Experimental Analysis Descriptive Statistics Inferential Statistics Sequential Testing

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Experimental Analysis

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Sequential Testing

Course Overview

- 1. Combinatorial Optimization, Methods and Models
- 2. General overview
- 3. Solver System and Working Environment
- 4. Construction Heuristics
- 5. Local Search: Components, Basic Algorithms
- 6. Local Search: Neighborhoods and Search Landscape
- 7. Efficient Local Search: Incremental Updates and Neighborhood Pruning
- 8. Stochastic Local Search & Metaheuristics
- 9. Methods for the Analysis of Experimental Results
- 10. Configuration Tools: F-race
- 11. Very Large Scale Neighborhoods

Examples: GCP, CSP, TSP, SAT, MaxIndSet, SMTWP, Steiner Tree, p-median, set covering

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DM811 Heuristics for Combinatorial Optimization

Lecture 15 Methods for Experimental Analysis

Marco Chiarandini

Department of Mathematics & Computer Science University of Southern Denmark

Outline

1. Experimental Analysis

2. Descriptive Statistics Performance Measures

Sample Statistics Scenarios of Analysis Guidelines for Presenting Data

3. Inferential Statistics Statistical Tests Experimental Design

4. Race: Sequential Testing

Contents and Goals

Provide a view of issues in Experimental Algorithmics

- Exploratory data analysis
- Presenting results in a concise way with graphs and tables
- Organizational issues and Experimental Design
- Basics of inferential statistics
- Sequential statistical testing: race, a methodology for tuning

The goal of Experimental Algorithmics is not only producing a sound analysis but also adding an important tool to the development of a good solver for a given problem.

Experimental Algorithmics is an important part in the algorithm production cycle, which is referred to as Algorithm Engineering

The Engineering Cycle

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Experimental Algorithmics

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In empirical studies we consider simulation programs which are the implementation of a mathematical model (the algorithm)

[McGeoch, 1996]

from http://www.algorithm-engineering.de/

Experimental Algorithmics

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Fairness Principle

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Goals

- Defining standard methodologies
- Comparing relative performance of algorithms so as to identify the best ones for a given application
- Characterizing the behavior of algorithms
- Identifying algorithm separators, *i.e.*, families of problem instances for which the performance differ
- Providing new insights in algorithm design

Fairness principle: being completely fair is perhaps impossible but try to remove any possible bias

- possibly all algorithms must be implemented with the same style, with the same language and sharing common subprocedures and data structures
- the code must be optimized, e.g., using the best possible data structures
- running times must be comparable, e.g., by running experiments on the same computational environment (or redistributing them randomly)



realistic

Definitions

The most typical scenario considered in analysis of search heuristics

Asymptotic heuristics with time/quality limit decided a priori

The algorithm \mathcal{A}^∞ is halted when time expires or a solution of a given quality is found.

Deterministic case: \mathcal{A}^{∞} on π returns a solution of cost x.

The performance of \mathcal{A}^{∞} on π is a scalar y = x.

Randomized case: \mathcal{A}^{∞} on π returns a solution of cost X, where X is a random variable. The performance of \mathcal{A}^{∞} on π is the univariate Y = X.

[This is not the only relevant scenario: to be refined later]

Generalization

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For each general problem Π (e.g., TSP, GCP) we denote by C_{Π} a set

(or class) of instances and by $\pi \in C_{\Pi}$ a single instance.

On a specific instance, the random variable Y that defines the performance measure of an algorithm is described by its probability distribution/density function

$$Pr(Y = y \mid \pi)$$

It is often more interesting to generalize the performance on a class of instances C_{Π} , that is,

$$Pr(Y = y, C_{\Pi}) = \sum_{\pi \in \Pi} Pr(Y = y \mid \pi) Pr(\pi)$$

Random Variables and Probability

Statistics deals with random (or stochastic) variables.

A variable is called random if, prior to observation, its outcome cannot be predicted with certainty.

The uncertainty is described by a probability distribution.

Discrete variables

Probability distribution:

 $p_i = P[x = v_i]$

Cumulative Distribution Function (CDF)

 $F(v) = P[x \le v] = \sum_{i} p_i$

Mean

$$\mu = E[X] = \sum x_i p_i$$

Variance

$$\sigma^2 = E[(X - \mu)^2] = \sum (x_i - \mu)^2 p_i$$

Continuous variables

Probability density function (pdf):

$$f(v) = \frac{dF(v)}{dv}$$

Cumulative Distribution Function (CDF):

$$F(v) = \int_{-\infty}^{v} f(v) dv$$

Mean

$$\mu = E[X] = \int x f(x) dx$$

Variance

$$\sigma^2 = E[(X - \mu)^2] = \int (x - \mu)^2 f(x) \, dx$$

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Sampling

In experiments,

- 1. we sample the population of instances and
- 2. we sample the performance of the algorithm on each sampled instance

If on an instance π we run the algorithm r times then we have r replicates of the performance measure Y, denoted Y_1, \ldots, Y_r , which are independent and identically distributed (i.i.d.), i.e.

$$Pr(y_1,\ldots,y_r|\pi) = \prod_{j=1}^r Pr(y_j \mid \pi)$$

$$Pr(y_1,\ldots,y_r) = \sum_{\pi \in C_{\Pi}} Pr(y_1,\ldots,y_r \mid \pi) Pr(\pi).$$

Instance Selection

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In real-life applications a simulation of $p(\pi)$ can be obtained by historical data.

In simulation studies instances may be:

- real world instances
- random variants of real world-instances
- online libraries
- randomly generated instances

They may be grouped in classes according to some features whose impact may be worth studying:

- type (for features that might impact performance)
- size (for scaling studies)
- hardness (focus on hard instances)
- application (e.g., CSP encodings of scheduling problems), ...

Within the class, instances are drawn with uniform probability $p(\pi) = c$

Objectives of the Experiments

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• Comparison:

bigger/smaller, same/different, Algorithm Configuration, Component-Based Analysis

• Standard statistical methods: experimental designs, test hypothesis and estimation





Statistical Methods

The analysis of performance is based on finite-sized sampled data. Statistics provides the methods and the mathematical basis to

- describe, summarizing, the data (descriptive statistics)
- make inference on those data (inferential statistics)

Statistics helps to

- guarantee reproducibility
- make results reliable (are the observed results enough to justify the claims?)
- extract relevant results from large amount of data

In the practical context of heuristic design and implementation (i.e., engineering), statistics helps to take correct design decisions with the least amount of experimentation

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- Comparison:

bigger/smaller, same/different, Algorithm Configuration, Component-Based Analysis

• Standard statistical methods: experimental designs, test hypothesis and estimation

Objectives of the Experiments

• Characterization:

Interpolation: fitting models to data Extrapolation: building models of data, explaining phenomena

• Standard statistical methods: *linear* and non linear regression model fitting



Measures and Transformations

1. Experimental Analysis

2. Descriptive Statistics Performance Measures Sample Statistics

Scenarios of Analysis Guidelines for Presenting Data

3. Inferential Statistics

Statistical Tests Experimental Designs

4. Race: Sequential Testing

Measures and Transformations

On a single instance

Computational effort indicators

- number of elementary operations/algorithmic iterations

 (e.g., search steps, objective function evaluations, number of visited nodes in the search tree, consistency checks, etc.)
- total CPU time consumed by the process (sum of *user* and *system* times returned by getrusage)

Solution quality indicators

- value returned by the cost function
- error from optimum/reference value
- (optimality) gap $\frac{UB-LB}{LB+\epsilon}$ (if max $\frac{UB-LB}{UB+\epsilon}$) ϵ is an infinitesimal for the case LB = 0 but $UB - LB \neq 0$
- ranks

On a single instance

Design: Several runs on an instance

	Algorithm 1	Algorithm 2	 Algorithm k
Instance 1	X_{11}	X_{21}	X_{k1}
	-	:	:
Instance 1	X_{1r}	X_{2r}	X_{kr}

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Measures and Transformations

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On a class of instances

Design A: One run on various instances

	Algorithm 1	Algorithm 2	 Algorithm k
Instance 1	X_{11}	X_{12}	X_{1k}
1	-		
Instance b	X_{b1}	X_{b2}	X_{bk}

Design B: Several runs on various instances

	Algorithm 1	Algorithm 2	 Algorithm k
Instance 1	X_{111}, \ldots, X_{11r}	X_{121}, \ldots, X_{12r}	X_{1k1},\ldots,X_{1kr}
Instance 2	X_{211}, \ldots, X_{21r}	X_{221},\ldots,X_{22r}	X_{2k1},\ldots,X_{2kr}
:	:	:	-
Instance b	X_{b11},\ldots,X_{b1r}	X_{b21},\ldots,X_{b2r}	X_{bk1}, \ldots, X_{bkr}

Measures and Transformations

On a class of instances

Computational effort indicators

- $\bullet\,$ no transformation if the interest is in studying scaling
- standardization if a fixed time limit is used
- geometric mean (used for a set of numbers whose values are meant to be multiplied together or are exponential in nature),
- otherwise, better to group homogeneously the instances

Solution quality indicators

Different instances imply different scales \Rightarrow need for an invariant measure

(However, many other measures can be taken both on the algorithms and on the instances [McGeoch, 1996])

Sampling



 $\widehat{\theta}$

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Sequential Testing

We work with samples (instances, solution quality) drawn from populations

Population	Random Sample
$P(x,\theta)$ -	 X^n
Parameter θ	Statistical Estimator

Measures and Transformations

On a class of instances (cont.)

Solution quality indicators

• Distance or error from a reference value (assume minimization case):

$$e_1(x,\pi) = rac{x(\pi) - ar{x}(\pi)}{\sqrt{\sigma(\pi)}}$$
 standard score

$$e_2(x,\pi) = rac{x(\pi) - x^{opt}(\pi)}{x^{opt}(\pi)}$$
 relative error

$$e_3(x,\pi) = \frac{x(\pi) - x^{opt}(\pi)}{x^{worst}(\pi) - x^{opt}(\pi)} \quad \text{invariant [Zemel, 1981]}$$

- optimal value computed exactly or known by construction
- surrogate value such bounds or best known values
- Rank (no need for standardization but loss of information)

Summary Measures

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Measures to describe or characterize a population

- Measure of central tendency, location
- Measure of dispersion

One such a quantity is

- a parameter if it refers to the population (Greek letters)
- a **statistics** if it is an *estimation* of a population parameter from the sample (Latin letters)

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Measures of central tendency

• Arithmetic Average (Sample mean)

 $\bar{X} = \frac{\sum x_i}{n}$

- *Quantile*: value above or below which lie a fractional part of the data (used in nonparametric statistics)
 - Median

 $\mathcal{M} = x_{(n+1)/2}$

• Quartile

 $Q_1 = x_{(n+1)/4}$ $Q_3 = x_{3(n+1)/4}$

• *q*-quantile

 \boldsymbol{q} of data lies below and $1-\boldsymbol{q}$ lies above

Mode

value of relatively great concentration of data (*Unimodal vs Multimodal* distributions)

Measure of dispersion

• Sample range

$$R = x_{(n)} - x_{(1)}$$

• Sample variance

$$s^2 = \frac{1}{n-1} \sum (x_i - \bar{X})^2$$

• Standard deviation

$$s = \sqrt{s^2}$$

• Inter-quartile range

 $IQR = Q_3 - Q_1$

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Boxplot and a probability density function (pdf) of a Normal N(0,1s2) Population. (source: Wikipedia)

[see also: http://informationandvisualization.de/blog/box-plot]



Scenarios

 $> \times < -runif(10,0,1)$ mean(x), median(x), quantile(x), quantile(x,0.25)range(x), var(x), sd(x), IQR(x)> fivenum(x) #(minimum, lower-hinge, median, upper-hinge, maximum) [1] 0.18672 0.26682 0.28927 0.69359 0.92343 > summary(x) > aggregate(x,list(factors),median) > boxplot(x)

- A. Single-pass heuristics
- B. Asymptotic heuristics: Two approaches:
 - 1. Univariate
 - 1.a Time as an external parameter decided a priori
 - 1.b Solution quality as an external parameter decided a priori
 - 2. Cost dependent on running time:

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Scenario A

Single-pass heuristics

Deterministic case: \mathcal{A}^{+} on class C_{Π} **Randomized case:** \mathcal{A}^{+} on class C_{Π} returns a solution of cost x with computational effort t (e.g., running time).

The performance of \mathcal{A}^{+} on class C_{Π} is the vector $\vec{y} = (x, t)$.

returns a solution of cost X with computational effort T, where X and T are random variables.

The performance of \mathcal{A}^{-} on class C_{Π} is the bivariate $\vec{Y} = (X, T)$.

Example

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Scenario:

- \triangleright 3 heuristics \mathcal{A}_1^{\dashv} , \mathcal{A}_2^{\dashv} , \mathcal{A}_3^{\dashv} on class C_{Π} .
- ▷ homogeneous instances or need for data transformation.
- > 1 or r runs per instance
- Interest: inspecting solution cost and running time to observe and compare the level of approximation and the speed.

Tools:

• Scatter plots of solution-cost and run-time

Multi-Criteria Decision Making



Needed some definitions on dominance relations

In Pareto sense, for points in \mathbf{R}^2

$\vec{x}^1 \preceq \vec{x}^2$	weakly dominates	$x_i^1 \leq x_i^2$ for all $i = 1, \ldots, n$
$\vec{x}^1 \parallel \vec{x}^2$	incomparable	neither $\vec{x}^1 \preceq \vec{x}^2$ nor $\vec{x}^2 \preceq \vec{x}^1$

Scenario B

Asymptotic heuristics

There are two approaches:

1.a. Time as an external parameter decided *a priori*. The algorithm is halted when time expires.

Deterministic case: \mathcal{A}^{∞} on class C_{Π} returns a solution of cost x.

Randomized case: \mathcal{A}^{∞} on class C_{Π} returns a solution of cost X, where X is a random variable.

The performance of \mathcal{A}^{∞} on class C_{Π}

is the univariate Y = X.

The performance of \mathcal{A}^{∞} on class C_{Π} is the scalar y = x.

Example

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Scenario:

- > 3 heuristics A₁[∞], A₂[∞], A₃[∞] on class C_Π.
 (Or 3 heuristics A₁[∞], A₂[∞], A₃[∞] on class C_Π without interest in computation time because negligible or comparable)
- ▷ homogeneous instances (no data transformation) or heterogeneous (data transformation)
- \triangleright 1 or r runs per instance
- \triangleright a priori time limit imposed
- ► Interest: inspecting solution cost

Tools:

- Histograms (summary measures: mean or median or mode?)
- Boxplots
- Empirical cumulative distribution functions (ECDFs)

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On a class of instances



On a class of instances



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Stochastic Dominance

Definition: Algorithm A_1 probabilistically dominates algorithm A_2 on a problem instance, iff its CDF is always "below" that of A_2 , *i.e.*:

$F_1(x) \le F_2(x), \qquad \forall x \in X$



R code behind the previous plots

We load the data and plot the comparative boxplot for each instance.

> load("TS.class-G.dataR")
> G[1:5,]
alg inst run sol time.last.imp tot.iter parz.iter exit.iter exit.time opt
1 TS1 G-1000-0.5-30-1.1.col 1 59 9.900619 5955 442 5955 10.02463 30
2 TS1 G-1000-0.5-30-1.1.col 2 64 9.736608 3880 130 3958 10.00062 30
3 TS1 G-1000-0.5-30-1.1.col 3 64 9.908618 4877 49 4877 10.03263 30
4 TS1 G-1000-0.5-30-1.1.col 4 68 9.948622 6996 409 6996 10.07663 30
5 TS1 G-1000-0.5-30-1.1.col 5 63 9.912620 3986 52 3986 10.04063 30
> library(lattice)
> bwplot(alg ~ sol inst,data=G)

If we want to make an aggregate analysis we have the following choices:

- maintain the raw data,
- transform data in standard error,
- transform the data in relative error,
- transform the data in an invariant error,
- transform the data in ranks.

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Transform data in standard error

- Maintain the raw data
- > par(mfrow=c(3,2),las=1,font.main=1,mar=c(2,3,3,1))
- > #original data
- > **boxplot**(sol~alg,**data**=G,horizontal=TRUE,main="Original data")

- > #standard error > T1 <- split(G\$sol,list(G\$inst))</pre>
- > T2 <- lapply(T1,scale,center=TRUE,scale=TRUE)
- > T3 <- unsplit(T2, list(G\$inst))
- > T4 <- split(T3,list(G\$alg))
- > T5 < stack(T4)
- > boxplot(values~ind,data=T5,horizontal=TRUE,main=expression(paste("Standard error: ", frac(x-bar(x),sqrt(sigma)))))
- > library(latticeExtra)
- > ecdfplot(~values,group=ind,data=T5,main=expression(paste("Standard error: ",frac(x-bar(x),sqrt(sigma)))))

> #standard error

- > G\$scale < -0
- > split(G\$scale, G\$inst) <- lapply(split(G\$sol, G\$inst), scale,center=TRUE,scale=TRUE)

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Transform the data in relative error

> #relative error

- > G\$err2 <- (G\$sol-G\$opt)/G\$opt
- > boxplot(err2~alg,data=G,horizontal=TRUE,main=expression(paste("Relative error: ",frac(x -x^(opt),x^(opt)))))

Transform the data in an invariant error

We use as surrogate of x^{worst} the median solution returned by the simplest algorithm for the graph coloring, that is, the ROS heuristic.

> #error 3

- > load("ROS.class-G.dataR")
- > F1 <- aggregate(F\$sol,list(inst=F\$inst),median)</pre>
- > F2 <- split(F1\$x,list(F1\$inst))
- > G\$ref <- sapply(G\$inst,function(x) F2[[x]])</pre>
- > G\$err3 <- (G\$sol-G\$opt)/(G\$ref-G\$opt)

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Scenario B

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Transform the data in ranks

> #rank

- > G\$rank <- G\$sol
- > split(G\$rank, G\$inst) <- lapply(split(G\$sol, D\$inst), rank)
- > bwplot(rank~reorder(alg,rank,median),data=G,horizontal=TRUE,main="Ranks") > ecdfplot(rank,group=alg,data=G,main="Ranks")

Asymptotic heuristics

There are two approaches:

1.b. Solution quality as an external parameter decided a priori. The algorithm is halted when quality is reached.

Deterministic case: \mathcal{A}^{∞} on class	Randomized case: \mathcal{A}^{∞} on class C_{Π}
C_{Π} finds a solution in running time t .	finds a solution in running time T ,
	where T is a random variable.
The performance of \mathcal{A}^{∞} on class C_{Π}	
is the scalar $y = t$.	The performance of \mathcal{A}^{∞} on class C_{Π}
	is the univariate $Y = T$.

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Dealing with Censored Data

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Asymptotic heuristics, Approach 1.b

- \triangleright Heuristic \mathcal{A}^{\dashv} stopped before completion or \mathcal{A}^{∞} truncated (always the case)
- ▶ Interest: determining whether a prefixed goal (optimal/feasible) has been reached

The computational effort to attain the goal can be specified by a cumulative distribution function F(t) = P(T < t) with T in $[0, \infty)$.

If in a run i we stop the algorithm at time L_i then we have a Type I right censoring, that is, we know either

- T_i if $T_i < L_i$
- or $T_i > L_i$.

Hence, for each run *i* we need to record $\min(T_i, L_i)$ and the indicator variable for observed optimal/feasible solution attainment, $\delta_i = I(T_i \leq L_i)$.

Example

Asymptotic heuristics, Approach 1.b: Example

- \triangleright An exact vs an heuristic algorithm for the 2-edge-connectivity augmentation problem.
- ▶ Interest: time to find the optimum on different instances.



Uncensored:



Censored:



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Scenario B

Example

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Asymptotic heuristics

There are two approaches:

2. Cost dependent on running time:

Deterministic case: \mathcal{A}^{∞} on π returns a current best solution xat each observation in t_1, \ldots, t_k .

The performance of \mathcal{A}^{∞} on π is the profile indicated by the vector $\vec{y} = \{x(t_1), \dots, x(t_k)\}.$ **Randomized case:** \mathcal{A}^{∞} on π produces a monotone stochastic process in solution cost $X(\tau)$ with any element dependent on the predecessors.

The performance of \mathcal{A}^{∞} on π is the multivariate $\vec{Y} = (X(t_1), X(t_2), \dots, X(t_k)).$

Scenario:

- \triangleright 3 heuristics \mathcal{A}_1^{∞} , \mathcal{A}_2^{∞} , \mathcal{A}_3^{∞} on instance π .
- \triangleright single instance hence no data transformation.
- $\triangleright r$ runs
- Interest: inspecting solution cost over running time to determine whether the comparison varies over time intervals

Tools:

• Quality profiles

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The performance is described by multivariate random variables of the kind $\vec{Y} = \{Y(t_1), Y(t_2), \dots, Y(l_k)\}.$

Sampled data are of the form $\vec{Y}_i = \{Y_i(t_1), Y_i(t_2), \dots, Y_i(t_k)\}$, $i = 1, \dots, 10$ (10 runs per algorithm on one instance)



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Sampled data are of the form $\vec{Y}_i = \{Y_i(t_1), Y_i(t_2), \dots, Y_i(t_k)\}$, $i = 1, \dots, 10$ (10 runs per algorithm on one instance)



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Summary

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The performance is described by multivariate random variables of the kind $\vec{Y} = \{Y(t_1), Y(t_2), \dots, Y(l_k)\}.$

Sampled data are of the form $\vec{Y}_i = \{Y_i(t_1), Y_i(t_2), \dots, Y_i(t_k)\}$, $i = 1, \dots, 10$ (10 runs per algorithm on one instance)



The median behavior of the two algorithms

Making Plots

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http://algo2.iti.uni-karlsruhe.de/sanders/courses/bergen/bergenPresenting.pdf
[Sanders, 2002]

- Should the experimental setup from the exploratory phase be redesigned to increase conciseness or accuracy?
- What parameters should be varied? What variables should be measured?
- How are parameters chosen that cannot be varied?
- Can tables be converted into curves, bar charts, scatter plots or any other useful graphics?
- Should tables be added in an appendix?
- Should a 3D-plot be replaced by collections of 2D-curves?
- Can we reduce the number of curves to be displayed?
- How many figures are needed?
- Should the x-axis be transformed to magnify interesting subranges?

Visualize your data for your analysis and for communication to others

Explore your data:

- make plots: histograms, boxplots, empirical cumulative distribution functions, correlation/scatter plots
- look at the numerical data and interpret them in practical terms: computation times, distance from optimum
- look for patterns

All the above both at a single instance level and at an aggregate level.

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- Should the x-axis have a logarithmic scale? If so, do the x-values used for measuring have the same basis as the tick marks?
- Is the range of x-values adequate?
- Do we have measurements for the right x-values, i.e., nowhere too dense or too sparse?
- Should the y-axis be transformed to make the interesting part of the data more visible?
- Should the y-axis have a logarithmic scale?
- Is it misleading to start the y-range at the smallest measured value? (if not too much space wasted start from 0)
- Clip the range of y-values to exclude useless parts of curves?
- Can we use banking to 45° ?
- Are all curves sufficiently well separated?
- Can noise be reduced using more accurate measurements?
- Are error bars needed? If so, what should they indicate? Remember that measurement errors are usually not random variables.

- Connect points belonging to the same curve.
- Only use splines for connecting points if interpolation is sensible.
- Do not connect points belonging to unrelated problem instances.
- Use different point and line styles for different curves.
- Use the same styles for corresponding curves in different graphs.
- Place labels defining point and line styles in the right order and without concealing the curves.
- Give axis units
- Captions should make figures self contained.
- Give enough information to make experiments reproducible.
- Golden ratio rule: make the graph wider than higher [Tufte 1983].
- Rule of 7: show at most 7 curves (omit those clearly irrelevant).
- Avoid: explaining axes, connecting unrelated points by lines, cryptic abbreviations, microscopic lettering, pie charts

Inferential Statistics

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- We work with samples (instances, solution quality)
- But we want sound conclusions: generalization over a given population (all possible instances)
- Thus we need statistical inference



Since the analysis is based on finite-sized sampled data, statements like

"the cost of solutions returned by algorithm ${\cal A}$ is smaller than that of algorithm ${\cal B}''$

must be completed by

```
"at a level of significance of 5\%".
```

A Motivating Example

- There is a competition and two stochastic algorithms \mathcal{A}_1 and \mathcal{A}_2 are submitted.
- We run both algorithms once on n instances.
 On each instance either A₁ wins (+) or A₂ wins (-) or they make a tie (=).

Questions:

- 1. If we have only 10 instances and algorithm A_1 wins 7 times how confident are we in claiming that algorithm A_1 is the best?
- 2. How many instances and how many wins should we observe to gain a confidence of 95% that the algorithm A_1 is the best?

1. Experimental Analysis

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- 3. Inferential Statistics Statistical Tests Experimental Designs
- 4. Race: Sequential Testing

Experimental Analysis

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A Motivating Example

- p: probability that A_1 wins on each instance (+)
- *n*: number of runs without ties
- Y: number of wins of algorithm \mathcal{A}_1

If each run is independent and consitent:

$$Y \sim B(n,p): \qquad \Pr[Y=y] = \binom{n}{y} p^y (1-p)^{n-y}$$



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Sequential Testing

2 How many instances and how many wins should we observe to gain a confidence of 95% that the algorithm A_1 is the best?

To answer this question, we compute the 95% quantile, *i.e.*, $y: \Pr[Y \ge y] < 0.05$ with p = 0.5 at different values of n:

n	10	11	12	13	14	15	16	17	18	19	20
y	9	9	10	10	11	12	12	13	13	14	15

This is an application example of sign test, a special case of binomial test in which p=0.5

1 If we have only 10 instances and algorithm A_1 wins 7 times how confident are we in claiming that algorithm A_1 is the best?

Under these conditions, we can check how unlikely the situation is if it were $p(+) \leq p(-)$.

If p = 0.5 then the chance that algorithm A_1 wins 7 or more times out of 10 is 17.2%: quite high!



Statistical tests

Experimental Analysis Descriptive Statistics Inferential Statistics Sequential Testing

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General procedure:

- Assume that data are consistent with a null hypothesis H_0 (e.g., sample data are drawn from distributions with the same mean value).
- Use a statistical test to compute how likely this is to be true, given the data collected. This "likely" is quantified as the p-value.
- Do not reject H₀ if the p-value is larger than an user defined threshold called level of significance α.
- Alternatively, (p-value $< \alpha$), H_0 is rejected in favor of an alternative hypothesis, H_1 , at a level of significance of α .

Preparation of the Experiments

• If the sample size is large enough (infinity) any difference in the means

Study factors until the improvement in the response variable is deemed

Note: If resources available for N runs then the optimal design is one run on

of the factors, no matter how small, will be significant

• Desired statistical power + practical precision \Rightarrow sample size

Variance reduction techniquesSame pseudo random seed

• Real vs Statistical significance

Sample Sizes

small

Experimental Design

Algorithms \Rightarrow Treatment Factor;

Instances \Rightarrow Blocking/Random Factor

Design A: One run on various instances (Unreplicated Factorial)

	Algorithm 1	Algorithm 2	 Algorithm k
Instance 1	X_{11}	X_{12}	X_{1k}
-	-		:
Instance b	X_{b1}	X_{b2}	X_{bk}

Design B: Several runs on various instances (Replicated Factorial)

	Algorithm 1	Algorithm 2	 Algorithm k
Instance 1	X_{111}, \ldots, X_{11r}	X_{121}, \ldots, X_{12r}	X_{1k1}, \ldots, X_{1kr}
Instance 2	X_{211}, \ldots, X_{21r}	X_{221},\ldots,X_{22r}	X_{2k1},\ldots,X_{2kr}
	:	-	-
Instance b	X_{b11},\ldots,X_{b1r}	X_{b21},\ldots,X_{b2r}	X_{bk1}, \ldots, X_{bkr}

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Outline

Experimental Analysis Descriptive Statistics Inferential Statistics Sequential Testing

1. Experimental Analysis

N instances [Birattari, 2004]

- 2. Descriptive Statistics Performance Measures Sample Statistics Scenarios of Analysis
- 3. Inferential Statistics Statistical Tests Experimental Design

4. Race: Sequential Testing

Unreplicated Designs

Procedure Race [Birattari 2002]:

repeat

Randomly select an unseen instance and run all candidates on it

Perform all-pairwise comparison statistical tests

Drop all candidates that are significantly inferior to the best algorithm **until** only one candidate left or no more unseen instances ;

- F-Race use Friedman test
- Holm adjustment method is typically the most powerful

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Experimental Analysis

Descriptive Statistics

Inferential Statistics

Sequential Testing

Sequential Testing

Experimental Analysis Descriptive Statistics Inferential Statistics Sequential Testing

References

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