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Surrogates and (mono-objective) optimization: a long-term relationship

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SAMCO workshop, Lorentz Center
This talk is about: surrogates AND mono-objective optimization but not about: surrogates, optimization without surrogates

Pre-requisite: basics of optimization algorithms & surrogate modeling

Language elements:

• surrogates = metamodels = response surfaces = function approximation = proxy = emulator = specific surrogate name

• specific surrogate names: polynomial (e.g., quadratic ...) response surface, Gaussian process or kriging, artificial neural networks, radial basis functions, splines, support vector machines, high-dimensional model reduction (HDMR), generalized additive model (GAM), ...

Surrogates and (mono-objective) optimization have had a long-term relationship because most optimization methods can be seen as having a surrogate inside.
Mono-objective optimization problem formulation

\[
\min_{x \in S} f(x)
\]

\(x\), \(n\) optimization variables

\(S\), search space, \(x \in S\), mainly a compact \([x^{LB}, x^{UB}]\) in \(\mathbb{R}^n\)

but many concepts apply to \(\mathbb{N}^n\)

\(f\), objective or cost function to minimize, \(f : S \rightarrow \mathbb{R}\)
Optimizing expensive functions

Optimization algorithms generate points $x \in S$ in order to approximate the solution to $\min_{x \in S} f(x)$

$$\min_{x \in S} f(x)$$

but $M$ (e.g., computing sub-task) is typically computationally intensive

$$f(M(x^t))$$

data exchange, flow proportional to line thickness
**Idea 1: replace expensive functions by surrogates**

f is too expensive (to do optimization, or more generally computer experiments). Let's replace it with a cheaper metamodel or surrogate = a statistical model of the physical model $f(M(x))$.

$$f(M(x)) \equiv f(x)$$

$$\min_{x \in S} f(x) \approx \min_{x \in S} s(x)$$

The computing cost of solving this problem is considered negligible (compared to $M$).

How to build $s()$? To be partly discussed.
Optimization has relied on surrogates for a long time

E.g., Newton's method

Step according to a local model (surrogate) of the function (quadratic on this example, corresponding to Newton method,

$$\nabla^2 f(x^t)(x^{t+1} - x^t) = -\nabla f(x^t)$$

May fail: cf. trust region methods

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Idea 2: surrogate criterion

$f$ is too expensive (to do optimization, or more generally computer experiments). Let's replace the original problem (the function) with a problem that leads to the same solution. It includes idea 1.

\[ f(M(x)) \equiv f(x) \quad \text{c}(x) \text{ not necessarily } \approx f(x) \]

\[
\min_{x \in S} f(x) \quad \iff \quad \min_{x \in S} c(x, s(x))
\]

\[ \text{O} \quad \begin{array}{c}
\min_{x \in S} c(x, s(x)) \\
\end{array} \quad \begin{array}{c}
\chi^t \\
\text{many} \\
\text{many} \\
\text{criterion}
\end{array}
\]

Ok as long as the best of the iterates $x^t$ leads to $\arg\min_{x \in S} f(x)$.

The cost of calculating $c(.)$ is negligible w.r.t. $f(.)$. $c(.)$ often based on $s(.)$.
To be discussed: how to build $c(.)$?
A first (naive) algorithm

- Use a quadratic polynomial surrogate
  \[ s(x; \theta) = \theta_1 + \theta_1 x_1 + \ldots + \theta_{n+1} x_n + \theta_{n+2} x_1 x_2 + \ldots + \theta_{(n+1)(n+2)/2} x_n^2 \]
  \[ = \sum_{i=1}^{(n+1)(n+2)/2} \Phi_i(x) \theta_i = \Phi(x) \theta \] (linear in \( \theta \))
  \[ \Phi(x) = [1, x_1, \ldots, x_n, x_1x_2, \ldots, x_{n-1}x_n, x_1^2, \ldots, x_n^2] \]

- Create a "design of experiments" (DoE):
  E.g., \( t \geq (n+1)(n+2)/2 \) points randomly chosen in \( S \)
  \[ \Rightarrow X \equiv \{x^i\}, F \equiv \{f(x^i)\}, i = 1, t \]

- Fit the surrogate to the DoE by minimizing its "empirical risk" (sum of squares error)
  \[ \theta^* = \arg \min_{\theta} \sum_{i=1}^t (f(x^i) - s(x^i; \theta))^2 \] (closed form solution exist for linear models)
  \[ \equiv \arg \min_{\theta} E(\theta, X, F) \]

- Minimize the surrogate
  \[ x^{t+1} = \arg \min_{x \in S} s(x; \theta^*) \]
A first (naive) algorithm: 1D expl (1)

$x^{t+1}$ is not a minimizer of $f()$

not enough points?
$x^{t+1}$ is still not a minimizer of $f()$

$s(\theta)$ too rigid (does not have the right functional form), cannot learn $f()$?
A first (naive) algorithm: 1D expl (3)

Surrogate = cubic spline (a piece-wise 3rd degree polynomial with interpolation and smoothness properties)

Ok in 1 or 2D for a rough approximation (no convergence accuracy), but an a priori space filling DoE is very expensive. Expl: a grid has a geometrically growing number of evaluations, \( \text{step}^n \).

→ Need a more greedy strategy, putting new evaluation points in the good regions of \( S \) (where \( f \) is low).
A second (naive) algorithm

- Use a flexible surrogate (interpolating, or neural net with universal approximation property)
- Create an initial DoE, \((X, F)\), with not too many points (at most linear in \(n, t \approx 3n\))

- While ( \(t < \text{budget} \) ) do
  - Fit surrogate to current DoE
    \[
    \theta^* = \arg \min_{\theta} \text{Error}(\theta, X, F)
    \]
  - Minimize the surrogate
    \[
    x^{t+1} = \arg \min_{x \in S} s(x; \theta^*)
    \]
  - Calculate \(f\) & update DoE
    \[
    X = \{X \cup x^{t+1}\}, \; F = \{F \cup f(x^{t+1})\}
    \]
  - \(t = t+1\)

- End while
A second (naive) algorithm: 1D Expl

( cubic spline surrogate )

Converges to a local optimum, at best ...
A second (naive) algorithm

... because it can stall at non stationary points, when

\[ s(x^{t+1}, \theta^*) = f(x^{t+1}) \]
We have replaced the costly

\[
\min_{x \in S} f(x)
\]

by the less costly yet not converging to stationary points

\[
\min_{x \in S} s(x; \theta^t)
\]

\[
\min_{\theta} E(\theta, X, F)
\]
We miss a control that the surrogate S is leading the optimizer O towards better regions of the design space.

Minimizing the surrogate is not a good enough criterion in itself to ensure that the DoE created by the iterations allows convergence to local or global optima.
The type of strategies that ensure that the surrogate is not misleading will shape this presentation:

- Context and introduction
- Surrogates and trust regions for local optimization
  - quadratic surrogates
  - any surrogates
- Stochastic optimization using surrogates
- Surrogates with embedded error estimates: kriging
- Ensembles of surrogates
  - unstructured
  - structured
Quadratic surrogates and trust regions (1)

Basic idea

Build a quadratic surrogate and monitor its validity in a ball around the current iterate. Define iterates by solving a minimization problem in the ball.

Motivations

- In high dimensions (say > 100), it may not be possible to learn flexible surrogates because of the needed number of points.
- Quadratic surrogates are rigid but may always approximate a twice differentiable function in a neighborhood (order 2 Taylor).
- The minimum of quadratic surrogates is analytically tractable.

Conn et al., *Introduction to derivative free optimization*, SIAM Publ., 2009.
Quadratic surrogates and trust regions  (2)

► Create an initial DoE \((X, F)\) of \(m\) points, \(n+2 \leq m \leq (n+1)(n+2)/2\) ...

► While (not stop) do
   ● Fit quadratic surrogate \(s()\) to current DoE
      \[ \theta^t = \arg \min_{\theta} \| \nabla_x^2 s(x; \theta) - \nabla_x^2 s(x; \theta^{t-1}) \|_F \] s.t. \(s(x^i; \theta) = f(x^i), \ i = 1, m\)
   ● Minimize the surrogate within the trust region
      \[ x' = \arg \min_{x \in S} s(x; \theta^t) \text{ such that } \|x - x^t\| \leq \Delta_t \]
   ● Calculate \(f\) & check validity of surrogate
      \[ \rho = \frac{(f(x^t) - f(x'))/\|s(x^t; \theta^t) - s(x'; \theta^t)\|}{s(x^t; \theta^t) - s(x'; \theta^t)} \]
   ● Update trust region radius, current iterate and DoE
      \[ \text{If } (\rho \geq \mu > 0) \{ \uparrow \Delta_t, \ x^{t+1} = x' \} \text{ else } \{ \downarrow \Delta_t, \ x^{t+1} = x^t \} \]
      Add \(x'\) and remove a point from \((X, F)\) depending on dist. to \(x^{t+1}\) and identifiability of \(\theta\), \(t = t + 1\)

► End while

* M.J.D. Powell, The BOBYQA algorithm for bound constrained optimization without derivatives, TR Cambridge, 2009
Surrogate usefulness controlled through trust region

The regularization scheme (minimization of Hessian distances) makes \( (n+1)(n+2)/2 \) points to determine the parameters of the quadratic surrogate not necessary and allows an \( O(n) \) optimization cost.

Identifiability of \( \theta \) : conditioning of the linear system that comes from 1st order optimality conditions of the surrogate fitting sub-problem.

Because the surrogate is local, trust region methods are local optimization methods.

The BOBYQA algorithm is a state-of-the-art derivative free method for bound constrained minimization. It has been tested up to dimension \( n=320 \).
Optimization with any surrogate converges to a stationary point if a trust region strategy is used and the gradient of the true function is fitted at data points.


Outline of the talk

• Context and introduction

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O
- (rank x's according to their f's)
- update internal state $\alpha^t$ of optimizer
- sample $\lambda$ new x's according to a pdf $d(x;\alpha^t)$

M
\{x^t, \ldots, x^{t+\lambda}\}

For expl., in CMA-ES $\alpha^t \equiv (m^t, C^t)$ where $d(x;\alpha^t)$ is $N(m^t, C^t)$

Two main implementations of surrogates in stochastic optimization:
- a) as a sampling filter mechanism
- b) as a generation-wise surrogate

following I.G. Loshchilov, (Surrogate-assisted evolutionary algorithms, PhD, 2013) but many other implementations, e.g.,
- Kern et al., 2006, local Meta-model CMA-ES
- Runarsson 2004 & 2006, approx. ranking & ordinal regression...
Surrogates and stochastic optimization
Filtered sampling (1)

Simplified ACM-ES algorithm

\[ \nu > \lambda > \lambda' \]

\[ \{x^t, \ldots, x^{t+\nu}\} \] many

\( f(M(x^{1:v})) \) \& \( (\lambda' - 1) \) \( f(M(x^I)) \)'s

E
- update data-base \((X, F)\)
- update surrogate,
\[ \theta^t = \arg \min_{\theta} E(\theta, X, F) \]

M
model

\( (\lambda' - 1) \) random in \( I \)

\( \lambda' - 1 \) \( x^I \)'s

\( \nu > \lambda > \lambda' \)

\( \nu > \lambda \) new \( x \)'s according to \( d(x; \alpha^t) \)

S
- keep the best and \( \lambda - 1 \) other \( x \)'s with proba.
~ (rank) \( s(x; \theta^t) \)
Surrogates and stochastic optimization
Filtered sampling (2)

Implementation issues

**Ordinal regression** (ranking SVM, Herbrich et al., 1999) surrogates for rank based optimizers in order to preserve invariance property w.r.t. any monotonous transformation of $f()$.

Do not mistake the data-base and the population. In ACM-ES, the data-base is made of the $30 \times \sqrt{n}$ to $70 \times \sqrt{n}$ most recently evaluated points.

Having a **probabilistic choice of the x's** for CMA population and for the data-base is necessary to preserve points diversity. Otherwise, think of $\nu \to \infty$, all the points would tend to the surrogate optimum, as strategy we have criticized.

Performance

Speed-ups going from 2 to 4 were observed for dimensions 2 to 40 except for the (difficult) Rastrigin function $\leftarrow$ exploration / intensification trade-off and there is no surrogate usefulness control in ACM-ES ($\nu$ is fixed).

Surrogates and stochastic optimization

Generation-wise surrogate (1)

Principle: optimize on the surrogate for $g$ iterations and then for 1 iteration on the true function. Adjust $g$ according to the surrogate error.
Surrogates and stochastic optimization
Generation-wise surrogate (2)

- Points diversity is automatically guaranteed by the use of the stochastic optimizer.
- This algorithm uses surrogate error to adjust some parameters (the surrogate life-length).

**Performance:**
~ saACM-ES, Loshchilov 2012, which has speed-ups of 2 to 3 for $n=2$ to 20 over CMA-ES (in a version where other surrogate hyper-parameters* are optimized by minimizing surrogate error).

* the difference between surrogate parameters and hyper-parameters is that the hyper-parameters are typically set outside of the surrogate specific functions. Expl: regularization constants $C$ in Support Vector Machines.
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A very short introduction to kriging (1)

Kriging = conditional Gaussian processes
Random process are the function pendant to random variables
A sample = a function of \( x \)
conditional = the samples are forced to go through the data points \( X,F \)

[see Rasmussen & Williams, GPML, 2006 for more explanations]
A very short introduction to kriging (2)

Statistical model of \( f(x) \) : \( F(x) \sim N(m(x), s^2(x)) \)

\[
\left( m(x) \pm 1.96 s(x) \right) \text{ mean and 95\% confidence interval}
\]
and $F$ is correlated in space,

$$c(x) = \left[ \text{Cov} (F(x), F(x^i)) \right]_{i=1,M}$$

$$C = \left[ \text{Cov} (F(x^i), F(x^j)) \right]_{i,j}$$

Kriging average: $m(x) = \mu + c^T(x) C^{-1} (f - \mu \ 1)$

Kriging variance: $s^2(x) = \sigma^2 - c^T(x) C^{-1} c(x)$

**Assumptions**: $f(x)$ is a sample of Gaussian process with a given parameterized (stationary) kernel $\text{Cov} (F(x), F(x'))$ = a function of $|x - x'|$ and parameters $\theta$ (length scale)

( not all functions are kernel functions )

(left) squared exponential, $\text{Cor} (F(x), F(x')) = \exp(-1/2*(|x-x'|/\theta)^2)$

(right) exponential, $\text{Cor} (F(x), F(x')) = \exp(-(|x-x'|/\theta))$ , $\theta=0.2$
Surrogates with embedded error criteria

The kriging prediction variance, \( s^2(x) \), opens the way to a large family of criteria for controlling the quality of the surrogate during optimization.

Optimizing with surrogate has
- a main goal : provide an iterate \( x^i \) with a low \( f(x^i) \)
- a secondary goal : have the DoE of iterates \( (X,F) \) allow a surrogate that is accurate in high performance regions of the design space.

but since we don't know a priori where are the good regions of the design space, this amounts to an intensification / exploration compromise.

Mise en abyme (multi-crit for mono-crit) : can be seen as the two criteria problem,

\[
\begin{align*}
\min_{x \in S} m(x) \\
\max_{x \in S} s(x)
\end{align*}
\]

although the next single criteria may be more meaningful ...

Expected Improvement criterion

A natural measure of progress: the improvement,

\[ I(x) = [f_{\min} - F(x)]^+ | F(x) = f(x), \text{ where } [.]^+ \equiv \max(0, .) \]

- The expected improvement is known analytically.
- It is a parameter free measure of the exploration-intensification compromise.
- Its maximization defines the EGO deterministic global optimization algorithm.

\[ EI(x) = s(x) \times [u(x) \Phi(u(x)) + \varphi(u(x))] , \text{ where } u(x) = \frac{f_{\min} - m(x)}{s(x)} \]

[ Jones et al., Efficient Global Optimization of expensive black-box functions, JOGO, 1998 ]
At each iteration, EGO adds to the $t$ known points the one that maximizes EI,

$$x^{t+1} = \arg \max_x EI(x)$$

then, the kriging model is updated ...
kriging-based approaches

EI criterion: example
kriging-based approaches

EI criterion : 6D example

Hartman function, \( f(x^*) = -3.32 \), 10 points in initial DoE

(DiceOptim, D. Ginsbourger et al., 2009)
Our first example of surrogate criterion, including progress on $f$ and construction of the DoE.

Computational complexity: for kriging, the error is typically minus the likelihood or the cross-validation error --> a $(t \times t)$ covariance matrix need to be inverted many times, $O(t^3)$. 
kriging-based approaches

A one-stage approach (1)

So far, optimization of the surrogate criterion and construction of the surrogate (as another optimization problem) have been separated.

One stage approach : maximize the likelihood of the data points conditional on an hypothetical optimum \((x,f^{\text{target}})\):

\[
\text{CL}(x, \theta) = \text{Prob}(X, F \mid (x, f^{\text{target}}), \theta) \quad \text{(closed form from the multivariate normal law family)}
\]

Jones, 2001 (cf. earlier).
**Pros**: \( x \) and the surrogate are chosen together, which partly removes the initial guess on the surrogate that decides which \( x \) is sampled.

**Cons**: guess on \( f_{\text{target}} \), the optimization problem is of larger dimension (\( n + \text{dim}(\theta) \)).
Some other kriging-based criteria

- Statistical lower bound $m(x) - \alpha s(x)$: Cox and John 1997
- Quantile improvement (for noisy functions): Picheny et al. 2013
- Multi-points EI: Ginsbourger et al., 2010
- Multi-points PI (many targets), statistical lower bounds (many $\alpha$'s): Jones 2001


V. Picheny, D. Ginsbourger, Y. Richet, G. Caplin, Quantile-based optimization of noisy computer experiments with tunable precision, Technometrics, 2013
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Structured ensemble of surrogates

Each of the kriging samples is seen as a possible surrogate (a set of surrogates indexed by the random event \( \omega \)).

There is a distribution of optima knowing \((X,F)\) of density \( p(x^* \mid X, F) \).
Structured ensemble of surrogates
An informational approach

Principle: the next iterate is the one that provides the most information on the location of optima

⇒ The next iterate, $x_{t+1}$, is the point that reduces the most the conditional entropy of

$$p(x^* | (X, F) \cup (x_t^{t+1}, F(x_t^{t+1}))) \equiv p^{t+1}(x^* | x_t^{t+1})$$

$$x_{t+1} = \arg \min_{x \in S} \int_{S} -p^{t+1}(u | x) \log(p^{t+1}(u | x)) \, du$$

Pros: a nice way to summarize the contribution of a lot of surrogates.

Cons: high computational complexity.

Unstructured ensemble of surrogates
Multiple points generation

The simplest way to use many (say $m$) surrogates is to generate one iterate per surrogate (with your favorite surrogate optimization technique) and keep them all.

For $i=1,m$ do

update $i$-th surrogate , $s^i(x)$ , with $\mathcal{X}, F$

$x^i = \arg \min_{x \in \mathcal{S}} c(x, s^i(x))$

$(\mathcal{X}, F) = (\mathcal{X}, F) \cup (x^i, f(x^i))$

End

Unstructured ensemble of surrogates
Synthesizing many surrogates

One can make one (hopefully better) surrogate of many surrogates by linear combination,

\[ \hat{s}(x) = \sum_{i=1}^{m} w_i^* s_i(x) \]

The simplest way to choose the weights \( w_i \) is to optimize them to minimize the squared error (matrix notation),

\[ w^* = \arg \min_{w \in \mathbb{R}^m} \| f(X) - s(X)w \|^2 \]

(normal equations) \( \Rightarrow w^* = [s(X)^T s(X)]^{-1} s(X)^T f(X) \)

optim. weighted prediction: \( \hat{s}(x) = \begin{pmatrix} f(X)^T s(X) \end{pmatrix} \left[ s(X)^T s(X) \right]^{-1} \begin{pmatrix} s_1(x) \\ \vdots \\ s_m(x) \end{pmatrix} \)

( compare to the kriging average formula: this is an interpolation in the space of surrogates instead of \( S \) )


Conclusions: what about multi-objective optimization

Surrogates and optimization have had long-term, yet increasingly intricated relationships. And we just discussed mono-objective optimization. With multi-objective optimization, the range of possibilities still grows ...

- What does it change to go from mono-objective optimization to multi-objective optimization?
- Should one build one surrogate per objective function independently? Not independently? Or a unique surrogate to learn something about Pareto optimality in the space of optimization variables?
- Multi-objective problems are more difficult than mono-objective ones, so they need more points to be sampled: are there any computational limitations that will be hit?
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