Using the rsm package

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1 Overview

The rsm package provides several useful functions to facilitate response-surface analysis. The primary one is the rsm function itself, which is an extension of 1m but with some enhancements. In specifying a model in rsm, the model formula is just like in 1m, but the response-surface portion of the model is specified using one or more of the special functions FO (first-order), TWI (two-way interactions), PQ (pure quadratic), or SO (second-order, an alias for all three of the previous functions, combined). The summary method for rsm results includes the usual regression summary (but with the coefficients compactly relabeled), an analysis of variance table with a lack-of-fit test, and additional information depending on the order of the model.

An important aspect of response-surface analysis is using an appropriate coding transformation of the data. The functions coded.data, as.coded.data, decode.data, code2val, and val2code facilitate these transformations; we simply provide formulas for the desired transformations. If a coded.data object is used in place of an ordinary data.frame in the call, to rsm, then appropriate additional output is provided in the summary and steepest outputs.

Of course, before we get to analysis, we need a good design for collecting the required data. The functions ccd and bbd are provided for generating two of the most popular classes of designs—central-composite designs (CCDs) and Box-Behnken designs (BBDs). In addition, ccd.pick allows one to take a quick look at various combinations of choices for CCDs and find the most suitable ones.

Auxiliary functions include steepest for finding a path of steepest ascent (for second-order models, this uses ridge analysis); and contour for obtaining a contour plot of the response surface.

2 Generating a design

Suppose that you want to experiment on a process with an aim to improving its yield. You have already done a little bit of preliminary experimentation and have identified five variables that you want to manipulate experimentally. Our plan is to develop a central-composite design, which consists of some blocks of "cube" or factorial points plus center points, and other blocks with "star" or axis points, plus center points. The cube points will be placed at positions ± 1 in coded units, and the axis points will be at $\pm \alpha$. Initially, we will only collect data on one or more of the cube blocks; then, after analyzing these data with a first-order model, we can either proceed to collect data on the other blocks and fit a second-order model, or if the fit is reasonably linear, we may want to forego the extra blocks and instead follow a path of steepest ascent.

There are a lot of choices to be made—how many center points, whether we have replications, whether the cube block(s) are fractional or full factorial, and what α to use. With response surfaces,

it is desirable to have a rotatable design (where the variance of the estimated response depends only on the distance from zero). On the other hand, a CCD is built in blocks, and it is a good idea to make the block effects independent of the effects needed to estimate the response surface. To help make good choices, we can run the function ccd.pick to explore some possibilities. Suppose that, for practical reasons, we want no more than 16 cube points in a block, but we'd also consider ones with only 8 cube points (e.g., all $2^5 = 32$ factor combinations dividing into 4 blocks of 8 each). Since there are 5 factors, there are $2 \times 5 = 10$ axis-point positions, but it might be worth considering replicating the axis points rather than having lots of center points. And we don't want the total number of runs in the design to be too excessive—say 65 at most.

2.1 Identifying a good design

Given these considerations we will run $\mathtt{ccd.pick}$ to obtain ideas for good designs. It will compute the α values needed for rotatability and orthogonal blocking for various combinations of numbers of cube points, center points, replications, etc., and show the best few after sorting in a specified order (by default, a measure of how well the two α s agree). Here is a suitable call based on the above discussion:

```
> library(rsm)
> ccd.pick(5, n.c = c(8, 16), blks.c = c(1, 2, 4), wbr.s = 1:2,
       restrict = "N <= 65")
   n.c n0.c blks.c n.s n0.s bbr.c wbr.s bbr.s
                                                       N alpha.rot alpha.orth
1
    16
           6
                    1
                       10
                              1
                                      1
                                             1
                                                    1 33
                                                           2.000000
                                                                        2.000000
                              2
2
    16
           8
                    1
                       10
                                      1
                                             1
                                                    1 36
                                                           2.000000
                                                                        2.000000
3
          10
                    1
                       10
                              3
                                      1
                                             1
                                                    1 39
                                                           2.000000
                                                                        2.000000
    16
4
                    2
                       20
                              1
                                             2
    16
           5
                                      1
                                                    1 63
                                                           2.000000
                                                                        2.000000
5
                    2
                              7
    16
           8
                       10
                                      1
                                             1
                                                    1 65
                                                           2.378414
                                                                        2.380476
                              7
6
     8
           4
                    4
                       10
                                      1
                                             1
                                                    1 65
                                                           2.378414
                                                                        2.380476
7
    16
           1
                    2
                       10
                              2
                                      1
                                             1
                                                    1 46
                                                           2.378414
                                                                        2.376354
                    2
                                             1
8
           5
                       10
                              5
                                      1
                                                           2.378414
                                                                        2.390457
    16
                                                    1 57
9
    16
            4
                    2
                       10
                              4
                                      1
                                             1
                                                    1 54
                                                           2.378414
                                                                        2.366432
           2
                                             1
10
     8
                    4
                       10
                              4
                                      1
                                                    1 54
                                                           2.378414
                                                                        2.366432
```

The first one listed has a total of N=33 runs; it has blks.c=1 cube block with n.c=16 cube points and n0.c=6 center points; and star block with n.s=10 axis points (wbr.s=1 at each position) and n0.s=1 center point; with these settings, the design is both rotatable and orthogonal if we use $\alpha=2$ for the axis-point positions. The 63-run design 4 is the only one shown where the axis points are replicated; it has two 16-point cube blocks with 5 center points each, and only one center point, but replicated axis points, in the star block. This design has the pleasing feature of requiring 21 runs in each block, ad it is both rotatable and orthogonal using $\alpha=2$. In the remaining designs, there is a slight discrepancy between the α s required for rotatability and orthogonality. Designs 5 and 6 have exactly the same number of runs, and differ only in where there are 2 blocks with 16 cube points or 4 blocks of 8 cube points. Design 10 has a slightly greater discrepancy between the α s than design 6, but fewer total runs.

2.2 Generating a CCD

Suppose that we decide to go with Design 1. To generate this design, the 16-run cube block is a half-fraction of the full 32-run design in 5 factors. This can be generated by confounding the main

effect of one factor with the 4-way interaction of the others. I flipped a coin and decided to use the negative of this interaction. The ccd function can generate and randomize the design:

```
> ccd(x1 + x2 + x3 + x4, x5 - x1 * x2 * x3 * x4, n0 = c(6, 1))
```

```
Block x1 x2 x3 x4 x5
C1.19
            1
               0
                  0
                      0
                         0
C1.2
           1
               1 -1 -1 -1
                             1
C1.18
           1
               0
                  0
                      0
                         0
                             0
C1.22
               0
                  0
                             0
           1
                      0
                         0
C1.8
           1
               1
                  1
                      1 -1
                             1
C1.10
            1
               1 -1 -1
                            -1
                         1
C1.15
            1 -1
                          1
C1.17
           1
               0
                  0
                      0
                         0
C1.5
           1 -1 -1
                      1 -1
                             1
C1.11
           1 -1
                  1 -1
                         1 - 1
C1.14
            1
               1 -1
                      1
                         1
                             1
C1.6
            1
               1 -1
                      1 -1 -1
C1.12
            1
               1
                  1 -1
                         1
C1.3
            1 -1
                  1
                    -1 -1
                             1
C1.16
               1
                  1
                      1
                         1 - 1
C1.4
               1
                  1 -1 -1 -1
           1
C1.9
           1 -1 -1
                    -1
                          1
                             1
C1.20
           1
               0
                  0
                      0
                         0
                             0
C1.13
           1 -1 -1
                      1
                         1 -1
C1.7
            1
             -1
                  1
                      1 -1 -1
C1.21
            1
               0
                  0
                      0
                         0
                             0
C1.1
           1 -1 -1 -1 -1
           2
                  0
S2.7
S2.11
           2
               0
                  0
                      0
                             0
                         0
S2.3
           2
               0 -2
                      0
                         0
                             0
           2
               2
S2.2
                  0
                      0
                         0
                             0
S2.9
           2
               0
                  0
                         0 -2
                      0
           2
                             2
S2.10
               0
                  0
                      0
                         0
S2.8
           2
               0
                  0
                      0
                         2
                             0
           2
               0
                  0
                      2
                             0
S2.6
                         0
S2.4
           2
               0
                  2
                      0
                         0
                             0
S2.1
           2
             -2
                  0
                      0
                         0
                             0
                 0 -2
               0
S2.5
```

By default, ccd chooses α for orthogonality. If we want to name the variables x1,x2,..., we can just give the number of variables instead of a formula in the first argument:

```
> ccd(4, x5 - x1 * x2 * x3 * x4, n0 = c(6, 1))
```

To generate design 4, we use the full 32-run design, but divided into blocks two blocks of 16 runs by confounding the 5-way interaction:

```
> des4 = ccd(5, , Block ~ x1 * x2 * x3 * x4 * x5, wbr = c(1, 2),
+ n0 = c(5, 1))
```

The wbr argument specifies within-block replications for cube blocks and star blocks, respectively. There is also a bbr argument for between-block replications (i.e. additional blocks with the same factor combinations).

The ccd call for generating design 5 would be similar to the one above, but no wbr argument is needed. For design 10 (or design 6), we need to block the 32 cube points into four sets of 8, by confounding two effects with blocks:

```
> des10 = ccd(5, , Block ~ c(x1 * x2 * x3, x3 * x4 * x5), n0 = c(2, + 4))
```

These designs, while having more total runs, may be preferred over design 1 because it is possible to run only one block (10 runs, compared with 22 runs with design 1) and still be able to estimate some first-order effects.

Because experimentation can be very expensive, it would be terrible to run the design only to find out you can't estimate all the effects. For that reason, ccd does a check to make sure we can do an analysis:

```
> bad.des = ccd(5, , Block ~ c(x1 * x2 * x3 * x4, x2 * x3 * x4 * x5), n0 = c(2, 4))

Warning in ccd(~x1 + x2 + x3 + x4 + x5, , Block ~ c(x1 * x2 * x3 * x4, x2 * : Some 1st or 2nd-order terms are aliased in the cube portion of this design
```

The problem here is that the generalized interaction between the two effects, $x_1x_2x_3x_4 \cdot x_2x_3x_4x_4 = x_1x_5$, is also confounded with blocks. Actually, by the time center points and axis points are added, x_1x_5 is only partially confounded; but this is still not a desirable design.

2.3 Box-Behnken designs

The bbd function is provided to generate Box-Behnken designs. These are fractional 3^k designs capable of fitting second-order models. Advantages are that they sometimes require fewer runs than a CCD, and each factor has only 3 levels instead of 5. Disadvantages are that they cannot be built-up in blocks like a CCD, and they are not rotatable. BBDs are available only for 3, 4, 5, 6, and 7 factors; and only 4- and 5-factor designs can be blocked orthogonally. Here is a BBD for 5 factors (by default, in two blocks)

```
> bbd5 = bbd(5, n0 = 1)
> nrow(bbd5)
[1] 42
```

In this case, some CCDs have fewer runs. However, the size of one block is comparable to that of the first design, and we could use it for first-order analysis.

3 Chemical reactor example

The provided dataset ChemReact comes from Table 7.7 of Myers and Montgomery (2002).

> ChemReact

```
Time
           Temp Block Yield
  80.00 170.00
                    B1
                        80.5
  80.00 180.00
                    B1
                        81.5
  90.00 170.00
                    B1
                        82.0
  90.00 180.00
                        83.5
                    B1
5
  85.00 175.00
                    B1
                        83.9
6
  85.00 175.00
                    B1
                        84.3
7
  85.00 175.00
                    В1
                        84.0
  85.00 175.00
                    B2
                        79.7
  85.00 175.00
                    B2
                        79.8
10 85.00 175.00
                    B2
                        79.5
11 92.07 175.00
                    B2
                        78.4
12 77.93 175.00
                    B2
                        75.6
13 85.00 182.07
                    B2
                        78.5
14 85.00 167.93
                    B2
                        77.0
```

The context is that block B1 of this data were collected first and analyzed, after which block B2 was added and a new analysis was done. Accordingly, we will illustrate the analysis in two stages.

3.1 Coding of predictors

First, though, we need to take care of coding issues. The data are provided in their original units, and the original experiment (block B1) used factor settings of Time = 85 ± 5 and Temp = 175 ± 5 , with three center points. Thus, the coded variables are $x_1 = (\text{Time} - 85)/5$ and $x_1 = (\text{Temp} - 175)/5$. Let's create a coded dataset with the appropriate codings. We do this via formulas:

```
> CR = coded.data(ChemReact, x1 ~ (Time - 85)/5, x2 ~ (Temp - 175)/5)
> CR[1:7, ]
  x1 x2 Block Yield
1 -1 -1
            В1
                80.5
2 -1
            В1
                81.5
      1
3
   1 -1
           В1
                82.0
4
   1
      1
           В1
                83.5
5
      0
                83.9
   0
            В1
6
   0
      0
            В1
                84.3
   0
      0
                84.0
            В1
```

```
Variable codings ...
x1 ~ (Time - 85)/5
x2 ~ (Temp - 175)/5
```

3.2 Analysis of initial block

The initial 7 runs are only good enough to estimate a first-order model. We will fit this by calling rsm just like we would 1m, but use the special function FO (first-order response surface) in the model formula:

```
> CR.rsm1 = rsm(Yield ~ FO(x1, x2), data = CR, subset = 1:7)
> summary(CR.rsm1)
Call:
rsm(formula = Yield ~ FO(x1, x2), data = CR, subset = 1:7)
Residuals:
                                                      7
      1
              2
                      3
                                      5
                                              6
-0.8143 -1.0643 -1.0643 -0.8143 1.0857 1.4857 1.1857
Coefficients:
            Estimate Std. Error t value Pr(>|t|)
(Intercept)
            82.8143
                         0.5472 151.346 1.14e-08 ***
              0.8750
                         0.7239
                                  1.209
                                           0.293
x1
              0.6250
                         0.7239
                                  0.863
                                           0.437
x2
Signif. codes: 0 âĂŸ***âĂŹ 0.001 âĂŸ**âĂŹ 0.01 âĂŸ*âĂŹ 0.05 âĂŸ.âĂŹ 0.1 âĂŸ âĂŹ 1
Residual standard error: 1.448 on 4 degrees of freedom
Multiple R-squared: 0.3555,
                                   Adjusted R-squared: 0.0333
F-statistic: 1.103 on 2 and 4 DF, p-value: 0.4153
Analysis of Variance Table
Response: Yield
            Df Sum Sq Mean Sq F value Pr(>F)
FO(x1, x2)
             2 4.6250 2.3125 1.1033 0.41534
Residuals
             4 8.3836 2.0959
Lack of fit 2 8.2969 4.1485 95.7335 0.01034
Pure error
             2 0.0867 0.0433
Direction of steepest ascent (at radius 1):
0.8137335 0.5812382
Corresponding increment in original units:
             Temp
    Time
4.068667 2.906191
```

Note that the summary includes a lack-of-fit test, and it is significant. We can try adding two-way interactions to see if it helps:

```
> CR.rsm1.5 = update(CR.rsm1, . ~ . + TWI(x1, x2))
> summary(CR.rsm1.5)
```

```
Call:
```

rsm(formula = Yield ~FO(x1, x2) + TWI(x1, x2), data = CR, subset = 1:7)

Residuals:

1 2 3 4 5 6 7 -0.9393 -0.9393 -0.9393 -0.9393 1.0857 1.4857 1.1857

Coefficients:

Estimate Std. Error t value Pr(>|t|)

(Intercept) 82.8143 0.6295 131.560 9.68e-07 *** 0.8750 0.8327 1.051 0.371 x1x2 0.6250 0.8327 0.751 0.507 0.1250 0.890 x1:x20.8327 0.150

Signif. codes: 0 âĂŸ***âĂŹ 0.001 âĂŸ**âĂŹ 0.01 âĂŸ*âĂŹ 0.05 âĂŸ.âĂŹ 0.1 âĂŸ âĂŹ 1

Residual standard error: 1.665 on 3 degrees of freedom

Multiple R-squared: 0.3603, Adjusted R-squared: -0.2793

F-statistic: 0.5633 on 3 and 3 DF, p-value: 0.6755

Analysis of Variance Table

Response: Yield

Df Sum Sq Mean Sq F value Pr(>F)
FO(x1, x2) 2 4.6250 2.3125 0.8337 0.515302
TWI(x1, x2) 1 0.0625 0.0625 0.0225 0.890202
Residuals 3 8.3211 2.7737
Lack of fit 1 8.2344 8.2344 190.0247 0.005221
Pure error 2 0.0867 0.0433

Stationary point of response surface:

x1 x2

-5 -7

Stationary point in original units:

Time Temp

60 140

Eigenanalysis:

\$values

[1] 0.0625 -0.0625

\$vectors

[,1] [,2]

[1,] 0.7071068 -0.7071068

[2,] 0.7071068 0.7071068

The lack of fit is still significant. Note that the summary output now shows a canonical analysis rather than the direction of steepest ascent, as the response surface now has second-order terms.

3.3 Analysis of combined blocks

The lack-of-fit results motivate us to collect additional runs at "star" points, plus some additional center points; these are the second block. In coded units, the data are

```
> CR[8:14, ]
```

```
x2 Block Yield
       x1
    0.000
           0.000
                     B2
8
                         79.7
9
    0.000
           0.000
                     B2
                         79.8
    0.000
           0.000
                         79.5
    1.414
           0.000
                     B2
                         78.4
11
12 -1.414
           0.000
                     B2
                         75.6
   0.000
           1.414
                     B2
                         78.5
13
    0.000 - 1.414
                     B2
                         77.0
```

```
Variable codings ... x1 ~ (Time - 85)/5
```

x2 ~(Temp - 175)/5

The choice of $\alpha = \sqrt{2}$ provides for rotatability, and the blocks are orthogonal as well. To do the analysis of the combined data, we should account for the block effect. We could fit a full second-order model by including F0, TWI, and PQ terms, but this is more easily done using S0 which generates all three sets of variables:

```
> CR.rsm2 = rsm(Yield ~ Block + SO(x1, x2), data = CR)
> summary(CR.rsm2)
```

Call:

```
rsm(formula = Yield ~ Block + SO(x1, x2), data = CR)
```

Residuals:

```
Min 1Q Median 3Q Max -0.19543 -0.09369 0.02157 0.06153 0.20457
```

Coefficients:

```
t value Pr(>|t|)
            Estimate Std. Error
(Intercept) 84.09543
                         0.07963 1056.067 < 2e-16 ***
BlockB2
            -4.45753
                                   -51.103 2.88e-10 ***
                         0.08723
                                    16.162 8.44e-07 ***
x1
             0.93254
                         0.05770
x2
             0.57771
                         0.05770
                                    10.013 2.12e-05 ***
x1:x2
                         0.08159
                                     1.532
                                              0.169
             0.12500
x1^2
            -1.30856
                         0.06006
                                   -21.786 1.08e-07 ***
                                  -15.541 1.10e-06 ***
x2^2
            -0.93344
                         0.06006
```

Signif. codes: 0 âĂŸ***âĂŹ 0.001 âĂŸ**âĂŹ 0.01 âĂŸ*âĂŹ 0.05 âĂŸ.âĂŹ 0.1 âĂŸ âĂŹ 1

```
Residual standard error: 0.1632 on 7 degrees of freedom
```

Multiple R-squared: 0.9981, Adjusted R-squared: 0.9964

F-statistic: 607.2 on 6 and 7 DF, p-value: 3.811e-09

Analysis of Variance Table

```
Response: Yield
```

	Df	Sum Sq	Mean Sq	F value	Pr(>F)
Block	1	69.531	69.531	2611.0950	2.879e-10
FO(x1, x2)	2	9.626	4.813	180.7341	9.450e-07
TWI(x1, x2)	1	0.063	0.063	2.3470	0.1694
PQ(x1, x2)	2	17.791	8.896	334.0539	1.135e-07
Residuals	7	0.186	0.027		
Lack of fit	3	0.053	0.018	0.5307	0.6851
Pure error	4	0.133	0.033		

Stationary point of response surface:

x1 x2 0.3722954 0.3343802

Stationary point in original units:

Time Temp 86.86148 176.67190

Eigenanalysis:

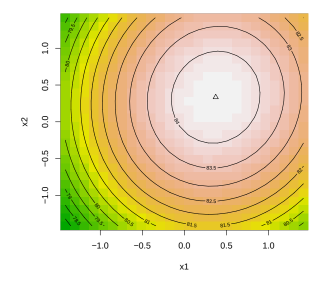
\$values

[1] -0.9233027 -1.3186949

\$vectors

This model fits well. The canonical analysis reveals that the stationary point is near the center of the experiment and that both eigenvalues are negative. This indicates that the fitted surface has a maximum at Time ≈ 86.9 , Temp ≈ 176.7 . We may visualize the response surface using the lm method for contour, provided with this package:

```
> contour(CR.rsm2, x2 ~ x1)
> points(0.372, 0.334, pch = 2)
```



4 Helicopter example

The provided dataset heli is presented in Table 12.5 of Box, Hunter, and Hunter (2005). It is also a central composite design in two blocks. There are four variables and 30 observations altogether. This is a coded.data object already; here are a few observations:

```
> heli[1:4, ]
```

Variable codings ...

```
x1 ~(A - 12.4)/0.6
```

x2 ~(R - 2.52)/0.26

 $x3 \sim (W - 1.25)/0.25$

x4 ~ (L - 2)/0.5

The response variable ave is the average flight time (in csec.) of four test runs each of paper helicopters made with different wing areas W, wing-length ratios R, body widths W, and body lengths L. The goal is to maximize flight time.

Like the Chemical Reaction data, the first block was analyzed first and then the star points were added. We'll skip the first part and go straight to the second-order analysis.

```
> heli.rsm = rsm(ave ~ block + SO(x1, x2, x3, x4), data = heli)
> summary(heli.rsm)

Call:
rsm(formula = ave ~ block + SO(x1, x2, x3, x4), data = heli)
```

```
Residuals:
```

Min 1Q Median 3Q Max -3.850 -1.579 -0.175 1.925 4.200

Coefficients:

```
Estimate Std. Error t value Pr(>|t|)
(Intercept) 372.80000
                        1.50638 247.481 < 2e-16 ***
block2
            -2.95000
                        1.20779 -2.442 0.028452 *
                        0.63656 -0.131 0.897707
x1
            -0.08333
                        0.63656 7.986 1.40e-06 ***
x2
             5.08333
                        0.63656 0.393 0.700429
x3
             0.25000
x4
                        0.63656 -9.557 1.63e-07 ***
            -6.08333
                        0.77962 -3.688 0.002436 **
x1:x2
            -2.87500
x1:x3
            -3.75000
                        0.77962 -4.810 0.000277 ***
x1:x4
             4.37500
                        0.77962 5.612 6.41e-05 ***
x2:x3
             4.62500
                        0.77962 5.932 3.66e-05 ***
x2:x4
            -1.50000
                        0.77962 -1.924 0.074926 .
x3:x4
            -2.12500
                        0.77962 -2.726 0.016410 *
                        0.60389 -3.374 0.004542 **
x1^2
            -2.03750
x2^2
            -1.66250
                        0.60389 -2.753 0.015554 *
                        0.60389 -4.202 0.000887 ***
x3^2
            -2.53750
x4^2
            -0.16250
                        0.60389 -0.269 0.791788
```

Signif. codes: 0 âĂŸ***âĂŹ 0.001 âĂŸ**âĂŹ 0.01 âĂŸ*âĂŹ 0.05 âĂŸ.âĂŹ 0.1 âĂŸ âĂŹ 1

Residual standard error: 3.118 on 14 degrees of freedom

Multiple R-squared: 0.9555, Adjusted R-squared: 0.9078

F-statistic: 20.04 on 15 and 14 DF, p-value: 6.54e-07

Analysis of Variance Table

Response: ave

	Df	Sum Sq	Mean Sq	${\tt F} \ {\tt value}$	Pr(>F)
block	1	16.81	16.81	1.7281	0.209786
FO(x1, x2, x3, x4)	4	1510.00	377.50	38.8175	1.965e-07
TWI(x1, x2, x3, x4)	6	1114.00	185.67	19.0917	5.355e-06
PQ(x1, x2, x3, x4)	4	282.54	70.64	7.2634	0.002201
Residuals	14	136.15	9.72		
Lack of fit	10	125.40	12.54	4.6660	0.075500
Pure error	4	10.75	2.69		

Stationary point of response surface:

x1 x2 x3 x4 0.8607107 -0.3307115 -0.8394866 -0.1161465

Stationary point in original units:

A R W L

12.916426 2.434015 1.040128 1.941927

Eigenanalysis:

\$values

[1] 3.258222 -1.198324 -3.807935 -4.651963

\$vectors

```
[,1] [,2] [,3] [,4]
[1,] 0.5177048 0.04099358 0.7608371 -0.38913772
[2,] -0.4504231 0.58176202 0.5056034 0.45059647
[3,] -0.4517232 0.37582195 -0.1219894 -0.79988915
[4,] 0.5701289 0.72015994 -0.3880860 0.07557783
```

This time, the situation is more complicated. Since the eigenvalues are of mixed sign, we have a saddle point. Here we obtain contour plots of each pair of variables, holding the other two fixed at their stationary values.

```
> par(mfrow = c(2, 3))
> contour(heli.rsm, ~x1 + x2 + x3 + x4, at = summary(heli.rsm)$canonical$xs)
```

The plots are shown in Figure 1. An important thing to note is that when the color underlay is used (as is the default), the color scale is consistent across all plots, facilitating appropriate visual comparisons.

Since we have not found a maximum, our next step might be to experiment along some path that seems promising of providing a higher response. In this particular example, the stationary point is within the experimental region, so we can regard it as reasonably well estimated. It is thus believable that the actual response function has a saddle point in the vicinity of our stationary point. The function canonical.path, by default, returns the path of steepest ascent each direction from the stationary point. This path is linear.

> canonical.path(heli.rsm)

```
dist
            x1
                   x2
                          xЗ
                                 x4 |
                                            Α
                                                    R
                                                                    L |
                                                                           yhat
  -5.0 -1.728
                1.921
                       1.419 -2.967 | 11.3632 3.01946 1.60475 0.5165 | 453.627
2
  -4.5 -1.469
                1.696
                       1.193 -2.682 | 11.5186 2.96096 1.54825 0.6590 | 438.150
3
  -4.0 -1.210
                       0.967 -2.397 | 11.6740 2.90246 1.49175 0.8015
                1.471
                                                                     1 424.302
4
  -3.5 -0.951
                1.246
                       0.742 -2.112 | 11.8294 2.84396 1.43550 0.9440 | 412.094
                       0.516 -1.827 | 11.9848 2.78546 1.37900 1.0865
5
  -3.0 -0.692
                1.021
                                                                     | 401.504
6
  -2.5 - 0.434
                0.795
                       0.290 -1.541 | 12.1396 2.72670 1.32250 1.2295
                                                                        392.534
7
  -2.0 - 0.175
                0.570
                       0.064 -1.256 | 12.2950 2.66820 1.26600 1.3720
                                                                        385.203
  -1.5 0.084
                0.345 -0.162 -0.971 | 12.4504 2.60970 1.20950 1.5145
                                                                        379.502
  -1.0 0.343 0.120 -0.388 -0.686 | 12.6058 2.55120 1.15300 1.6570 | 375.429
10 -0.5 0.602 -0.105 -0.614 -0.401 | 12.7612 2.49270 1.09650 1.7995
                                                                     1 372.986
   0.0 0.861 -0.331 -0.839 -0.116 | 12.9166 2.43394 1.04025 1.9420
                                                                        372.172
                             0.169 | 13.0720 2.37544 0.98375 2.0845 |
12
   0.5 1.120 -0.556 -1.065
                                                                        372.987
13
   1.0 1.378 -0.781 -1.291
                              0.454 | 13.2268 2.31694 0.92725 2.2270 | 375.428
        1.637 -1.006 -1.517
                              0.739 | 13.3822 2.25844 0.87075 2.3695
15
        1.896 -1.232 -1.743
                              1.024 | 13.5376 2.19968 0.81425 2.5120
   2.5 2.155 -1.457 -1.969
                              1.309 | 13.6930 2.14118 0.75775 2.6545 | 392.538
```

```
17
         2.414 -1.682 -2.195
                               1.594 | 13.8484 2.08268 0.70125 2.7970 | 401.498
    3.0
         2.673 -1.907 -2.421
                                       14.0038 2.02418 0.64475 2.9395
18
    3.5
                               1.879
                                                                         412.088
                                       14.1592 1.96568 0.58850 3.0820
         2.932 -2.132 -2.646
                               2.164 |
                                                                         424.295
19
    4.0
         3.190 -2.358 -2.872
                               2.449 | 14.3140 1.90692 0.53200 3.2245
20
    4.5
                                                                       | 438.140
         3.449 -2.583 -3.098
                               2.734 | 14.4694 1.84842 0.47550 3.3670
21
    5.0
```

We should conduct additional experimental runs along this path and see where we get the most improvement in the observed response.

Had the stationary point been more distant, it would be more of an extrapolation from the range of the experiment, and thus it would not be a good starting point for further experimentation. That is, for a distant stationary point, a steepest-ascent method makes more sense. For second-order surfaces, the steepest function uses ridge analysis to determine an appropriate path:

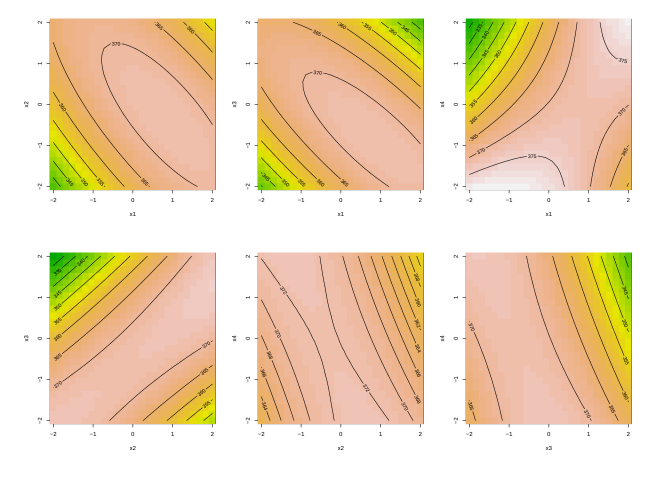


Figure 1: Contour plots for heli data.

> steepest(heli.rsm)

```
Path of steepest ascent from ridge analysis:
                  x2
                        x3
                           0.000 | 12.4000 2.52000 1.25000 2.0000 | 372.800
   0.0 0.000 0.000 0.000
   0.5 -0.127 0.288 0.116 -0.371 | 12.3238 2.59488 1.27900 1.8145 | 377.106
   1.0 -0.351 0.538 0.312 -0.700 | 12.1894 2.65988 1.32800 1.6500 | 382.675
   1.5 -0.595 0.775 0.526 -1.009 | 12.0430 2.72150 1.38150 1.4955 | 389.783
   2.0 -0.846 1.007 0.745 -1.309 | 11.8924 2.78182 1.43625 1.3455 | 398.485
5
   2.5 -1.101 1.237 0.966 -1.605 | 11.7394 2.84162 1.49150 1.1975 | 408.819
6
7
   3.0 -1.356 1.465 1.189 -1.897 | 11.5864 2.90090 1.54725 1.0515 | 420.740
   3.5 -1.613 1.693 1.413 -2.188 | 11.4322 2.96018 1.60325 0.9060 | 434.322
   4.0 -1.870 1.920 1.637 -2.477 | 11.2780 3.01920 1.65925 0.7615 | 449.497
   4.5 -2.127 2.147 1.862 -2.766 | 11.1238 3.07822 1.71550 0.6170 | 466.323
11 5.0 -2.385 2.373 2.086 -3.054 | 10.9690 3.13698 1.77150 0.4730 | 484.750
```

This gives a path that starts at the *origin* in the coded variables, rather than the stationary point.

5 Miscellaneous notes and examples

5.1 Coded data

Use coded.data as shown in the Chemical reactor example to convert a dataset that has its predictors in raw units. If the dataset is already in coded units, you may embed the coding information using as.coded.data:

```
> dat = expand.grid(t = c(-1, 1), w = -1:1)
> dat = as.coded.data(dat, t ~ (Thickness - 3.5)/0.5, w ~ (Width -
      12)/2)
> dat
   t
1 -1 -1
2
  1 -1
3 -1 0
 1
5 -1 1
  1
Variable codings ...
t \sim (Thickness - 3.5)/0.5
w \sim (Width - 12)/2
> decode.data(dat)
  Thickness Width
1
          3
               10
2
          4
               10
3
          3
               12
```

```
12
4
          4
          3
5
                14
6
          4
                14
> code2val(c(t = -0.5, w = 0.25), attr(dat, "codings"))
Thickness
               Width
     3.25
               12.50
   The design-generation functions ccd and bbd also support coding:
> des = bbd(Finish ~ x1 + x2 + x3, coding = list(x1 ~ (Time - 60)/10,
      x2 \sim (Feedrate - 2.2)/0.4, x3 \sim (Speed - 2000)/250))
> des[1:3, ]
  x1 x2 x3 Finish
3 -1 1 0
                NA
     0 -1
5 -1
                NA
  1 0 -1
                NA
Variable codings ...
x1 \sim (Time - 60)/10
x2 ~ (Feedrate - 2.2)/0.4
x3 \sim (Speed - 2000)/250
> decode.data(des[1:3, ])
  Time Feedrate Speed Finish
3
    50
             2.6
                  2000
                            NA
5
    50
             2.2
                  1750
                            NA
6
    70
             2.2
                  1750
                            NA
```

5.2 Contour plots

The contour method provided by this package works for any lm object, not just response surfaces. By default, it overlays the contour plot on an image plot using terrain colors. Arguments provide for the image portion to be disabled or the colors changed if desired.

To make contour work, it was necessary to obtain the data used by a lm object. The standard function get_all_vars does not make it very easy, and model.frame incorporates transformations and expands polynomials and factors. The provided function model.data makes it very easy to obtain just the variables included in the model formula. For example, following the first-order model for the chemical reactor example,

```
> model.data(CR.rsm1, lhs = TRUE)
  Yield x1 x2
  80.5 -1 -1
1
  81.5 -1
3
  82.0
         1 -1
  83.5
4
         1
5
  83.9
         0
            0
6
            0
  84.3
         0
7
   84.0
        0
            0
```

References

Box, G.E.P., Hunter, J.S., and Hunter, W.G. (2005), Statistics for Experimenters: Design, Innovation, and Discovery (2nd ed.), New York: Wiley-Interscience.

Myers, R. H. and Montgomery, D. C. (2002), Response Surface Methodology: Process and Product Optimization Using Designed Experiments (2nd ed.), New York: Wiley-Interscience.

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