Response surface methodology revisited
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If you believe that this document breaches copyright, please contact us providing details, and we will remove access to the work immediately and investigate your claim.
Response Surface Methodology (RSM) searches for the input combination that optimizes the simulation output. RSM treats the simulation model as a black box. Moreover, this paper assumes that simulation requires much computer time. In the first stages of its search, RSM locally fits first-order polynomials. Next, classic RSM uses steepest descent (SD); unfortunately, SD is scale dependent. Therefore, Part 1 of this paper derives scale independent ‘adapted’ SD (ASD) accounting for covariances between components of the local gradient. Monte Carlo experiments show that ASD indeed gives a better search direction than SD. Part 2 considers multiple outputs, optimizing a stochastic objective function under stochastic and deterministic constraints. This part uses interior point methods and binary search, to derive a scale independent search direction and several step sizes in that direction. Monte Carlo examples demonstrate that a neighborhood of the true optimum can indeed be reached, in a few simulation runs.

1 INTRODUCTION

RSM was invented by Box and Wilson (1951) for finding the input combination that minimizes the output of a real, non-simulated system. They ignored constraints. Also see recent publications such as Box (1999), Khuri and Cornell (1996), Myers (1999), and Myers and Montgomery (1995).

Later on, RSM was also applied to random simulation models, treating these models as black boxes (a black box means that there is no gradient information available; see Spall (1999)). Classic articles are Donohue, Houck, and Myers (1993, 1995); recent publications are Irizarry, Wilson, and Trevino (2001), Kleijnen (1998), Law and Kelton (2000, pp. 646-655), Neddermeijer et al. (2000), and Safizadeh (2002).

Technically, RSM is a stagewise heuristic that searches through various local (sub)areas of the global area in which the simulation model is valid. We focus on the first stage, which fits first-order polynomials in the inputs, per local area. This fitting uses Ordinary Least Squares (OLS) and estimates the SD path, as follows.

Let \( \hat{d}_j \) denote the value of the original (non-standardized) input \( j \) with \( j = 1, \ldots, k \). Hence \( k \) main or first-order effects (say) \( \beta_j \) are to be estimated in the local first-order polynomial approximation. For this estimation, classic RSM uses resolution-3 designs, which specify the \( n \times k \times 1 \) input combinations to be simulated. (Spall (1999) proposes to simulate only two combinations in his ‘simultaneous perturbation stochastic approximation’ or SPSA.) These input/output (I/O) combinations give the OLS estimates \( \hat{\beta}_j \), and the SD path uses the local gradient

\[
\hat{\beta}_1, \ldots, \hat{\beta}_k
\]

Unfortunately, RSM suffers from two well-known problems; see Myers and Montgomery (1995, pp. 192-194): (i) SD is scale dependent; (ii) the step size along the SD path is selected intuitively. For example, in a case study, Kleijnen (1993) uses a step size that doubles the most important input.

Our research contribution is the following. In Part 1 (§§2-3) we derive ASD; that is, we adjust the estimated first-order factor effects through their estimated covariance matrix. We prove that ASD is scale independent. In most of our Monte Carlo experiments with simple test functions, ASD indeed gives a better search direction. Note that we examine only the search direction, not the other elements of classic RSM. In Part 2 (§§4-5) we consider multiple outputs, whereas classic RSM assumes a single output. We optimize a stochastic objective function under multiple stochastic and deterministic constraints. We derive a scale
independent search direction – inspired by interior point methods - and several step sizes - inspired by binary search. This search direction namely scaled and projected SD, is a generalization of classic RSM’s SD. We then combine these search direction and step sizes into an iterative heuristic. Notice that if there are no binding constraints at the optimum, then classic RSM combined with ASD might suffice.

The remainder of this paper is organized as follows. For the unconstrained problem, §2 derives ASD, its mathematical properties, and its interpretation. §3 compares the search directions SD and ASD, by means of Monte Carlo experiments. §4 derives a novel heuristic combining a search direction and a step size procedure for constrained problems. §5 studies the performance of the novel heuristic by means of Monte Carlo experiments. §6 gives conclusions.

Note that this paper summarizes two separate papers, namely Kleijnen, Den Hertog, and Angün (2002), and Angün et al. (2002), which give all mathematical proofs and additional experimental results.

2 ADAPTED STEEPEST DESCENT

RSM uses the following approximation:

$$y = \beta_0 + \sum_{j=1}^{k} \beta_j d_j + e$$

(1)

where \( y \) denotes the predictor of the expected simulation output; \( e \) denotes the noise consisting of intrinsic noise caused by the simulation’s pseudo-random numbers (PRN) plus lack of fit. RSM assumes white noise; that is, \( e \) is normally, identically, and independently distributed with zero mean \( \mu_e \) and constant variance \( \sigma_e^2 \).

The OLS estimator of the \( q \) parameters \( \beta_0, \ldots, \beta_k \) in (1) is

$$\hat{\beta} = (XX')^{-1}Xw$$

(2)

with \( X \): \( N \times q \) matrix of explanatory variables including the ‘dummy’ variable with constant value 1; \( X \) is assumed to have linearly independent columns

\( N \times q \times m_i \): number of simulation runs

\( m_i \): number of replicates at input combination \( i \), with \( m_i \neq 0 \)

\( n \): number of different, simulated input combinations with \( n \geq N \).

\( w \): vector with \( N \) simulation outputs \( w_{t,r} \)

\( q \neq 1, \ldots, m_i \)

The noise in (1) may be estimated through the mean squared residual (MSR):

$$\hat{s}^2 = \frac{\hat{\epsilon}_i^2}{N-q}$$

where \( \hat{\epsilon}_i \) follows from (1) and (2):

$$\hat{\epsilon}_i = \hat{\beta}_0 + \sum_{j=1}^{k} \hat{\beta}_j d_{i,j}.$$

Kleijnen et al. (2002) derives the design point that minimizes the variance of the regression predictor:

$$\hat{d}_0 = \tilde{C}^{-1}b$$

where \( \tilde{C} = \hat{s}^2 \) is the covariance matrix of \( \hat{\beta}_0 \), which equals \( \hat{\beta} \) excluding the intercept \( \hat{\beta}_0 \):

$$\text{cov} (\hat{\beta}) = s^2 (XX')^{-1}$$

(4)

where \( a \) is a scalar, \( b \) a \( k \)-dimensional vector, and \( C \) a \( k \times k \) matrix.

Now we consider the one-sided \( 1\gamma \) confidence interval ranging from \( \hat{y}_0 \) to

$$\hat{y}_{\max} = x\hat{\beta} + \sum_{i=1}^{N} \tilde{t}_{\gamma,q} \hat{s} \sqrt{(x'XX')^{-1}x}$$

(5)

where \( x \neq 0 \), \( \tilde{t}_{\gamma,q} \) and \( \tilde{t}_{\gamma,q} \) denotes the \( 1\gamma \) quantile of the \( t \) distribution with \( N \) \( q \) degrees of freedom.

ASD selects \( d^2 \), the design point that minimizes the maximum output predicted through (5) (this gives both a search direction and a step size). Kleijnen et al. (2002) derives

$$d^2 = \tilde{C}^{-1}b - \tilde{C}^{-1}\hat{\beta}_0$$

(6a)

where \( \tilde{C}^{-1}b \) is derived from (4), \( \tilde{C}^{-1}\hat{\beta}_0 \) is the ASD direction, and \( \gamma \) is the step size:

$$\gamma = \frac{a \cdot \hat{\beta}_0 \cdot \tilde{C}^{-1}b}{(\tilde{t}_{N-q,\gamma}^2 \hat{s})^2 \cdot \hat{\beta}_0 \cdot \tilde{C}^{-1}b}$$

(6b)
We mention the following mathematical properties and interpretations of ASD.

The first term in (6a) means that the ASD path starts from the point with minimal predictor variance. The second term means that the classic SD direction $\hat{\beta}_0$ (second term’s last factor) is adjusted for the covariance matrix of $\hat{\beta}_0$, see $C$ in (4). The step size $?\beta$ is quantified in (6b).

Kleijnen et al. (2002) proves that ASD is scale independent.

In case of large signal/noise ratios $\hat{\beta}_0 \var/ \beta_jj$, the denominator under the square root in (6b) is negative so (6) does not give a finite solution for $d^+$. Indeed, if the noise is negligible, we have a deterministic problem, which our technique is not meant to address (many other researchers - including Conn et al. (2000) - study optimization of deterministic simulation models).

In case of a small signal/noise ratio, no step is taken. Kleijnen et al (2002) further discusses two subcases: (i) the signal is small; (ii) the noise is big.

3 COMPARISON OF ASD AND SD THROUGH MONTE CARLO EXPERIMENTS

To compare the ASD and SD search directions, we perform Monte Carlo experiments. The Monte Carlo method is an efficient and effective way to estimate the behavior of search techniques applied to random simulations; see Kleijnen et al. (2002).

We limit the example to two inputs, so $k = 2$. We generate the simulation output $w$ through a second-order polynomial with white noise:

$$w = \beta_0 + \beta_1 d_1 + \beta_2 d_2 + \beta_{1,1} d_1^2 + \beta_{2,2} d_2^2 + \beta_{1,2} d_1 d_2 + e. \quad (7)$$

The response surface (7) holds for the global area, for which we take the unit square: $?1?d_1?1$ and $?1?d_2?1$. (We have already seen that RSM fits first-order polynomials locally.)

In the local area we use a one-at-a-time design, because this design is non-orthogonal - and in practice designs in the original inputs are not orthogonal (see Kleijnen et al. (2002)). The specific local area is in the lower corner of Figure 1. To enable the computation of the MSR, we simulate one input combination twice: $m_1 = 2$.

![Figure 1: Tilted Ellipsoid Contours $E[w|d_1, d_2]$ with Global and Local Experimental Areas](image)

We consider a specific case of (7): $\beta_0 = \beta_1 = \beta_2 = 0$, $\beta_{1,1} = 2$, $\beta_{1,2} = -2$, $\beta_{2,2} = -1$, so the contour functions (for example, iso-cost curves) form ellipsoids tilted relative to the $d_1$ and $d_2$ axes. Hence, (7) has as its true optimum $d^* = 0,0$.

After fitting a first-order polynomial, we estimate the SD and ASD paths starting from $d_0 = (0.85, -0.95)$, explained above (4).

In this Monte Carlo experiment we know the truly optimal search direction, namely the vector (say) $g$ that starts at $d_0$ and ends at the true optimum $0,0$. So we compute the angle (say) $?\hat{\beta}$ between the true search direction $g$ and the estimated search direction $p$:

$$?\hat{\beta} = \arccos \frac{g \cdot p}{|g| |p|}. \quad (8)$$

Obviously, the smaller $?\hat{\beta}$ is, the better the search technique performs.

To estimate the distribution of $?\hat{\beta}$ defined in (8), we take 100 macro-replications. Figure 2 shows a bundle of 100 $p$’s around $g$ when $s = 0.1$. In each macro-replicate, we apply SD and ASD to the same I/O data $w, d_1, d_2$. We characterize the resulting empirical $?\hat{\beta}$ distribution through several statistics, namely its average, standard deviation, and specific quantiles; see Table 1.
Table 1: Statistics in case of Interactions, for ASD and SD’s Estimated Angle Error $\hat{\theta}$ (in Degrees)

<table>
<thead>
<tr>
<th>Statistics</th>
<th>$s_\theta$ ? 0.10</th>
<th>$s_\theta$ ? 0.25</th>
</tr>
</thead>
<tbody>
<tr>
<td>ASD</td>
<td>9.72</td>
<td>16.01</td>
</tr>
<tr>
<td>SD</td>
<td>10.14</td>
<td>17.33</td>
</tr>
<tr>
<td>Average</td>
<td>9.68</td>
<td>16.02</td>
</tr>
<tr>
<td>Standard deviation</td>
<td>6.23</td>
<td>8.99</td>
</tr>
<tr>
<td>Median (50% quantile)</td>
<td>12.37</td>
<td>21.12</td>
</tr>
<tr>
<td>75% quantile</td>
<td>15.66</td>
<td>27.05</td>
</tr>
<tr>
<td>95% quantile</td>
<td>17.41</td>
<td>30.08</td>
</tr>
<tr>
<td>0% quantile</td>
<td>0.85</td>
<td>1.46</td>
</tr>
</tbody>
</table>

Further, we perform the Monte Carlo experiment for two noise values: $s_\epsilon$ is 0.10 or 0.25. We use the same PRN for both values. In case of high noise, the estimated search directions may be very wrong. Nevertheless, ASD still performs better; see Table 1.

In general, ASD performs better than SD, unless we focus on outliers; see Kleijnen et al. (2002).

4 MULTIPLE RESPONSES: INTERIOR POINT AND BINARY SEARCH APPROACH

In Part 1, we assumed a single response of interest – denoted by $w$ in (2). Now we consider a more realistic situation, namely the simulation generates multiple outputs. For example, an academic inventory simulation defines $w$ in (2) as the sum of inventory-carrying, ordering, and out-of-stock costs, whereas a practical simulation minimizes the sum of the expected inventory-carrying and ordering costs provided the service probability exceeds a pre-specified value.

In RSM, there have been several approaches to multiresponse optimization. Khuri (1996) surveys most of these approaches (including desirability functions, generalized distances, and dual responses). Angün et al. (2002) discusses drawbacks of these approaches. To overcome these drawbacks, we propose the following alternative based on mathematical programming.

We select one of the responses as the objective and the remaining (say) $z$ responses as constraints. The SD search would soon hit the boundary of the feasible area formed by these constraints, and would then creep along this boundary. Instead, our search starts in the interior of the feasible area and avoids the boundary; see Barnes (1986) on Karmarkar’s algorithm for linear programming.

Note that our approach has the additional advantage of avoiding areas in which the simulation model is not valid and may even crash.

Formally, our problem becomes:

$$\begin{align*}
\text{minimize} & \quad E[w_0, \ldots, w_h] \\
\text{subject to} & \quad E[w_h; \ldots, a_h] \geq 0, \ldots, z \geq 1
\end{align*}$$

where $d$ is the vector of simulation inputs, $l$ and $u$ the deterministic lower and upper bounds on $d$, $a_h$ the right-hand-side value for the $h$th constraint, and $w_h; \ldots, a_h$ is the response $h$.

Note that probabilities (for example, service percentages) can be formulated as expected values of indicator functions. Further, the multiple simulation responses are correlated, since they are estimated through the same PRN fed into the same simulation model.

As in Part 1, we locally fit a first-order polynomial – but now for each response; see (1). However, the noise $\epsilon$ is now multi-variate normal – still with zero means but now with covariance matrix (say) $S$. Yet, since the same design is used for all $z$ responses, the GLS estimator reduces to the OLS estimator; see Ruud (2000, p. 703).

Therefore we still use (2), but to the symbol $\hat{\beta}$ we add the subscript $h$. Further, for $h$? $h$? $0, \ldots, z$ we estimate $S$ through the analogue of (3):

$$s_{h?h}^2 = \frac{\sum (w_{ht} - \hat{y}_{ht}) (w_{ht} - \hat{y}_{ht})}{(k+1)}.$$  \hspace{1cm} (10)

We introduce $B$? $h_1, \ldots, h_z$? $？$?, where $b_h$ $h$? $1, \ldots, z+1$? denotes the vector of OLS estimates $\hat{\beta}_{0:h}$ (excluding the intercept $\hat{\beta}_{0:h}$) for the $h$th response.

Adding slack vectors $s, r,$ and $v$, we obtain

$$\begin{align*}
\text{minimize} & \quad E[w_0, \ldots, w_h] \\
\text{subject to} & \quad E[w_h; \ldots, a_h] \geq 0, \ldots, z
\end{align*}$$

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We introduce $B$? $h_1, \ldots, h_z$? $？$?, where $b_h$ $h$? $1, \ldots, z+1$? denotes the vector of OLS estimates $\hat{\beta}_{0:h}$ (excluding the intercept $\hat{\beta}_{0:h}$) for the $h$th response.

Adding slack vectors $s, r,$ and $v$, we obtain
\[
\begin{align*}
\text{minimize} & \quad d \mathbf{b}_0 \\
\text{subject to} & \quad \mathbf{B}d + s^2c_w^2d + r^2u, d, v \leq l, \quad (11)
\end{align*}
\]

where \( \mathbf{b}_0 \) denotes the vector of OLS estimates \( \hat{\mathbf{B}}_{0.0} \) (excluding the intercept \( \hat{\mathbf{B}}_{0.0} \)) for \( \mathbf{w}_0 \), and \( c \) is the vector with components \( c_h, a_h, \hat{\mathbf{B}}_{0,0}, h \). \( s, r, v \) \( > 0 \)

Through (11) we obtain a local linear approximation for (9). Then using ideas from interior point methods - more specifically the affine scaling method - Angün et al. (2002) derives the following search direction:

\[
\mathbf{p} = (\mathbf{B}^T \mathbf{S}^{-2} \mathbf{R}^{-2} + \mathbf{V}^{-2} \mathbf{b}_0
\]

where \( \mathbf{S}, \mathbf{R}, \) and \( \mathbf{V} \) are diagonal matrices with the current estimated slack vectors \( \mathbf{s}, \mathbf{r}, \mathbf{v} \). Obviously, \( \mathbf{R} \) and \( \mathbf{V} \) in (12) are known as soon as the deterministic input \( d \) to the simulation is selected; \( \mathbf{S} \) in (12) is estimated from the simulation output, not from the local approximation. Unlike SD, the search direction (12) is scale independent (the inverse of the matrix within the parentheses in (12) scales and projects the estimated SD direction, \( \mathbf{b}_0 \)).

Having estimated a search direction for a specific starting point through (12), we must next select a step size. Actually, we run the simulation model for several step sizes in that direction, as follows.

First, we compute the maximum step size assuming that the local approximation (11) holds globally:

\[
?_{\text{max}} = \max \{0, \min \{?_1, ?_2, ?_3\} \}
\]

where

\[
?_1 = \min \{|c_h, \mathbf{h}|^2d^2|/\mathbf{p}^\mathbf{b}_h, h?| 1, ..., z?1, \mathbf{p}^\mathbf{b}_h ? 0\}
\]

\[
?_2 = \min \{|a_j, \mathbf{d}^2|/\mathbf{p}^j, j?1, ..., k), \mathbf{p}_j ? 0\}
\]

\[
?_3 = \min \{|d_j, \mathbf{d}^2|/\mathbf{p}^j, j?1, ..., k), \mathbf{p}_j ? 0\}
\]

To increase the probability of staying within the interior of the feasible region, we take only 80% of \( ?_{\text{max}} \) as our maximum step size.

The subsequent step sizes are inspired by binary search, as follows. We systematically halve the current step size along the search direction. At each step, we select as the best point the one with the minimum value for the simulation objective \( \mathbf{w}_0 \), provided it is feasible. We stop the search in a particular direction after a user-specified number of iterations (say) \( G ? 3 \). For details see Angün et al. (2002).

For all these steps we use common random numbers (CRN), in order to better test whether the objective improves. Moreover, we test whether the other \( z?1 \) responses remain within the feasible area: we test the slack vector \( s \), introduced in (11). These \( z \) tests use ratios instead of absolute differences, to avoid scale dependence.

A statistical complication is that these ratios may not have finite moments. Therefore we test their medians (not their means). For these tests we use Monte Carlo sampling, which takes negligible computer time compared with the expensive simulation runs. This Monte Carlo takes (say) \( K ? 1000 \) samples from the assumed distributions with means and variances estimated through the simulation; in the numerical example of the next section we assume \( z \) normal distributions ignoring correlations between these responses. From these Monte Carlo samples we compute slack ratios.

We formulate pessimistic null-hypotheses; that is, we ‘accept’ an input combination only if it gives a significantly lower objective value and all its \( z?1 \) slacks imply a ‘feasible’ solution. Actually, our interior point method implies that a new slack value is a percentage – say, 20% - of the old slack value; our pessimistic hypotheses make our acceptable area smaller than the original feasible area in (9).

After we have run \( G \) simulations along the search path, we find a ‘best’ solution - so far. Now we wish to re-estimate the search direction \( \mathbf{p} \) defined in (12). Therefore we again use a resolution-3 design in the \( k \) factors. We still use CRN; actually, we take the same seeds as we used for the very first local exploration. We save one (expensive) run by using the best combination found so far, as one of the combinations for the design.

We stop the whole search when either the computer budget is exhausted or the search returns to an old combination twice. When the search returns to an old combination for the first time, we use a new set of seeds.

5 MONTE CARLO EXPERIMENTS FOR MULTIPLE RSM

Like in Part 1 (§3), we study the novel procedure - explained in §4 - by means of a Monte Carlo example. As in §3, we assume globally valid functions quadratic in two inputs, but now we consider three responses; moreover, we add deterministic ‘box’ constraints for the two inputs:
where the noise has $s_0 = 1$, $s_1 = 0.15$, $s_2 = 0.4$, and correlations $\rho_{0,1} = 0.6$, $\rho_{0,2} = 0.3$, $\rho_{1,2} = 0.1$.

It is easy to derive the analytical solution as $d^* = [1.24, 0.52]$ with a mean objective value of 22.96 approximately.

We select the initial local area shown in the lower left corner of Figure 3. We run 100 macro-replicates; Figure 3 displays the macro-replicate that gives the median result for the objective; that is, 50% of the macro-replicates have worse objective values. In this figure we have $d^* = [1.46, 0.49]$ and an estimated objective of 25.30 approximately.

**Table 2: Estimated Objective and Slacks over 100 Macro-replicates**

<table>
<thead>
<tr>
<th></th>
<th>Criterion 1</th>
<th>Criterion 2</th>
<th>Criterion 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>10th quantile</td>
<td>0.04</td>
<td>0.03</td>
<td>0.03</td>
</tr>
<tr>
<td>25th quantile</td>
<td>0.06</td>
<td>0.12</td>
<td>0.15</td>
</tr>
<tr>
<td>50th quantile</td>
<td>0.10</td>
<td>0.25</td>
<td>0.29</td>
</tr>
<tr>
<td>75th quantile</td>
<td>0.19</td>
<td>0.43</td>
<td>0.49</td>
</tr>
<tr>
<td>90th quantile</td>
<td>0.18</td>
<td>0.61</td>
<td>0.50</td>
</tr>
</tbody>
</table>

Our conclusion is that the heuristic reaches the desired neighborhood of the real optimum in a relatively small number of simulation runs. Once the heuristic reaches this neighborhood, it usually stops at a feasible point.

**CONCLUSIONS**

In Part 1 of this paper we addressed the problem of searching for the simulation input combination that minimizes the output. RSM is a classic technique for tackling this problem, but it uses SD, which is scale dependent. Therefore we devised adapted SD (ASD), which corrects for the covariances of the estimated gradient components. ASD is scale independent. Our Monte Carlo experiments demonstrate that - in general - ASD gives a better search direction than SD.

In Part 2, we account for multiple simulation responses. We use a mathematical programming approach; that is, we minimize one random objective under random and deterministic constraints. As in classic RSM, we locally fit functions linear in the simulation inputs. Next we apply interior point techniques to these local approximations, to estimate a search direction. This direction is scale independent. We take several steps into this direction, using binary search and statistical tests. Then we re-estimate these local linear functions, etc. Our Monte Carlo experiments demonstrate that our method indeed approaches the true optimum, in relatively few runs with the (expensive) simulation model.

**REFERENCES**


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