

DALL: Davidon's Algorithm for Log Likelihood Maximization — A subroutine for statistical model builders —

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Abstract: A subroutine for the log likelihood maximization is proposed. It is an implementation of Davidon's variance algorithm with a numerical derivative evaluation procedure. It is designed for use of statistical model builders. It is originally coded in FORTRAN. Now C version is available. Numerical examples are given.

Key Words: Statistical modeling, log likelihood, AIC, Numerical optimization, Davidon's variance algorithm, Numerical differentiation, MLE, FORTRAN, parallel computation, Open Market Licence

1 Introduction

By the introduction of AIC (Akaike, 1973) it is recognized that many important statistical problems could be formulated as model selection problems. Necessary statistical modeling is the task not only of expert statisticians, but of all those who concern with the data analysis. Here the log likelihood maximization is the key operation. Typical steps for the statistical problem solving are as follows:

1. model building
2. log likelihood maximization
3. AIC computation and model selection
4. interpretation of the results.

The hindrance there, if any, is the difficulty of numerical maximization of the likelihood.

Numerical procedures should be used for the maximization. Traditional statistical procedures seem to be designed so that the computing cost will be as low as possible. However, taking into account the recent development of computing technology, we need not be afraid of some computation. We may relax our attachment to the analytical methods, so that we will be able to concentrate our attention more on the physical aspect of data.

The optimization procedure should be one that works without user supplied gradient evaluation algorithm. The run-time cost could be reduced by the gradient evaluation algorithm. However, it is usual that the necessary effort to write down the algorithm is substantial and it is not rare that the obtained algorithm contradicts the algorithm for the calculation of the function value. When the analysts is groping for a good model to describe a given phenomenon, the expectation of the possible trouble often suppresses the desire to develop some better ideas.

The purpose of the present article is to propose one working procedure which meets the above stated requirements. We do not claim that ours is the best of all the optimization techniques in the literature. We do not even try to compare the performances of methods. We just make public our procedure and encourage every data analyst to participate in the model building game. Those who are interested in other methods are referred to, for example, Douke & Asano (1989), Kowalik & Osborne (1968), Künzi et al. (1968), Lavi & Vogl (1966) and Powell (1981).

2 DALL

Read "DALL" so that it rhymes with "call". Following the above reasoning, we proposed, in 1996, a FORTRAN subroutine DALL for numerical maximization of the log likelihood function which was based on Davidon's algorithm (Davidon, 1968) and numerical differentiation. The choice of the Davidon's algorithm was based on the experience of the second author in the implementation of the maximum likelihood procedure for ARMA time series model in TIMSAC-74 (Akaike et al., 1985). The subroutine BILL for Davidon's algorithm is a FORTRAN77 version of the maximization procedure in TIMSAC-74. The construction and coding of DALL was mainly done by the first author. Now we propose a new FORTRAN version and an equivalent C version. The flow chart of DALL is given as Figure 1.

2.1 Davidon's Algorithm

Assume that a function ϕ has the form

$$\phi(\mathbf{x}) = \phi_0 + \frac{1}{2}(\mathbf{x} - \mathbf{x}_0)^T V_0^{-1}(\mathbf{x} - \mathbf{x}_0), \quad (1)$$

where V_0^{-1} is a negative definite matrix, then the relation between the gradient at a point \mathbf{x} and the maximizing point is given by

$$\mathbf{g} = V_0^{-1}(\mathbf{x} - \mathbf{x}_0). \quad (2)$$

That is, if the variance V_0 of the function (1) is known and the gradient at a point is known the maximizing point is readily calculated by

$$\mathbf{x} - V_0 \mathbf{g}. \quad (3)$$

On the other hand, if gradients at sufficient number of points are known V_0 can be estimated. Assuming that the object function is well approximated by a quadratic function, Davidon's variance algorithm (Davidon, 1968), a typical quasi-Newton method, is constructed on these two facts.

Notes to Figure 1

- (1) The gradient is computed numerically by

$$g_i = \frac{\phi(\mathbf{x} + \text{step}(i) \mathbf{e}_i) - \phi(\mathbf{x} - \text{step}(i) \mathbf{e}_i)}{2 \text{step}(i)} \quad (i = 1, 2, \dots, IP) \quad (4)$$

where \mathbf{e}_i is the unit vector of the i -th axis and IP is the dimension of the parameter vector $\boldsymbol{\theta}$. This computation can (and should, if possible) be executed parallelly.

- (2) The difference between *SPHI* and the maximum value is given by

$$\frac{1}{2}(\mathbf{x}_* - \mathbf{x}_0)^T V_0^{-1}(\mathbf{x}_* - \mathbf{x}_0),$$

which is equal to

$$\frac{1}{2} \mathbf{g}_*^T V_0 \mathbf{g}_*. \quad (5)$$

RHO is the twice of an estimate of this value. It is implicitly assumed that the variance V_n is sufficiently close to V_0 .

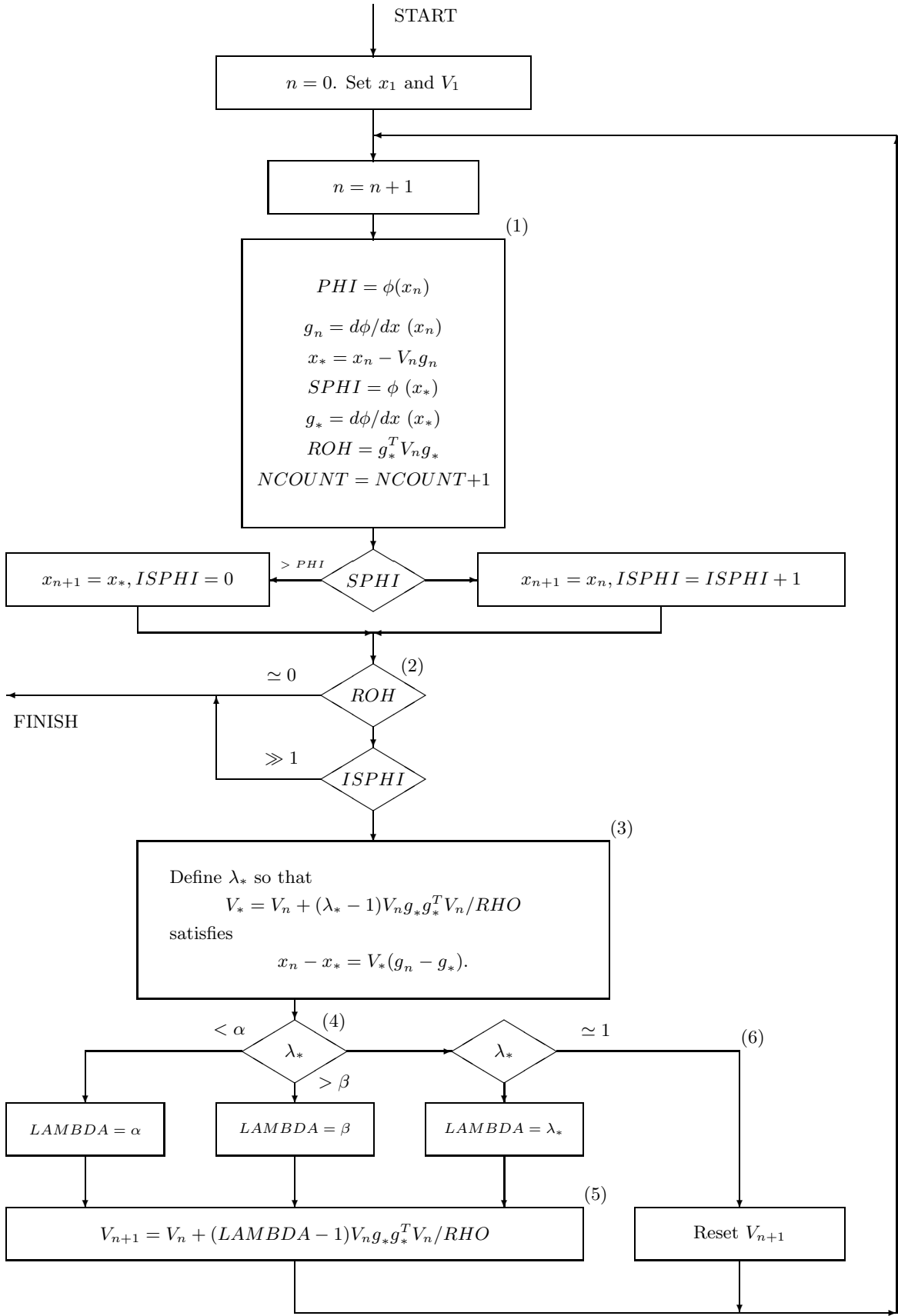


Figure 1. Flow chart of DALL

(3) Under the present assumptions, if $V_n = V_0$ then $\mathbf{g}_* = 0$ and the procedure ends. The fact that $\mathbf{g}_* \neq 0$ is an evidence of $V_n \neq V_0$. If (1) is the correct form, V_0 satisfies

$$\mathbf{x}_n - \mathbf{x}_* = V_0(\mathbf{g}_n - \mathbf{g}_*). \quad (6)$$

Define $\gamma = -\mathbf{g}_n^T V_n \mathbf{g}_* / RHO$, then $\lambda_* = |\gamma / (\gamma + 1)|$.

- (4) If $LAMBDA = 1$, V_n is not improved. If $LAMBDA = 1$ is far from 1 the modification might be too drastic, especially if the assumed form (1) of the function is dubious.

The careful choice of α and β , which appear in step 3.(c), is important. The choice is discussed by Davidon(1968) and $\alpha = 10^{-3}$ and $\beta = 10$ are suggested as reasonable choices. However, based on our own experience of the use of the procedure we set α and β equal to 0.25 and 4.0, respectively.

- (5) Generally, the shape of a K -dimensional matrix A is known by calculating $\{A\mathbf{b}_1, A\mathbf{b}_2, A\mathbf{b}_3, \dots, A\mathbf{b}_K\}$, where $\{\mathbf{b}_1, \mathbf{b}_2, \mathbf{b}_3, \dots, \mathbf{b}_K\}$ is an orthogonal system of K vectors. If we choose the basis so that $\mathbf{b}_1 = V_n \mathbf{g}_*$, we have relations

$$V_{n+1} \mathbf{b}_j = V_n \mathbf{b}_j \quad (j = 2, \dots, K). \quad (7)$$

- (6) This path is not a part of Davidon's original algorithm.

2.2 DALL

Explanations hereafter are for the FORTRAN version. It should not be too difficult to translate the following text into words of C.

Usage Prepare

```
subroutine model(a,ip,el,lel)
```

to compute the log likelihood, initial guess 'a' of parameter and step sizes step(1), step(2), ..., step(ip) for numerical differentiation, and

```
call dall (model,a,ip,el,g vd,encd,step,limit,lpsw),
```

then the variable 'el' gives the maximum log likelihood. The specification for the make of 'model' is as follows:

- The number of free parameters is given in 'ip'.
- Parameter values are set in 'a'.
- Set log likelihood value in 'el'.
- Set 'lel'=0, if given 'a' is out of domain of the log likelihood. Otherwise set 'lel'=1.

The meaning of other arguments are summarized in Table 1.

Table 1 Arguments of DALL

Argument	Meaning
model	input. subroutine name.
a(ip)	input/output. parameters.
ip	input. number of free parameter.
el	output. maximum log likelihood.
g(ip)	output. final gradient estimate.
vd(ip,ip+5)	output/work area.
encd	output. end code.
step(ip)	input. step size vector for the numerical differentiation.
limit	input. NCOUNT limit. limit=0 indicates no limit.
lpsw	input. output level control (see Table 3)

Outputs Arguments for outputs are picked up in Table 2. Table 3 summarizes output control by variable ‘lpsw’.

Table 2 Results of DALL

Argument	Contents
a(ip)	estimated parameters.
el	maximum loglikelihood estimation.
g(ip)	final gradient estimate.
vd(ip,ip)	final inverse Hessian estimate.
endc	end code. NCOUNT + LCODE/100. see Table 4.

Table 3 Output control

lpsw	Output
0	No print out, with exceptions of fatal errors.
1	Summary print out only
2	Minimum report from the outer loop of the procedure
3	Full report from the inner loop of the procedure

When ‘lpsw’ is greater than 1, “grobal profile of the log likelihood” is given. This ‘graph’ shows the profile of the log likelihood along the bee-line from the initial point to the final point.

When ‘lpsw’ is set equal to 0, user has to check ‘endc’ variable to know the status. ‘endc’ is defined by

$$\text{endc} = \text{NCOUNT} + \text{LCODE}/100$$

where LCODE has the meaning summarized in Table 4.

Table 4 LCODE

lcode	Meaning
0	Davidon’s algorithm ended successfully with $RHO < \varepsilon$
50	NCOUNT reached given limit. Computation is not completed.
70	Davidon’s algorithm reached dead end with $LAMBDA < \lambda_{\text{lim}}$
71	Davidon’s algorithm reached dead end with $ISPHI = ISPHI_{\text{lim}}$
72	\mathbf{x}_* can’t be evaluated.
85	Bad initial guess! ‘model’ can’t be evaluated at the initial point \mathbf{x}_1 .
90	Setting of lel variable in ‘model’ is incorrect.
95	Step vector has zero element. Numerical differentiation impossible.

3 Numerical Examples

3.1 One dimensional optimization

This example gives a side view of the performance of DALL. Assume that we conducted coin tossing n times and got m times “top”. Then the log likelihood function of the probability p of getting “top” is given by

$$\ell(p) = m \log p + (n - m) \log(1 - p). \quad (8)$$

Actually MLE is analytically calculated to be $\hat{p} = m/n$.

The example is for the case $n = 12$, $m = 76$ and then $\hat{p} = 0.158$. We can see how Dall finds the maximum value starting from 0.9 as the initial guess in Fig.2. It is seen that quadratic function model

$$\phi(\mathbf{x}) = \phi_* + \frac{1}{2}(\mathbf{x} - \mathbf{x}_*)^T V_n^{-1}(\mathbf{x} - \mathbf{x}_*). \quad (9)$$

fitted to the data is improved gradually. The position of the tangent point of the graph of $\ell(p)$ and the graph of the model is $\mathbf{x}_{\text{NCOUNT}}$. Fig.3 and Fig.4 show the program and its output, respectively.

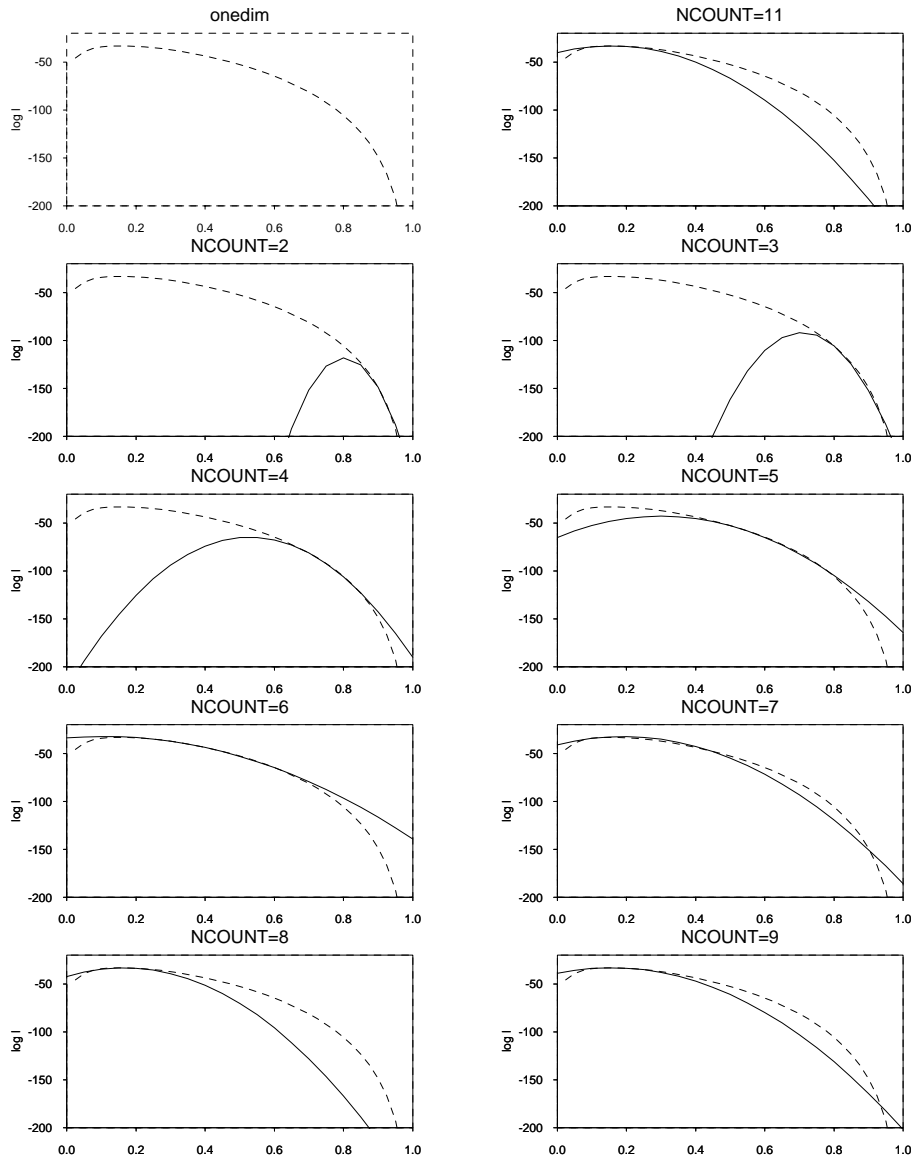


Figure 2 Numerical maximization of the log likelihood of a binomial model.

```

implicit real*8 (a-h,o-z)
parameter (ipmax=5)
dimension g(ipmax), vd(ipmax,ipmax+5)
dimension a1(ipmax),step(ipmax)
common /cmodel/ m,m1
external model0

m = 76
m1 = 12
do 1 i=1,ipmax
1  step(i) = 0.001d0

ip1=1
a1(1) = 0.9d0
call dall(model0,a1,ip1,e1,g,vd,endcode,step,0,3)
stop
end

subroutine model0(par,np,e1,l1)
implicit real*8 (a-h,o-z)
common /cmodel/ m,m1
dimension par(np)
l1 = 0
di = par(1)
if(di .le. 0.d0) return
if(di .ge. 1.d0) return

```

```

f = m1 * dlog(di) + (m-1) * dlog(1.d0-di)
lel=1
return
end

```

Figure 3. Program for binomial model fitting.

```

D1:===== log likelihood maximization =====
D1:number of processor = 1
D1:limit = 0
D2:Hessian reset count = 0
D2:value= -148.63
D2:point
0.90000D+00
B3: NCOUNT PHI SPHI-PHI RHO LAMBDA-1 ISPHI
B3: 2 -148.63 42.2388 -14.8632 0.97 0
B3: 3 -106.39 23.5797 -12.5120 1.89 0
B3: 4 -82.81 27.0428 -11.3286 1.27 0
B3: 5 -55.77 18.4616 -5.3661 0.84 0
B3: 6 -37.31 3.3336 -5.3506 -0.42 0
B3: 7 -33.97 0.5519 -0.5487 -0.30 0
B3: 8 -33.42 0.2532 -0.0321 0.40 0
B3: 9 -33.17 0.0192 -0.0012 -0.14 0
B2:Bill ended with abs(log LAMBDA)= 0.15 < 0.20 = ramlim: code = 70
D2:val. dif.= 115.48
D2:NCOUNT= 9
D2:Hessian reset count = 1
D2:value= -33.15
D2:point
0.15657D+00
B3: NCOUNT PHI SPHI-PHI RHO LAMBDA-1 ISPHI
B3: 11 -33.15 0.0005 0.0000 0.0000 0
B2:Bill ended with abs(RHO) = 0.00000 < 0.00010 = rhomin: code = 0
D2:val. dif.= 0.00
D2:NCOUNT= 11
D1:Bill ended with abs(RHO) < rhomin.
D2:===== Summary =====
D1:value= -33.15
D1:point
0.15789D+00
D2:final gradient
0.51524D-02
D1:val. dif.= 115.48
D2:global profile
-150.00 -125.00 -100.00 -75.00 -50.00 -25.00 0.00
1 | | | | | | |
2 | *XXXXXX | | | | | | |
3 | XXXXXX*XXXX | | | | | | |
4 | XXXX*XX | | | | | | |
5 | XX*XX | | | | | | |
6 | XX*X | | | | | | |
7 | X*X | | | | | | |
8 | X* | | | | | | |
9 | * | | | | | | |
10 | * | | | | | | |
11 | * | | | | | | |
D1:Hessian reset count = 1
D1:end code =11.00
D1:===== end of maximization =====

```

Figure 4. A result of binomial model fitting.

3.2 Two-dimensional Optimization

This example gives a ‘bird’s eye view’ of the performance of DALL. The following function is an artificial function. Not a real log likelihood function.

$$f(x, y) = -\frac{1}{2}\{r(x^2 + x + y)^2 + (x^2 + 3x + y)^2\} \quad (10)$$

Here, $r = 40.0$. The function takes the maximum value 0.0 at point B(0,0) shown in the top-left panel of Fig.5-1. The initial guess is taken at point A(-2.5, -3.0). Two contour curves in dotted line at 1.0 and 10.0 below the highest point are drawn in the figure.

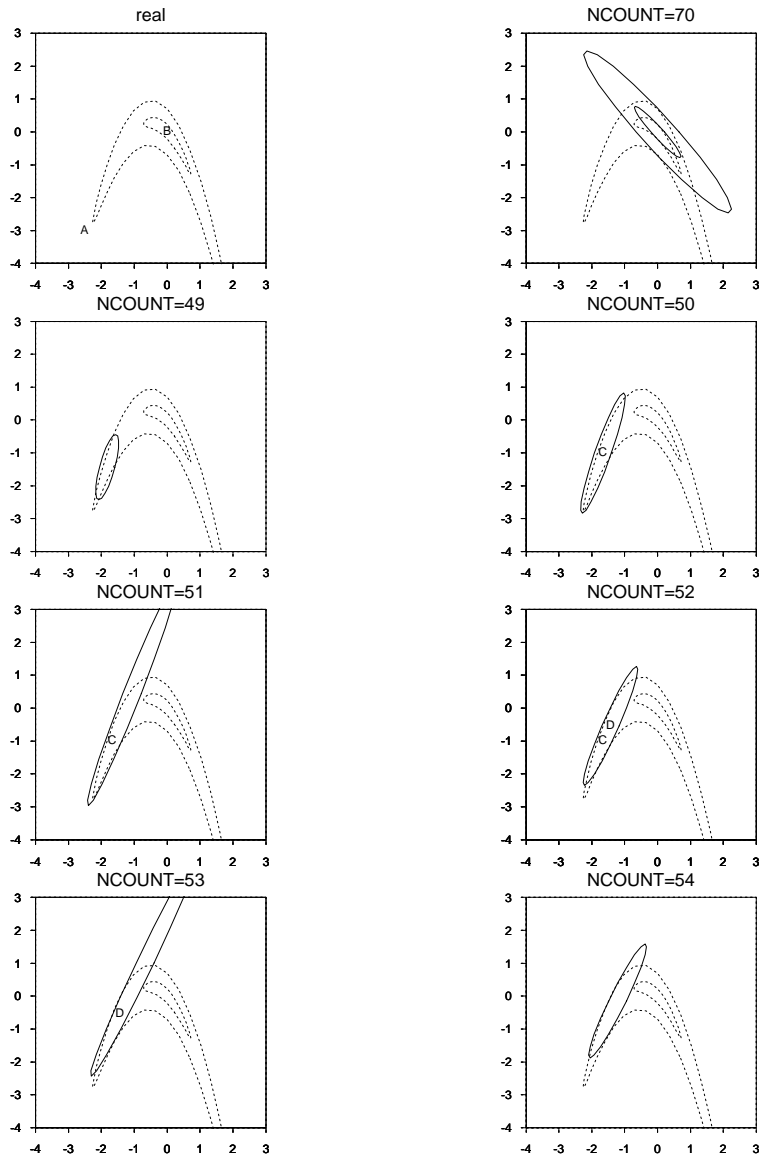


Figure 5-1. Search process by Davidon's algorithm.

Character 'C' in the panel titled $NCOUNT = 51$, indicates the position of the point \mathbf{x}_{51} . The solid oval is the contour line at the level of $\phi_* - 10.0$ of the model(9) estimated based on information obtained up to this moment. The function value at point \mathbf{x}_* , the center of the oval, is less than the value at \mathbf{x}_{51} and it turned out that this model is not good and \mathbf{x}_{52} is set equal to \mathbf{x}_{51} . The model is modified to get one shown in the figure titled $NCOUNT = 52$, the function value at $\mathbf{x}_*(D)$ is larger than that at \mathbf{x}_{52} and \mathbf{x}_{53} is set equal to \mathbf{x}_* . Iterating this process (see Figs. 5-2 and 5-3), we get the result shown in the figure titled $NCOUNT=70$ at top-right panel. Two ovals in solid line in the figure are contour lines at levels of $\phi_* - 1.0$ and $\phi_* - 10.0$. It is clear that fitting of the last model is poor globally in this case. But the local structure of the object function is captured by the model.

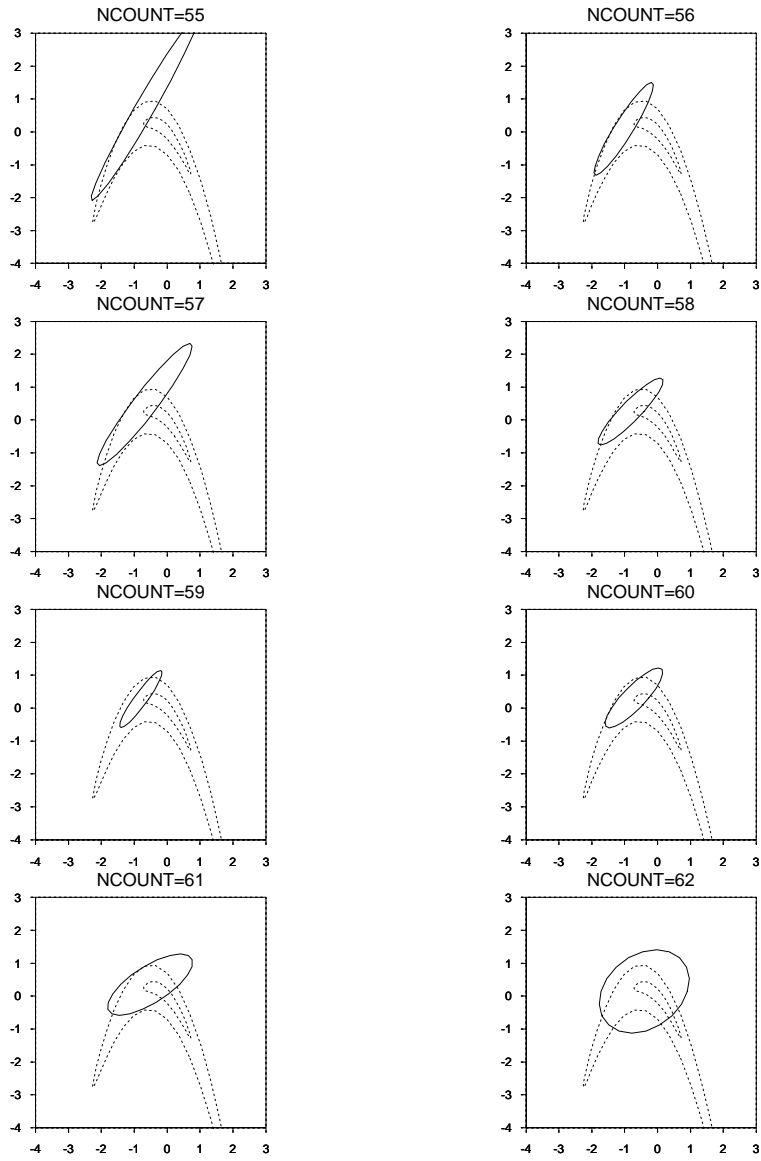


Figure 5-2. (continued)

The print out of DALL is shown in Figure 3. The “graph” composed of * and X at the end of the figure shows profile of the object function along the line from point A to B. Since this “graph” does not show the function values at points $\mathbf{x}_1, \mathbf{x}_2, \dots$, it does not increase monotonically. Seeing this “graph” we can get some information if the object function has an easy shape or not.

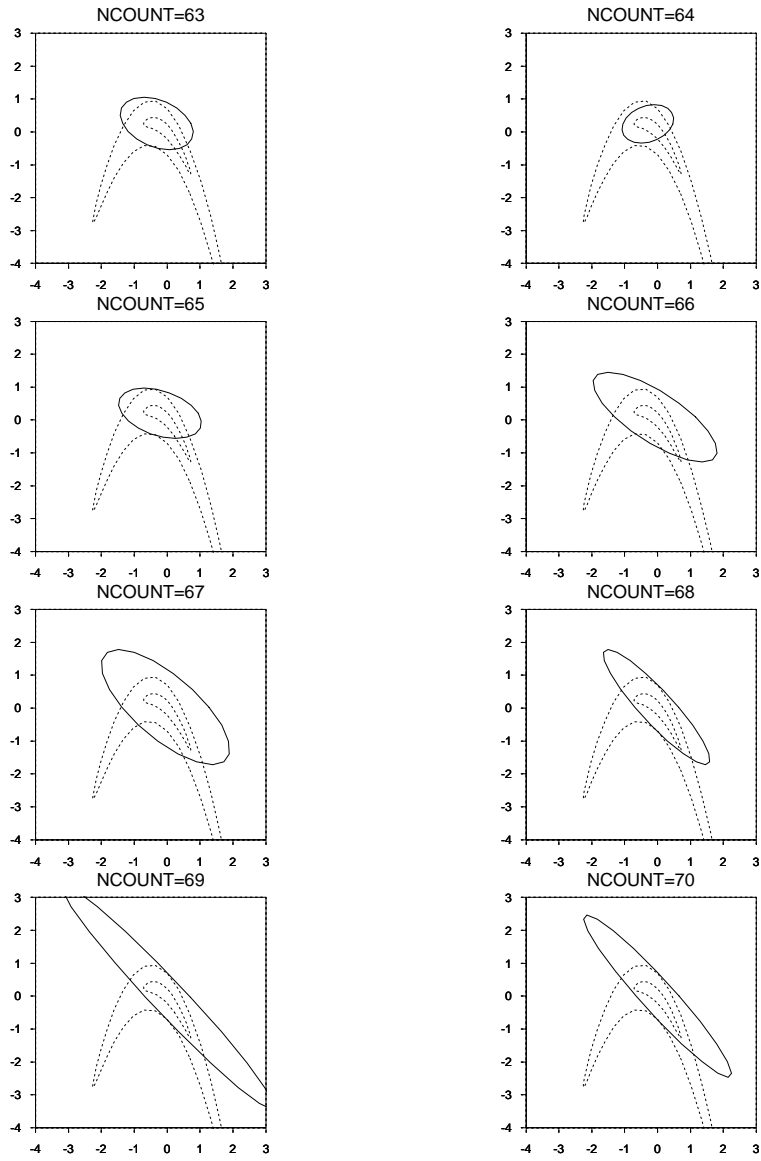


Figure 5-3. (continued)

```

D2:Hessian reset count = 11
D2:value= -7.60
D2:point
-0.19124D+01 -0.15012D+01
B3: NCOUNT PHI SPHI-PHI RHO LAMBDA-1 ISPHI
B3: 47 -7.60 0.1857 -2.2604 -0.50 0
B3: 48 -7.41 0.6270 -0.0819 2.73 0
B3: 49 -6.79 0.2991 -0.2897 3.00 0
B3: 50 -6.49 1.1112 -0.9496 3.00 0
B3: 51 -5.38 -1.0834 -49.6538 -0.75 0
B3: 52 -5.38 1.3701 -1.0523 3.00 0
B3: 53 -4.01 -8.7070 -39.8665 -0.75 0
B3: 54 -4.01 1.0407 -0.5745 2.85 0
B3: 55 -2.97 -2.2384 -73.4456 -0.75 1
B3: 56 -2.97 0.7719 -0.4114 2.26 0
B3: 57 -2.19 0.2893 -4.6355 -0.75 0
B3: 58 -1.90 0.4668 -0.3263 -0.75 0
B3: 59 -1.44 0.0776 -0.0683 3.00 0
B3: 60 -1.36 0.2411 -0.1684 3.00 0
B3: 61 -1.12 0.4728 -0.2189 3.00 0
B3: 62 -0.65 0.0453 -3.1124 -0.75 0
B3: 63 -0.60 0.3932 -0.0996 -0.71 0
B3: 64 -0.21 0.0271 -0.0233 3.00 0
B3: 65 -0.18 0.0763 -0.0422 2.51 0
B3: 66 -0.10 0.0882 -0.0166 0.92 0
B3: 67 -0.02 0.0025 -0.0915 -0.75 0
B3: 68 -0.01 0.0123 -0.0015 3.00 0
B3: 69 0.00 -0.0001 -0.0065 -0.52 0
B3: 70 0.00 0.0014 0.0000 0.0000 0
B2:Bill ended with abs(RHO) = 0.00003 < 0.00010 = rhomin: code = 0

```

```

D2:val. dif.=          7.60
D2:NCOUNT=          70
D1:Bill ended with abs(RHO) < rhomin.
D2:===== Summary =====
D1:value=          0.00
D1:point
      -0.97975D-06 -0.80570D-03
D2:final gradient
      0.34650D-01  0.33076D-01
D1:val. dif.=          20.28
D2:global profile
      -40.00      -30.00      -20.00      -10.00      0.00      10.00
      1 |-----|-----|-----|-----|-----|-----|
      2 |          |          |          |          |          |          |
      3 |          |          |          |          |          |          |
      4 |          |          |          |          |          |          |
      5 |          |          |          |          |          |          |
      6 |          |          |          |          |          |          |
      7 |          |          |          |          |          |          |
      8 |          |          |          |          |          |          |
      9 |          |          |          |          |          |          |
     10 |          |          |          |          |          |          |
     11 |          |          |          |          |          |          |
D1:Hessian reset count = 11
D1:end code =70.00
D1:===== end of maximization =====

```

Figure 6 Output of DALL

3.3 Constrained Gaussian model fitting

Let x_1, \dots, x_n be data. The log likelihood of a normal distribution model with respect to the data is given by

$$l_x(\mu_x, \sigma_x^2) = -\frac{n}{2} \log 2\pi - \frac{n}{2} \log \sigma_x^2 - \frac{1}{2\sigma_x^2} \sum_{i=1}^{n_x} (x_i - \mu_x)^2. \quad (11)$$

For data y_1, \dots, y_n ;

$$l_y(\mu_y, \sigma_y^2) = -\frac{n}{2} \log 2\pi - \frac{n}{2} \log \sigma_y^2 - \frac{1}{2\sigma_y^2} \sum_{i=1}^{n_y} (y_i - \mu_y)^2. \quad (12)$$

MLE's of μ_x and σ_x^2 are

$$\hat{\mu}_x = \frac{1}{n} \sum_{i=1}^n x_i$$

and

$$\hat{\sigma}_x^2 = \frac{1}{n_x} \sum_{i=1}^{n_x} (x_i - \hat{\mu}_x)^2,$$

Similar expression for μ_y, σ_y^2 can be given.

For data $x = \{0.73, -0.06, 1.04, 2.29, 0.51, -0.45, 1.03, 0.44, 0.02, 0.11, -2.42\}$ MLE's are

$$\begin{aligned} \hat{\mu}_x &= 0.2945 \\ \hat{\sigma}_x^2 &= 1.2267 \end{aligned}$$

and MLE's for data $y = \{0.10, 0.56, -1.11, -0.48, 3.46, -2.39, 0.36, 4.56\}$ are

$$\begin{aligned} \hat{\mu}_y &= 0.6325 \\ \hat{\sigma}_y^2 &= 4.6491. \end{aligned}$$

The log likelihood for the model with the constraint $\mu_x = \mu_y$ is given by

$$\begin{aligned} l_{x+y}(\mu_{x+y}, \sigma_x^2, \sigma_y^2) &= -\frac{n}{2} \log 2\pi - \frac{n}{2} \log \sigma_x^2 - \frac{1}{2\sigma_x^2} \sum_{i=1}^{n_x} (x_i - \mu_{x+y})^2 \\ &\quad - \frac{n}{2} \log 2\pi - \frac{n}{2} \log \sigma_y^2 - \frac{1}{2\sigma_y^2} \sum_{i=1}^{n_y} (y_i - \mu_{x+y})^2 \\ &= l_x(\mu_{x+y}, \sigma_x^2) + l_y(\mu_{x+y}, \sigma_y^2). \end{aligned} \quad (13)$$

The problem of analytical maximization of the function is equivalent to solving an algebraic equation of order three. Instead, let DALL solve the problem. Prepare subroutines shown in Fig.7.

Subroutines llx, lly and llxy, are composed so that they compute log likelihood for parameter value specifies by the array p of length m. As eq.(13) show the subroutin lly is composed so that it calls llx and lly. The main routine and results aare shown as Figures 8 and 9, respectively. From the resultant relation

$$AIC_x + AIC_y = 37.46 + 39.00 = 76.46 > 74.62 = AIC_{xy}$$

supports the assumption $\mu_x = \mu_y$.

```

subroutine llx(p,m,el,lel)
implicit real*8 (a-h,o-z)
dimension p(m)
common /i721/ n
common /r721/ x(11)
data pi /3.14159265d0/
lel=0
if(p(2) .gt. 0.d0) then
  sum=0.d0
  do 1 i=1,n
    sum=sum+(x(i)-p(1))**2
1  continue
  el= -n*0.5*( dlog(pi*2)+dlog(p(2)) ) - 0.5*sum/p(2)
  lel=1
end if
return
end

c
subroutine lly(p,m,el,lel)
implicit real*8 (a-h,o-z)
dimension p(m)
common /i722/ n
common /r722/ x(8)
data pi /3.14159265d0/
lel=0
if(p(2) .gt. 0.d0) then
  sum=0.d0
  do 1 i=1,n
    sum=sum+(x(i)-p(1))**2
1  continue
  el= -n*0.5*( dlog(pi*2)+dlog(p(2)) ) - 0.5*sum/p(2)
  lel=1
end if
return
end

c
subroutine llxy(p,m,el,lel)
implicit real*8 (a-h,o-z)
dimension p(m),px(2),py(2)
px(1)=p(1)
px(2)=p(2)
call llx(px,2,elx,lel)
if(lel .eq. 0) return
py(1)=p(1)
py(2)=p(3)
call lly(py,2,ely,lel)
if(lel .eq. 0) return
el=elx+ely
return
end

c
block data ex72
implicit real*8 (a-h,o-z)
common /i721/ n1
common /r721/ x1(11)
common /i722/ n2
common /r722/ x2(8)
data n1/11/
data x1/ 0.73, -0.06, 1.04, 2.29, 0.51,
* -0.45, 1.03, 0.44, 0.02, 0.11,
* -2.42 /
data n2/8/
data x2/ 0.10, 0.56, -1.11, -0.48, 3.46,
* -2.39, 0.36, 4.56 /
end

```

Figure 7. Subroutines to compute log likelihood of Gaussian models and constrained Gaussian model.

```

parameter (ipmax=4)
implicit real*8( a-h,o-z )

```

```

dimension ax(ipmax),ay(ipmax),axy(ipmax)
dimension g(ipmax), vd(ipmax,ipmax+5),step(ipmax)
external llx,lly,llx
c
myid = 0
do 1 i=1,ipmax
1 step(i) = 0.001d0
  ipx=2
  ax(1) = 0.d0
  ax(2) = 1.d0
  call dall(llx,ax,ipx,el,g,vd,encd,step,0,3)
  if(myid .eq. 0) then
    write(6,*) ' meanx(mle)=',ax(1)
    write(6,*) ' variancex(mle)=',ax(2)
    write(6,*) ' maximum likelihood=',el
    write(6,*) ' number of parameter=',ipx
    aic= -2*el + 2*ipx
    write(6,'('' AICx='',F10.2)') aic
  end if
  ipy=2
  ay(1) = 0.d0
  ay(2) = 1.d0
  call dall(lly,ay,ipy,el,g,vd,encd,step,0,3)
  if(myid .eq. 0) then
    write(6,*) ' meany(mle)=',ay(1)
    write(6,*) ' variancex(mle)=',ay(2)
    write(6,*) ' maximum likelihood=',el
    write(6,*) ' number of parameter=',ipy
    aic= -2*el + 2*ipy
    write(6,'('' AICy='',F10.2)') aic
  end if
  ipxy=3
  axy(1) = (ax(1) + ay(1))*0.5d0
  axy(2) = ax(2)
  axy(3) = ay(2)
  call dall(llxy,axy,ipxy,el,g,vd,encd,step,0,3)
  if(myid .eq. 0) then
    write(6,*) ' meanxy(mle)=',axy(1)
    write(6,*) ' variancedx1(mle)=',axy(2)
    write(6,*) ' variancex(mle)=',axy(3)
    write(6,*) ' maximum likelihood=',el
    write(6,*) ' number of parameter=',ipxy
    aic= -2*el + 2*ipxy
    write(6,'('' AICxy='',F10.2)') aic
  end if
stop
end

```

Figure 8. A main program to fit Gaussian distribution model and constrained Gaussian distribution model.

```

D1:===== log likelihood maximization =====
D1:number of processor = 1
D1:limit = 0
D2:Hessian reset count = 0
D2:value = -17.33
D2:point
0.00000D+00 0.10000D+01 SPHI-PHI RHO LAMBDA-1 ISPFI
B3: NCOUNT 2 -17.33 0.5980 -0.0019 0.08 0
B2:Bill ended with abs(log LAMBDA) = 0.08 < 0.20 = ramlim: code = 70
D2:val. dif. = 0.60
D2:NCOUNT = 2
D2:Hessian reset count = 1
D2:value = -16.73
D2:point
0.29455D+00 0.11927D+01 SPHI-PHI RHO LAMBDA-1 ISPFI
B3: NCOUNT 5 -16.73 0.0022 0.0000 0.0000
B2:Bill ended with abs(RHO) = 0.00001 < 0.00010 = rhomin: code = 0
D2:val. dif. = 0.00
D2:NCOUNT = 5
D1:Bill ended with abs(RHO) < rhomin.
D2:===== Summary =====
D1:value = -16.73
D1:point
0.29455D+00 0.12249D+01
D2:final gradient
-0.17764D-11 0.67384D-02
D1:val. dif. = 0.60
D2:global profile
-17.50 -17.25 -17.00 -16.75 -16.50 -16.25 -16.00
  || | | | | | | | | | | |
  || *X | | | | | |
  || XX* | | | | | |
  ||  | | | | | |
  || | | X*X | | | | | |
  || | | | | | | | |
  || | | | X*X | | | | | |
  || | | | | | X* | | | | | |
  || 5 | | | | | | | |

```

```

6
10
11
D1:Hessian reset count = 1
D1:end code = 4.00
D1:==== end of maximization =====
meanx(mle)= 0.29454545454562
variancex(mle)= 1.2248781111300
maximum likelihood= -16.732202448716
number of parameter= 2
AICx= 37.46

```

```

D1:==== log likelihood maximization =====
D1:number of processor = 1
D1:limit = 0

```

```

D2:Hessian reset count = 0
D2:value= -27.55
D2:point
0.00000D+00 0.10000D+01
B3: NCOUNT PHI SPHI-PHI RHO LAMBDA-1 ISPHI
B3: 2 -27.55 5.8549 -1.0351 0.0861 0
B3: 3 -21.679 1.3469 -0.7011 1.0801 0
B3: 4 -20.332 1.4238 -0.4925 1.0901 0
B3: 5 -18.922 0.7238 -0.2774 1.0901 0
B3: 6 -18.20 0.0 0.1403 0.0 0
B3: 7 -17.76 0.0 0.0503 0.0 0
B3: 8 -17.51 0.0 0.0149 0.0 0
B3: 10 -17.50 0.0015 -0.0001 0.0 0

```

```

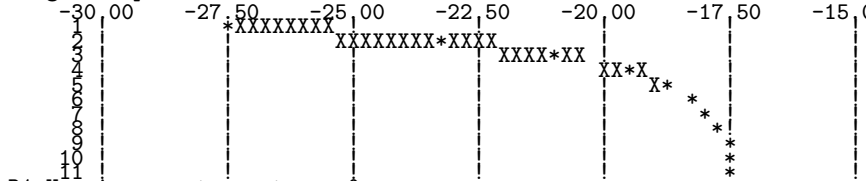
B2:Bill ended with abs(RHO) = 0.00006 < 0.00010 = rhomin: code = 0
D2:val dif.= 10.05
D2:NCOUNT= 10
D1:Bill ended with abs(RHO) < rhomin.
D2:==== Summary =====
D1:value= -17.50
D1:point

```

```

D2:final gradient
0.63250D+00 0.46284D+01
-0.44409D-10 0.38577D-02
D1:val dif.= 10.05
D2:global profile
-30.00 -27.50 -25.00 -22.50 -20.00 -17.50 -15.00

```



```

D1:Hessian reset count = 0
D1:end code = 10.00
D1:==== end of maximization =====
meany(mle)= 0.6325000002579
variancex(mle)= 4.6284090071286
variancex(mle)= 4.7290641711741
maximum likelihood= -17.498215714685
number of parameter= 2
AICy= 39.00

```

```

D1:==== log likelihood maximization =====
D1:number of processor = 1
D1:limit = 0

```

```

D2:Hessian reset count = 0
D2:value= -34.38
D2:point
0.46352D+00 0.12249D+01 0.46284D+01
B3: NCOUNT PHI SPHI-PHI RHO LAMBDA-1 ISPHI
B3: 2 -34.38 0.0698 -0.0023 -0.47 0
B3: 3 -34.31 0.0009 -0.0003 1.09 0
B3: 4 -34.31 0.0003 0.0000 0.0000 0

```

```

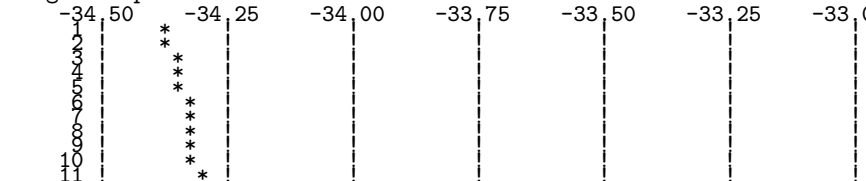
B2:Bill ended with abs(RHO) = 0.00000 < 0.00010 = rhomin: code = 0
D2:val dif.= 0.07
D2:NCOUNT= 4
D1:Bill ended with abs(RHO) < rhomin.
D2:==== Summary =====
D1:value= -34.31
D1:point

```

```

D2:final gradient
0.34827D+00 0.12291D+01 0.47291D+01
0.22816D-04 0.19078D-02 0.14178D-03
D1:val dif.= 0.07
D2:global profile
-34.50 -34.25 -34.00 -33.75 -33.50 -33.25 -33.00

```



```

D1:Hessian reset count = 0
D1:end code = 4.00
D1:==== end of maximization =====
meanxy(mle)= 0.34826773312855
variancex(mle)= 1.2290783247532
variancex(mle)= 4.7290641711741
variancex(mle)= 4.7290641711741
maximum likelihood= -34.312209336033
number of parameter= 3
AIC3= 74.62

```

4 Distributed files

DALL package summarized in Table 5 can be obtained at <http://www.ism.ac.jp/software/ismlib/soft.e.html>

Table 5. ARdock package

file name	contents
README	Latest announcement
example3.f	log likelihood subroutines and main program for the third example and DALL routine in FORTRAN
gauss.c	log likelihood subroutines and main program for the third example in C
dall_pack.c	DALL routine in C
nrutil.h	a header file required by dall_pack.c
Makefile	sample makefile for C version for Unix environment
dall.ps	post script file of this monograph

Note: FORTRAN version DALL routine contained in this file is the one extracted from ARdock package. Those users who are interested in using DALL on a parallel computer are invited to down-load the source code of ARdock package.

ARdock, an Auto-Regressive model analyzer

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(<http://www.ism.ac.jp/software/ismlib/soft.e.html>)

So far, the test of DALL is successfully done in environments summarized in the following Table.

Table 6. Computation Environments

Hardware	Operating System	Compiler
HITACHI SR8000	HI-UX/MPP	FORTRAN90
HP9000V2250	HP-UX 11.0	HP Fortran 90
IBM R/6000 SP	AIX Version 4	AIX XL Fortran
SGI Origin 2000/4-DS-I	IRIX Release 6.4 IP27	FORTRAN90
SPARCstation 5	Sun OS 4.1.4-JLE 1.1.4	g77 version0.5.23
PC-9801 nx/C	MS-DOS 5.00A-H	Microsoft FORTRAN Version 4.01

4.1 Licencing policy

We make public the source code of DALL under the open market licence policy proposed by the Institute of Statistical Mathematics.

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DALL A subroutine for statistical model builders

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(1) <http://www.ism.ac.jp/software/ismlib/soft.e.html>

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