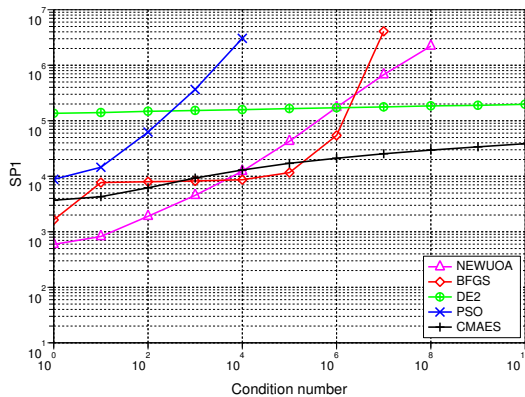
CMA-ES

July, 2011 70 / 80

Comparison to BFGS, NEWUOA, PSO and DE

f non-convex, non-separable (rotated) with varying condition number α

Sqrt of sqrt of rotated ellipsoid dimension 20, 21 trials, tolerance $1e-09$, eval max $1e+07$



BFGS (Broyden et al 1970)
NEWUOA (Powell 2004)
DE (Storn & Price 1996)
PSO (Kennedy & Eberhart 1995)
CMA-ES (Hansen & Ostermeier 2001)

$f(x) = g(x^T H x)$ with
 H full
 $g : x \mapsto x^{1/4}$ (for **BFGS** and **NEWUOA**)
 g any order-preserving = strictly increasing function (for all other)

SP1 = average number of objective function evaluations¹⁰ to reach the target function value of $g^{-1}(10^{-9})$

¹⁰ Auger et.al. (2009): Experimental comparisons of derivative free optimization algorithms, SEA

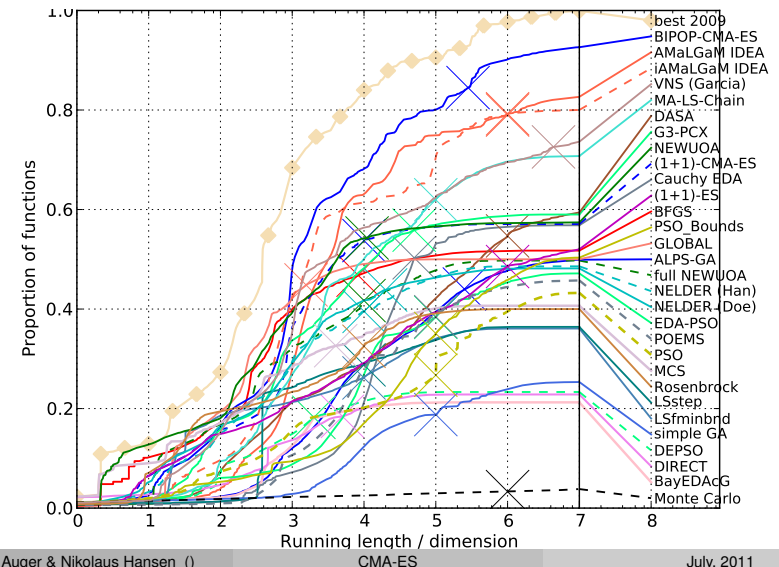
Anne Auger & Nikolaus Hansen ()

CMA-ES

July, 2011 71 / 80

Comparison during BBOB at GECCO 2009

24 functions and 31 algorithms in 20-D



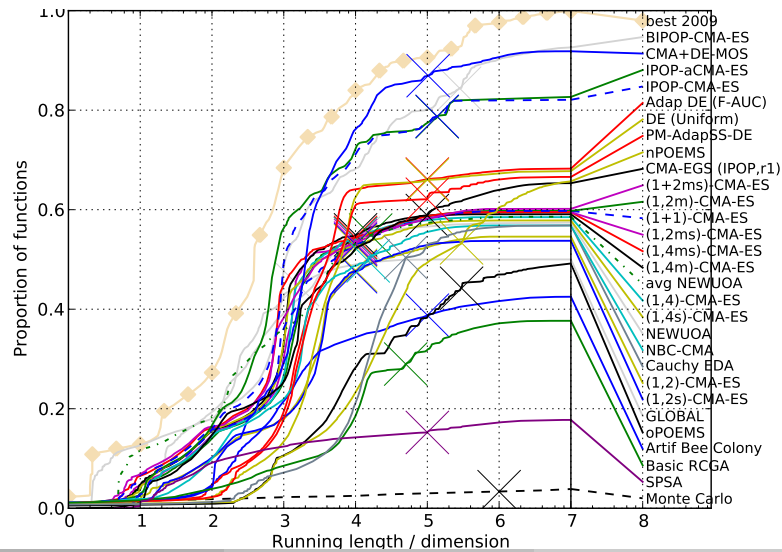
Anne Auger & Nikolaus Hansen ()

CMA-ES

July, 2011 72 / 80

Comparison during BBOB at GECCO 2010

24 functions and 20+ algorithms in 20-D



Anne Auger & Nikolaus Hansen ()

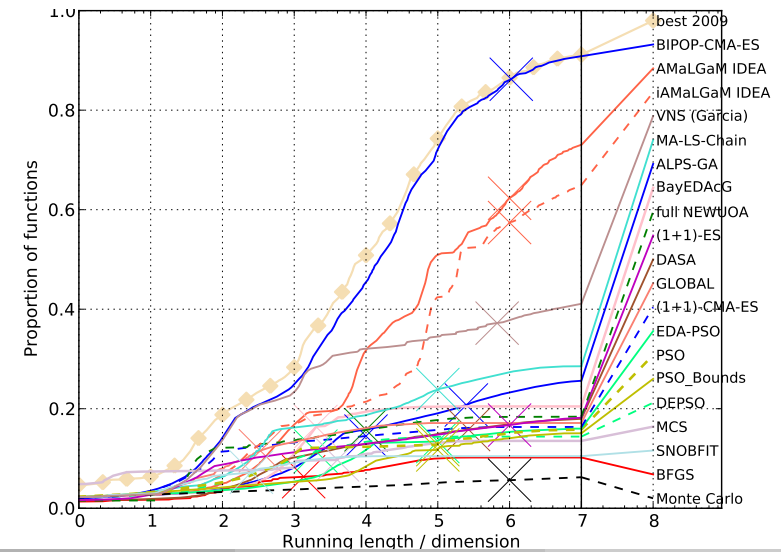
CMA-ES

July, 2011

73 / 80

Comparison during BBOB at GECCO 2009

30 **noisy** functions and 20 algorithms in 20-D



Anne Auger & Nikolaus Hansen ()

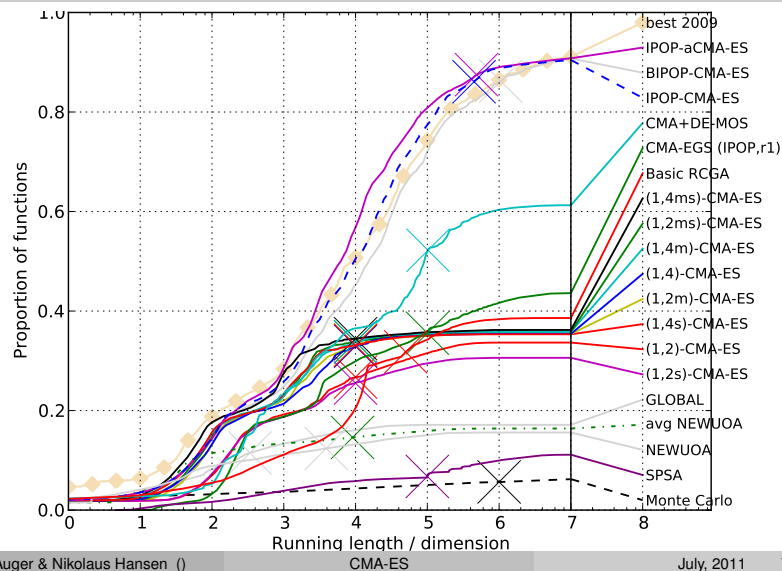
CMA-ES

July, 2011

74 / 80

Comparison during BBOB at GECCO 2010

30 **noisy** functions and 10+ algorithms in 20-D



Anne Auger & Nikolaus Hansen ()

CMA-ES

July, 2011

75 / 80

- 1 Problem Statement
- 2 Evolution Strategies
- 3 Step-Size Control
- 4 Covariance Matrix Adaptation
- 5 Theoretical Foundations
- 6 Experiments
- 7 Summary and Final Remarks

Anne Auger & Nikolaus Hansen ()

CMA-ES

July, 2011

76 / 80

The Continuous Search Problem

Difficulties of a non-linear optimization problem are

- dimensionality and non-separability
demands to exploit problem structure, e.g. neighborhood
- ill-conditioning
demands to acquire a second order model
- ruggedness
demands a non-local (stochastic?) approach

Approach: population based stochastic search, coordinate system independent and with second order estimations (covariances)

Limitations

of CMA Evolution Strategies

- **internal CPU-time:** $10^{-8}n^2$ seconds per function evaluation on a 2GHz PC, tweaks are available
100 000 f -evaluations in 1000-D take 1/4 hours internal CPU-time
- better methods are presumably available in case of
 - partly separable problems
 - specific problems, for example with cheap gradients
specific methods
 - small dimension ($n \ll 10$)
for example Nelder-Mead
 - small running times (number of f -evaluations $\ll 100n$)
model-based methods

Main Features of (CMA) Evolution Strategies

- 1 Multivariate normal distribution to generate new search points
follows the maximum entropy principle
- 2 Rank-based selection
implies invariance, same performance on $g(f(x))$ for any increasing g
more invariance properties are featured
- 3 Step-size control facilitates fast (log-linear) convergence
based on an evolution path (a non-local trajectory)
- 4 Covariance matrix adaptation (CMA) increases the likelihood of previously successful steps and can improve performance by orders of magnitude
the update follows the natural gradient
 $\mathbf{C} \propto \mathbf{H}^{-1} \iff$ adapts a variable metric
 $\iff$ new (rotated) problem representation
 $\implies f(\mathbf{x}) = g(\mathbf{x}^T \mathbf{H} \mathbf{x})$ reduces to $g(\mathbf{x}^T \mathbf{x})$

Source code for CMA-ES in C, Java, Matlab, Octave, Scilab, Python is available at

http://www.lri.fr/~hansen/cmaes_inmatlab.html