Optimization with R -Tips and Tricks

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Introduction

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Optimization

"optimization: an act, process, or methodology of making something (such as a design, system, or decision) as fully perfect, functional, or effective as possible; specifically: the mathematical procedures (such as finding the maximum of a function) involved in this."

- Merriam-Webster Online Dictionary, 2017 (*)

Forms of optimization (cf. Netspeak: "? optimization"):

- ► Code / program / system optimization
- ► Search / website / server . . . optimization
- ▶ Business / process / chain . . . optimization
- ► Engine / design / production optimization
- (*) First Known Use: 1857

Mathematical Optimization

A mathematical optimization problem consists of maximizing (or minimizing) a real objective function on a defined domain:

Given a set $A \subseteq R^n$ and a function $f: A \to R$ from A to the real numbers, find an element $x_0 \in A$ such that $f(x_0) \leq f(x)$ for all x in an *environment* of x_0 .

Typical problems:

- finding an optimum will be computationally expensive
- different types of objective functions and domains
- ▶ need to compute the optimum with very high accuracy
- ▶ need to find a global optimum, restricted resources
- etc.

Classification of Optimization Tasks

- Unconstrained optimization
- ▶ Nonlinear least-squares fitting (parameter estimation)
- ▶ Optimization with constraints
- ▶ Non-smooth optimization (e.g., minimax problems)
- ► Global optimization (stochastic programming)
- ► Linear and quadratic programming (LP, QP)
- ► Convex optimization (resp. SOCP, SDP)
- ► Mixed-integer programming (MIP, MILP, MINLP)
- ► Combinatorial optimization (e.g., graph problems)

Optimization in Statistics

- Maximum Likelihood
- ► Parameter estimation
- ► Quantile and density estimation
- ► LASSO estimation
- ► Robust regression
- ► Nonlinear equations
- ► Geometric programming problems
- ▶ Deep Learning / Support Vector Machines
- ► Engineering and Design, e.g. optimal control
- ▶ Operations Research, e.g. network flow problems
- ► Economics, e.g. portfolio optimization

100+ Packages on the Optimization TV

adagio alabama BB boot bvls cccp cec2005benchmark cec2013 CEoptim clpAPI CLSOCP clue cmaes cmaesr copulaedas cplexAPI crs dclone DEoptim DEoptimR desirability dfoptim ECOSolveR GA genalg GenSA globalOptTests glpkAPI goalprog GrassmannOptim gsl hydroPSO igraph irace isotone kernlab kofnGA lbfgs lbfgsb3 limSolve linprog localsolver LowRankQP lpSolve lpSolveAPI matchingMarkets matchingR maxLik mcga mco minpack.lm minqa neldermead NlcOptim nleqslv nlmrt nloptr nls2 NMOF nnls onls optimx optmatch parma powell pso psoptim qap quadprog quantreg rcdd RCEIM Rcgmin rCMA Rcplex RcppDE Rcsdp Rdsdp rgenoud Rglpk rLindo Rmalschains Rmosek rneos ROI Rsolnp Rsymphony Rvmmin scs smoof sna soma subplex tabuSearch trust trustOptim TSP ucminf

Goals for this Talk

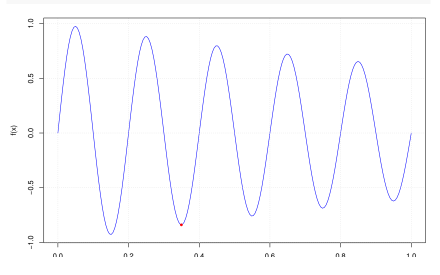
- Overview of (large, rapidly changing, still incomplete) set of tools for solving optimization problems in R
- ► Appreciation of the types of problems and types of methods to solve them
- Advice on setting up problems and solvers
- Suggestions for interpreting results
- ► Some almost real-world examples

Unfortunately, there is no time to talk about the new and exciting developments in *convex optimization* and optimization *modelling languages*.

Unconstrained Optimization

1-dimensional Example

```
f <- function(x) exp(-0.5*x) * sin(10*pi*x)
curve(f, 0, 1, n = 200, col=4); grid()
opt <- optimize(f, c(0, 1))
points(opt$minimum, opt$objective, pch = 20, col = 2)</pre>
```



Univariate (1-dim.) Minimization

```
optimize(f = , interval = , ..., lower = min(interval),
   upper = max(interval), maximum = FALSE,
   tol = .Machine$double.eps^0.25)

optim(par = , fn = , gr = NULL, ...,
   method = "Brent",
   lower = -Inf, upper = Inf)

optimizeR(f, lower, upper, ..., tol = 1e-20,
   method = c("Brent", "GoldenRatio"),
   maximum = FALSE,
   precFactor = 2.0, precBits = -log2(tol) * precFactor,
   maxiter = 1000, trace = FALSE)
```

optim() and Friends

Methods / Algorithms:

- ▶ Nelder-Mead downhill simplex method
- ▶ **BFGS** "variable metric" quasi-Newton method
- ▶ **CG** conjugate gradient method
- ► L-BFGS-B Broyden-Fletcher-Goldfarb-Shannon
- ▶ **Brent** univariate minimization, same as optimize
- ► **SANN** Simulated Annealing [don't use !]

Nelder-Mead

Nelder-Mead iteratively generates a sequence of simplices to approximate a minimal point.

At each iteration, the vertices of the simplex are ordered according to their objective function values and the simplex 'distorted' accordingly.

- ► **Sort** function values on simplex
- ▶ **Reflect** compute the reflection point
- **Expand** compute the expansion point
- ► Contract (outside | inside)
- ▶ **Shrink** the simplex

Stop when the simplex is small enough ('tolerance').

Showcase Rosenbrock

As a showcase we use the *Rosenbrock function*, defined for $n \ge 2$. It has has a very flat valley leading to its minimal point.

$$f(x_1,\ldots,x_n)=\sum_{i=2}^n[100(x_{i+1}-x_i^2)^2+(1-x_i)^2]$$

The global minimum obviously is (1, ..., 1) with value 0.

```
fnRosenbrock <- function (x) {
    n <- length(x)
    x1 <- x[2:n]; x2 <- x[1:(n - 1)]
    sum(100 * (x1 - x2^2)^2 + (1 - x2)^2)
}</pre>
```

Available in package adagio as fnRosenbrock(), with exact gradient grRosenbrock().

Nelder-Mead in Action

Nelder-Mead Simplex search over Banana Eunction

Figure 1: Source: de.wikipedia.org

optim() w/ Nelder-Mead

##

9707

NA

Nelder-Mead Solvers

dfoptim

adagio

pracma [new]

```
anms(fn, x0, ...,
    tol = 1e-10, maxfeval = NULL)
```

Gradient-Based Approaches

Exploiting the direction of "steepest descent" as computed by the negative gradient $-\nabla f(x)$ of a multivariate function.

- ► Steepest descent $d_k = -\nabla f(x_k)$
- ► Conjugate Gradient (GC)

$$d_k = -\nabla f(x_k) + \beta_k d_{k-1}$$
, $d_0 = -\nabla f(x_0)$, e.g., $\beta_k = \frac{||\nabla f(x_{k+1})||}{||\nabla f(x_k)||}$ (Fletcher and Reeves).

▶ **BFGS** and **L-BFGS-B** $d_k = -H_f(x_k)^{-1} \nabla f(x_k)$, $H_f(x)$ Hessian of f in x.

Adaptive Nelder-Mead

anms in *pracma* implements a new (Gao and Han, 2012) adaptive Nelder-Mead algorithm, adapting to the size of the problem (i.e., dimension of the objective function).

Line Searches

Given a function $f: R^n \to R$ and a direction $d \in R^n$, a line search method **approximately** minimizes f along the line $\{x + t \ d \mid t \in R\}$.

Armijo-Goldstein inequality: $0 < c, \nu < 1$

$$f(x_0 + t^*d) \le f(x_0) + c \nu^k f'(x_0; d), \quad k = 0, 1, 2, ...$$

(Weak) Wolf condition: $0 < c_1 < c_2 < 1$

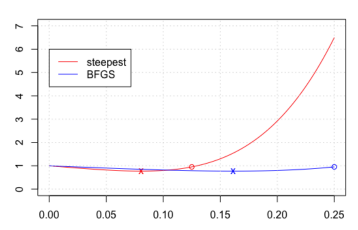
$$f(x_k + t_k d_k) \leq f(x_k) + c_1 t_k f'(x_k; d_k)$$

$$c_2f'(x_k;d_k) \leq f'(x_k + t_k d_k;g_k)$$

Rosenbrock with Line Search

Steepest descent direction vs. BFGS direction Wolfe line search these two directions

Wolfe line search



optim() w/ BFGS

BFGS and L-BFGS-B

The Broyden-Fletcher-Goldfarb-Shanno (BFGS) algorithm

Iteration: While $\|\nabla f_k\| > \epsilon$ do

- ▶ compute the search direction: $d_k = -H_k \nabla f_k$
- proceed with line search: $x_{k+1} = x_k + \alpha d_k$
- ▶ Update approximate Hessian inverse: $H_{k+1} \approx H_f(x_{k+1})^{-1}$

L-BFGS – low-memory BFGS stores matrix H_k in O(n) storage.

BFGS-B - BFGS with bound constraints ('active set' approach).

Best optim() usage

- use only method = "L-BFGS-B"
 (faster, more accurate, less memory, bound constraints)
- ▶ use factr = 1e-10 for tolerance, default 1e07
- set maxit = 50*d ... 50*d^2 (default is 100)
- use dfoptim or pracma for gradients (if you don't have an analytical or exact gradient)
- ► look carefully at the output

More BFGS Packages

▶ **Ibfgsb3** interfaces the Nocedal et al. 'L-BFGS-B.3.0' (2011) (FORTRAN) minimizer with bound constraints.

BUT: Options like "maximum number of function calls" are not accessible. (And the result is returned as 'invisible'.)

```
sol <- lbfgsb3(par, fn, gr = NULL, lower=-Inf, upper=Inf)
sol</pre>
```

▶ **Ibfgs** interfaces the 'libBFGS' C library by Okazaki with Wolfe line search (based on Nocedal).

BUT: Bound constraints are not accessible through the API.

```
lbfgs(fn, gr, par, invisible=1)
```

ucminf with Rosenbrock

```
fn <- adagio::fnRosenbrock; gr <- adagio::grRosenbrock</pre>
sol <- ucminf::ucminf(rep(0, 100), fn, gr, control=list(max
list(par=sol$par, value = sol$value, conv = sol$conv, mess
## $par
   ##
## $value
## [1] 1.223554e-15
##
## $conv
## [1] 1
##
## $mess
## [1] "Stopped by small gradient (grtol)."
```

More quasi-Newton type Algorithms

- stats::nlm [don't ever use!]
- stats::nlminb [PORT routine]

trustOptim::trust.optim [trust-region approach] no linesearch, suitable for sparse Hessians

▶ ucminf::ucminf [BFGS + line search + trust region]

```
ucminf(par, fn, gr = NULL, ..., control = list(), hessi
```

More John Nash Work

Thorough implementation of quasi-Newton solvers in pure R.

- ► Rcgmin ("conjugate gradient"")
- ► **Rvmmin** ("variable metric"")
- Rtnmin ("truncated Newton")

Apply, test, and compare different nonlinear optimization solvers for smooth, possibly bound constrained multivariate functions:

optimx, optimr, or optimrx?

Comparison of Nonlinear Solvers

```
method
                  value fevals gevals convd xtime
BFGS
            3.127628e-21
                            291
                                   98
                                          0 0.003
CG
           1.916095e-12
                          1107
                                   408
                                          0 0.010
Nelder-Mead 8.147198e+00
                          1501
                                   NA
                                          1 0.008
                                          0 0.001
L-BFGS-B
            5.124035e-10
                            78
                                   78
nlm
                                          0 0.002
           4.342036e-13
                            NA
                                   55
nlminb
           4.243607e-18
                           121
                                   97
                                          0 0.002
           5.124035e-10
                            78
                                   78
                                          0 0.029
lbfgsb3
                                          0 0.004
Rcgmin
           3.656125e-19
                            300
                                  136
Rtnmin
           5.403094e-13
                           105
                                   105
                                          0 0.013
           2.935561e-27
                           116
                                   72
                                          0 0.007
Rvmmin
ucminf
          1.470165e-15
                            77
                                   77
                                          0 0.002
                                          0 0.022
           3.614733e-11
                          1814
                                   NA
newuoa
                                          0 0.025
bobyqa
           6.939585e-10
                           2142
                                   NA
nmkb
           9.099242e-01
                          1500
                                   NA
                                          1 0.083
hjkb
           8.436900e-07
                           4920
                                   NA
                                          0 0.033
           9.962100e-13
lbfgs
                            NA
                                   NA
                                          0 0.001
```

Central-difference Formula

```
\nabla f(x) = (\frac{f(x)}{\partial x_1}, \dots, \frac{f(x)}{\partial x_n}) and \frac{df(x)}{dx}(x) \approx \frac{f(x+h) - f(x-h)}{2 \cdot h}
```

pracma::grad

```
function (f, x0, heps = .Machine$double.eps^(1/3), ...)
{
    # [...input checking...]
    n <- length(x0)
    hh <- rep(0, n)
    gr <- numeric(n)
    for (i in 1:n) {
        hh[i] <- heps
        gr[i] <- (f(x0 + hh) - f(x0 - hh))/(2 * heps)
        hh[i] <- 0
    }
    return(gr)
}</pre>
```

Excurse: Computing Gradients

- manually
- ► symbolically: package **Deriv**
- numerically: packages numDeriv or pracma

```
gr <- function(x) numDeriv::grad(fn, x) # simple, or:
gr <- function(x) pracma::grad(fn, x, heps=6e-06) # ce</pre>
```

complex step derivation

```
gr <- function(x) pracma::grad_csd(fn, x)</pre>
```

▶ automated differentiation [not yet available]

Optimization with Constraints

Constraints

- ▶ box/bound constraints: $I_i \le x_i \le u_i$ [trick: the 'transfinite' approach]
- ▶ linear inequality constraints: $Ax \le 0$
- ▶ linear equality constraints: Ax = b [trick: the 'hyperplane' approach]
- quadratic constraints
- ▶ inequality constraints in general
- equality and inequality constraints

Example: 'Transfinite' Approach

[1] 7.594813

Minimize the Rosenbrock function in 10 dimensions with $0 \le x_i \le 0.5$.

```
Tf <- adagio::transfinite(0, 0.5, 10)
h <- Tf$h; hinv <- Tf$hinv
p0 <- rep(0.25, 10)
f <- function(x) fn(hinv(x)) # f: R^n --> R
g <- function(x) pracma::grad(f, x)

sol <- lbfgs::lbfgs(f, g, p0, epsilon=1e-10, invisible=1)
hinv(sol$par); sol$value

## [1] 0.5000000000 0.2630659827 0.0800311137 0.0165742342
## [6] 0.0102120052 0.0102084108 0.0102042121 0.0100040850</pre>
```

The 'transfinite' Trick

If the solver does not support bound constraints $l_i \le x_i \le u_i$, the *transfinite* approach will do the trick.

Generate a smooth (surjective) function $h: \mathbb{R}^n \to [l_i, u_i]$, e.g.

$$h: x_i \rightarrow l_i + (u_i - l_i)/2 \cdot (1 + \tanh(x_i))$$

and optimize the composite function g(x) = f(h(x)), i.e.

$$g: R^n \to [I_i, u_i] \to R$$

$$x^* = \operatorname{argmin}_{x} g(x) = f(h(x))$$

then $x_{min} = h(x^*)$ will be a minimum of f in $[I_i, u_i]$.

Linear Inequality Constraints

Optimization with linear constraints only: $Ax \ge 0$ (or $Ax \le 0$)

```
constrOptim(theta, f, grad, ui, ci, mu = 1e-04, control
    method = if(is.null(grad)) "Nelder-Mead" el
    outer.iterations = 100, outer.eps = 1e-05,
    hessian = FALSE)
```

- ▶ ui %*% theta ci >= 0 corresponds to $Ax \ge 0$
- ▶ Bounds formulated as linear constraints (even $x_i \ge 0$)
- ▶ theta must be in the interior of the feasible region
- ▶ Inner iteration still calls optim

Recommendation: Do not use constrOptim. Instead, use an 'augmented Lagrangian' solver, e.g. alabama::auglag.

Trick: Linear Equality Constraints

Task: min!
$$f(x_1,...,x_n)$$
 s.t. $Ax = b$

Let $b_1, ..., b_m$ be a basis of the *nullspace* of A, i.e. $Ab_i = 0$, and x_0 a special solution $Ax_0 = b$. Define a new function $g(s_1, ..., s_m) = f(x_0 + s_1b_1 + ... + s_mb_m)$ and solve this as a minimization problem *without* constraints:

$$s = \operatorname{argmin} g(s_1, ..., s_m)$$

Then $xmin = x_0 + s_1b_1 + ... + s_mb_m$ is a (local) minimum.

Augmented Lagrangian Approach

Task: $\min f(x)$ s.t. $g_i(x) \ge 0, h_i(x) = 0$

Define the $augmented\ Lagrangian\ function\ L$ as

$$L(x, \lambda; \mu) = f(x) - \sum_{i} \lambda_{j} h_{j}(x) + \frac{1}{2\mu} \sum_{i} h_{j}^{2}(x)$$

The inequality constraints $g_i(x) \ge 0$ are included by introducing slack variables s_i and replacing the inequality constraints with

$$g_i(x)-s_i=0, \quad s_i\geq 0$$

The bound constraints are treated differently (e.g., through the LANCELOT algorithm).

Example: Linear Equality

```
A <- matrix(1, 1, 10)
                                       \# x1 + \ldots + xn = 1
N <- pracma::nullspace(A)</pre>
                                       # size 10 9
x0 \leftarrow qr.solve(A, 1)
                                       #Ax = 1
fun <- function(x) fn(x0 + N \% x) # length(x) = 9
sol <- ucminf::ucminf(rep(0, 9), fun)</pre>
xmin <- c(x0 + N \% *\% sol\$par)
xmin; sum(xmin)
    [1] 0.559312323 0.314864715 0.102103618 0.013695782
##
    [6] 0.003318010 0.003316801 0.003316309 0.003252102
## [1] 1
fn(xmin)
## [1] 7.421543
```

Augmented Lagrangian Solvers

alabama

► NLoptr

```
auglag(x0, fn, gr = NULL, lower = NULL, upper = NULL,
hin = NULL, hinjac = NULL, heq = NULL, heqjac =
localsolver = c("COBYLA"), localtol = 1e-6, ine
nl.info = FALSE, control = list(), ...)
```

- RsoInp
- ▶ **NIcOptim** (Sequential Quadratic programming, SQP)
- ▶ Rdonlp2 (removed from CRAN, see R-Forge's Rmetrics)

Example with alabama::auglag

```
Minimize the Rosenbrock function with constraints x_1 + \ldots + x_n = 1 and 0 \le x_i \le 1 for all i = 1, \ldots, n.

fheq <- function(x) sum(x) - 1
fhin <- function(x) c(x)

sol <- alabama::auglag(rep(0, 10), fn, gr, heq = fheq, hin control.outer = list(trace = FALSE, method = "Iprint(sol$par, digits=5)

## [1] 5.5707e-01 3.1236e-01 1.0052e-01 1.3367e-02 3.## [6] 3.3082e-03 3.3071e-03 3.3069e-03 3.2854e-03 -7.

sum(sol$par)
```

[1] 1

Quadratic Optimization

The *nloptr* Package (NLopt Library)

► COBYLA (Constrained Optimization By Linear Approximation)

slsqp (Sequential Quadratic Programming, SQP)

```
slsqp(x0, fn, gr = NULL, lower = NULL, upper = NULL,
hin = NULL, hinjac = NULL, heq = NULL, heqjac = NULL,
nl.info = FALSE, control = list(), ...)
```

auglag (Augmented Lagrangian)

```
auglag(x0, fn, gr = NULL, lower = NULL, upper = NULL,
hin = NULL, hinjac = NULL, heq = NULL, heqjac =
localsolver = c("COBYLA", "LBFGS", "MMA", "SLSC
nl.info = FALSE, control = list(), ...)
```

Quadratic Programming

Quadratic Programming (QP) is the problem of optimizing a quadratic expression of several variables subject to linear constraints.

Minimize
$$\frac{1}{2}x^{T}Qx + c^{T}x$$
s.t.
$$Ax < b$$

where Q is a symmetric, positive (semi-)definite $n \times n$ -matrix, c an n-dim. vector, A an $m \times n$ -matrix, and b an m-dim. vector.

For some solvers, linear equality constraints are also allowed.

Example: The enclosing ball problem

Quadratic Solvers

Standard solver for quadratic problems in R is solve.QP in package quadprog. The matrix Q has to be positive definite.

solve.QP(Dmat, dvec, Amat, bvec, meq=0, factorized=FALS

Package	Function	Matrix	Timings
quadprog	solve.QP	pdef	1
kernlab	ipop	spdef	50
LowRankQP	LowRankQP	spdef	2
DWD	solve_QP_SOCP	pdef	9500
coneproj	qprog	pdef	_

Nonsmoothness: Minimax Problems

Functions defined as maximum are not smooth and cannot be optimized through a straightforward gradient-based approach.

Task:
$$\min! f(x) = \max(f_1(x), \dots, f_m(x))$$

Instead, define a smooth function $g(x_1, ..., x_n, x_{n+1}) = x_{n+1}$ and minimze it under constraints

$$x_{n+1} \ge f_i(x_1,\ldots,x_n)$$
 for all $i=1,\ldots,m$

The solution $(x_1, \ldots, x_n, x_{n+1})$ returns the minimum point $xmin = (x_1, \ldots, x_n)$ as well as the minimal value $fmin = x_{n+1}$.

[Cf. the example in Chapter ?? in the bookdown text.]

Nonsmooth Optimization

Least Squares Solvers

Linear Least-squares

A linear least-squares (LS) problem means solving $\min \|Ax = b\|_2$, possibly with bounds or linear constraints.

The function qr.solve(A, b) from Base R solves over- and underdetermined linear systems in the least-squares sense.

- nnls (Lawson-Hansen algorithm) linear LS with non-negative/-positive constraints
- ▶ bvls (Stark-Parker algorithm) linear LS with bound constraints I < x < u</p>
- pracma::lsqlincon(A, b, ...)
 linear LS with linear equality and inequality constraints (applies
 a quadratic solver)

'Stabilized' Nonlinear LS

Modern nonlinear LS solvers use the Levenberg-Marquardt method (not Gauss-Newton) to minimize sums of squares.

minpack.lm

```
nlsLM(formula, data = parent.frame(), start, jac = NULI
    algorithm = "LM", control = nls.lm.control(),
    lower = NULL, upper = NULL, trace = FALSE, ...)
```

► nlmrt

```
nlxb(formula, start, trace=FALSE, data, lower=-Inf, upp
    masked=NULL, control, ...)
```

Cf. also **pracma**::lsqnonlin(fun, x0, options = list(), ...)

Nonlinear Least-squares

The standard nonlinear LS estimator for model parameter, given some data. in Base R is:

Problems:

- ▶ too small or zero residuals
- "singular gradient" error message (R-help, Stackoverflow)
- ▶ too many local minima, proper starting point (cf. nls2 with random or grid-based start points)
- bounds require the 'port' algorithm (Port library) (recommended anyway)

Tip: Rosenbrock as LS Problem

Redefine Rosenbrock as vector-value function:

```
fn <- function(x) {
    n <- length(x)
    x1 <- x[2:n]; x2 <- x[1:(n - 1)]
    c(10*(x1 - x2^2), 1 - x2)
}</pre>
```

and now apply **pracma**'s lsqnonlin:

Quantile Regression

```
Median (or: L^1) Regression: \min! \sum |y - Ax| (aka "least absolute deviation" (LAD) regression)
```

quantreg

```
rq(formula, tau = 0.5, data, subset, weights, na.action,
  method = "br", model = TRUE, contrasts, ...)
```

pracma

```
L1linreg(A, b, p = 1, tol = 1e-07, maxiter = 200)
```

solves the linear system Ax = b in an L^p sense, i.e. minimizes the term $\sum |b - Ax|^p$ (for 0) by applying an "iteratively reweighted least square" (IRLS) method.

DE Solvers

Differential Evolution (DE) is a relatively simple genetic algorithm variant, specialized for real-valued functions (10-20 dims).

▶ DEoptim

► RcppDE

```
DEoptim(fn, lower, upper, control = DEoptim.control(),
```

▶ DEoptimR

Global Optimization

CMA-ES Solvers

Covariance Matrix Adaptation – Evolution Strategy (CMA-ES) is an evolutionary algorithm for continuous optimization problems (adapting the covariance matrix). *It is quite difficult to implement*, but is applicable to dimensions up to 50 or more.

► Packages that contain CMA-ES solvers:

cmaes cmaesr

rCMA

parma::cmaes Rmalschains

adagio::pureCMAES

More Evolutionary Approaches

- Simulated Annealing (SA)GenSA
- Genetic Algorithms (GA)GA, genalg, SOMA, rgenoud
- Particle Swarm Optimization (PSO)pso, psoptim, hydroPSO

NMOF: DEopt, GAopt, PSopt

NLoptr: crs2lm, direct, mlsl, isres, stogo

Future Developments

The gloptim Package

Package **gloptim** incorporates and compares 25 stochastic solvers. The following is a typical output, here only showing the results of CMA-ES and DE solvers for the 'Runge' problem:

	solver	package	fmin	time
1	purecmaes	adagio	0.06546780	43.583
2	cmaes	parma	0.06546780	23.523
3	${\tt cmaoptim}$	\mathtt{rCMA}	0.06546780	91.257
4	malschains	Rmalschains	0.06546781	76.457
5	deopt	NMOF	0.06546876	75.809
6	deoptimr	${\tt DEoptimR}$	0.06549435	57.712
7	simplede	adagio	0.06573988	84.000
8	cma_es	cmaes	0.07430865	7.208
9	cmaes	cmaesr	0.07503498	8.305
22	cppdeoptim	RcppDE	6.82525344	17.050
23	deoptim	${ t DEoptim}$	7.28454226	39.287

ROI - R Optimization Infrastructure

Available Plugins:

glpk, symphony, quadprog, ipop, ecos, scs, nloptr, cplex, ...

CVXR

CVXR provides an R modeling language for convex optimization problems (announced UseR!2016, not yet ready).

Example: Estimating a discrete distribution, e.g.

```
max! \sum_{i=1}^{m} -w_i \log w_i

s.t. w_i \ge 0, \sum w_i = 1, X^T w = b

library(CVXR)

w <- Variable(m)

obj <- SumEntries(Entr(w)) # entropy function

constr <- list(w >= 0, SumEntries(w) == 1, t(X) %*% w == b)

pro <- Problem(Maximize(obj), constr)

sol <- solve(pro)

sol$w
```

Using the NEOS Solvers

"The **NEOS Server** https://neos-server.org/neos/ is a free internet-based service for solving numerical optimization problems. [It] provides access to more than 60 state-of-the-art [free and commercial] solvers."

rneos: XML-RPC Interface to NEOS

Using Julia Solvers

Ipopt (Interior Point OPTimizer) is a software package for large-scale nonlinear optimization (with nonlinear equality and inequality constraints).

- difficult to install (extra components needed)
- ► ECLIPSE license (not allowed on CRAN?)

There is an easy-to-install *Ipopt.jl* package for Julia.

With the R packages XR and XRJulia (John Chambers, 2016) it will be possible to utilize this with a new R package ipoptilr.

Epilogue

"What can go wrong?"

- ► Modell, constraints, gradients, . . .
- ► Local: bad starting values Global: no guaranteed optimum
- ► Applying appropriate solvers
- ► Setting solver controls
- ► Special problems, e.g.

 Non-smooth objective functions, noise, . . .
- Understanding solver output (and error messages)
 convergence, accuracy, no. of loops and function calls
- Checking results

References

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- ► Cortez, P. (2014). **Modern Optimization With R**. Use R! Series, Springer Intl. Publishing, Switzerland.

[&]quot;Most methods work most of the time." - John Nash