# MINUIT Tutorial 

## Function Minimization

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## 1 Introduction

### 1.1 The motivation

A large class of problems in many different fields of research can be reduced to the problem of finding the smallest value taken on by a function of one or more variable parameters. Examples come from fields as far apart as industrial processing (minimization of production costs) and general relativity (determination of geodesics by minimizing the path length between two points in curved space-time). But the classic example which occurs so often in scientific research is the estimation of unknown parameters in a theory by minimizing the difference ( $\chi^{2}$ ) between theory and experimental data. In all these examples, the function to be minimized is of course determined by considerations proper to the particular field being investigated, which will not be addressed here. The main goal is to study the problem of minimization.

### 1.2 Minimization, maximization and optimization

Although traditionally one speaks of function minimization, some authors refer to maximization. Of course the two are entirely equivalent since one can be converted to the other by changing the sign of the function. Thus the problems of minimizing $\chi^{2}$, maximizing likelihood, minimizing cost, or maximizing efficiency can all be considered as minimization (or maximization). To avoid committing himself, it is now fashionable to speak of optimization, to cover both cases. This unfortunately causes confusion with optimization in control theory where the principle techniques are analytical (calculus of variations) and hence bear little relationship to the numerical methods used in function minimization as treated here.

To add to the confusion there is the term "programming", which is also used to mean minimization (usually specified as linear programming, non-linear programming, or mathematical programming), a historical usage dating from the time when programmers in the modern sense did not exist, and computer users were not programming but coding.

Other terms used for minimization are extremization and hill-climbing. Since this can also be used to mean other things, the general conclusion is that in this field you can not tell a book from its title. While waiting for general agreement as to what the subject should be called, here it will be refered to as function minimization.

### 1.3 Definition of the problem

Given a function $F(\mathbf{x})$, the general problem is to find the value of the variable $\mathbf{x}$ for which the function $F(\mathbf{x})$ takes on its smallest value. (As pointed out above, this is entirely equivalent to finding the $\mathbf{x}$ for which the function $-F(\mathbf{x})$ takes on its largest
value, but for consistency only minimiziation will be considered here.) The rules of the game are the following:

1. The function $F(\mathbf{x})$ is assumed not to be known analytically, but is specified by giving its value at any point $\mathbf{x}$.
2. The allowed values of the variable $\mathbf{x}$ may be restricted to a certain range, in which case one speaks of constrained minimization. Here only unconstrained problems are refered to.
3. In some cases additional information about the function $F$ may be available, such as the numerical values of the derivative $\partial F / \partial \mathbf{x}$ at any point $\mathbf{x}$. Such knowledge cannot in general be assumed, but should be used when possible.
4. The function $F(\mathbf{x})$ is repeatedly evaluated at different points $\mathbf{x}$ until its minimum value is attained.

The method which finds the minimum (within a given tolerance) after the fewest function evaluations is the best. Occasionally other considerations may be important, such as the amount of storage required by the method or the amount of computation required to implement the method, but normally the dominating factor will be the time spent in evaluating the function.

### 1.4 Definition of a minimum

The theorems of elementary calculus say that the function $F(\mathbf{x})$ must take on its smallest value at a point where either:

1. all derivatives $\partial F / \partial \mathbf{x}=0$ (a stationary point), or
2. some derivative $\partial F / \partial \mathbf{x}$ does not exist (a cusp), or
3. the point $\mathbf{x}$ is on the boundary of the allowed region (an edge point).

Although it is sometimes found useful to consider points satisfying the above properties, this approach of considering essentially the analytical properties of the function is clearly not well adapted to the rules of the game as outlined above. Indeed, when one considers that there may be any number of stationary points, cusps, and edge points, all of which may be arbitrarily hard to find by simply sampling the function value, the whole problem begins to appear hopeless unless some simplifying assumptions are made.

The usual simplification consists in abandoning the attempt to find the global minimum and being satisfied with the local minimum. A local minimum may be defined as a point $\mathbf{x}_{0}$, where for all points $\mathbf{x}$ in some neighbourhood arround $\mathbf{x}_{0} F(\mathbf{x})>F\left(\mathbf{x}_{0}\right)$.

Now the situation looks much brighter since the very definition of a local minimum suggests a general strategy for finding one: one varies $\mathbf{x}$ by small steps in a direction which causes $F$ to decrease, and continue until $F$ increases in all allowed directions from some point $\mathbf{x}_{0}$. This does not yet tell one how to vary $\mathbf{x}$, but at least it suggests that a solution can be found.

Here only unconstrained local minimization will be considered, unless otherwise stated. The problem of global minimization will be treated in ??? section 6 .

### 1.5 The shape of the function - Taylor's series

With a view to making an intelligent minimizing method, it is of interest to consider what one might reasonably expect about the behaviour of $F$. If $F$ represents a physically meaningful function, one would certainly expect all the derivatives of $F$ to exist everywhere in the region of interest. Under these conditions one can write down the Taylor's series expansion for $F$ about some point $\mathbf{x}_{1}$ :

$$
F(\mathbf{x})=F\left(\mathbf{x}_{1}\right)+\mathbf{g}^{\mathrm{T}}\left(\mathbf{x}-\mathbf{x}_{1}\right)+\frac{1}{2}\left(\mathbf{x}-\mathbf{x}_{1}\right) \mathbf{V}\left(\mathbf{x}-\mathbf{x}_{1}\right)^{\mathrm{T}}+\ldots,
$$

where $\mathbf{g}$ is the gradient vector

$$
\mathbf{g}_{i}=\frac{\partial F}{\partial x i}
$$

and the matrix $\mathbf{V}$ is defined by

$$
\mathbf{V}_{i j}=\frac{\partial^{2} F}{\partial x_{i} \partial x_{j}}
$$

all evaluated at $\mathbf{x}_{1}$. Note the difference between $x_{i}$ (the $i^{\text {th }}$ variable of $\mathbf{x}$ ) and $\mathbf{x}_{i}$ (the position vector $\mathbf{x}$ at point $i$ ).

Although one does not know anything a priori about the convergence of this series, one does know that as the distance $\left(\mathbf{x}-\mathbf{x}_{1}\right)$ becomes smaller, the higher order terms become less important, so that one would expect that predicitons based on the loworder terms should not be very wrong, at least for small steps.

The first term of the above series is constant, so it will not tell much about where to look for a minimum. The second term is proportional to the gradient $\mathbf{g}$, telling in which direction the function is decreasing fastest, but since it is linear in $\mathbf{g}$, it does not predict a minimum and therefore does not say what stepsize to take. Moreover, when approaching the minimum $\mathbf{g} \rightarrow 0$ (if it exists) so one will have to go further and consider the next term. The third, or quadratic term describes a parabolic behaviour and is therefore the lowest term to predict a minimum. Unlike $\mathbf{g}$ one can expect $\mathbf{V}$ to be roughly constant over small regions, since it would be exactly constant if higher-order terms were zero.

One class of problems needs be mentioned in which the above analysis would not hold at all. This is the field known as linear programming, which limits itself to minimizing functions which are linear in the parameters, subject to constraints which are also linear. A linear function can not have a minimum in the sense described above (a stationary point) but must take on its minimum at a constraint boundary (edgs point). For such problems the description of the constraints therefore takes on greater importance than the analysis of the function itself, and will not be considered here.

### 1.6 Non-existence of optimum in general

Although we will study and comparing different minimization algorithms (recipes), the reader should be warned at the outset that in the strict sense of the rules of the game as stated in section ??? 1.3 above, it will not be possible to show any algorithm to be superior to any other for all functions. In principle at least, no matter how bad one algorithm is, or how good another, one can always find a function which will be minimized faster by the bad method than by the good one. One should keep such essentially theoretical considerations in mind, but should not be overly discouraged by them. In particular, certain objective criteria will merge for comparing methods even though the principal criterion - minimization speed - depends on the function. In the past there has been an overemphasis on such objective criteria in an attempt to find the ideal universal minimization algorithm. More recently, the tendency is to adapt the algorithm to the function, even to the point of introducing a superalgorithm which would choose a sub-algorithm appropriate to the function at hand. Such questions of global strategy will be considered later.

The reader should also be warned that in presenting particular algorithms details will be often omitted which are unimportant to an understanding of the algorithm although they may be crucial in actually making it work. The original references should therefore be consulted before programming such algorithms.

### 1.7 The role of the computer

While the subject here is essentially a mathematically one, it has been profoundly influenced by the existence of high-speed electronic computers that it would certainly be unfair not to mention them here. Indeed, real progress in the solving of largescale practical problems has come only since the 1960's, although much of the basic theory dates back to Newton's time or even earlier. This is, of course, because of the renewed interest in numerical minimization techniques for use on computers. As it is no longer even thinkable to use these techniques for hand calculations, it is best to place itself immediately in the computer context and to conceive of the function $F(\mathbf{x})$ rather as a function object which returns a value of $F$ for given input values of x .

One unpleasant consequence of the computer-oriented approach is that one will often have to worry about rounding-off errors in the function value due to the finite word length of digital computers. In addition there may be problems of overflow and underflow. In a real program for minimization or analysis general functions, all numerical operations must be protected against such numerical exceptions, and this typically represents more than half of the computer code, sometimes nearly all of it.

## 2 One-dimensional minimization

### 2.1 Usefulness in $n$-dimensional problems

We will first consider functions of just one variable, since some general problems can be seen more easily in this simplest case and also because some $n$-variable algorithms contain steps which require one-dimensional minimization. The one-variable problem is therefore both instructive and useful even though our prime consideration will be that of more complex problems.

### 2.2 Grid search

The most elementary search technique consists in choosing $k$ equally spaced points within the range of the parameter $x$, evaluating the function at each of the points, and retaining the lowest value found. If the spacing between points is $\Delta x$, one of the points is sure to be within $\Delta x / 2$ of the true minimum, although in principle it may not be the point corresponding to the lowest value. Still, if the function does not vary too wildly over the distances of the order of $\Delta x$, one generally assumes that this method gives the minimum within a range of about $\Delta x$.

Of course the grid search method has some difficulties. It is not directly applicable to the usual case where the range of $x$ is infinite. But in this case a simple remedy is to choose a reasonable range in the middle of the allowed range, and later to shift the sampling range if the minimum comes out at an end point.

The most serious objection to the grid method is its inefficiecy. Given the assumption that $F$ does not vary too much over a distance of $\Delta x$, many of the function evaluations are certainly unnecessary, namely those that are in regions where the function value is known to be large. In other words, the algorithm takes no account of what it has learned about the function. This inefficiency becomes more striking, in fact prohibitive, when extended to many variables.

On the other hand, this method has the prized virtues of extreme simplicity and absolute stability. It always converges within the desired tolerance in a known number of steps and is quite insensitive to the detailed behaviour of the function.
The efficiency of the grid method may be greatly improved by proceeding in several stages, using a smaller range and smaller step size in each succeeding stage. In this way each stage takes account of the least value found in the preceding stage, and the method can be said to converge in the usual sense of increasing accuracy due to decreasing step size. In the next section we consider optimum ways to arrange staging in order to obtain the fastest decrease in step size.


Fig. 1


Fig. 2


Fig. 3

### 2.3 Fibonacci and golden section searches

In order to optimize the grid search, we want to minimize the number of function evaluations per stage, compatible with maintaining a constant reduction of a factor $t$ in the step sizes at each stage. This will yield the fastest reduction in step size. One function evalution tells us nothing about the possible location of a minimum, but as long as we restrict ourselves to local minima in a given range of $x$, two points are sufficient as shown in fig. 1. If $F\left(x_{1}\right)<F\left(x_{2}\right)$, then there must be at least one local minimum somewhere in the range $0<x<x_{2}$. Now in this new range, we already have one point $\left(x_{1}\right)$, so that a further reduction in range is possible with only one new function evaluation, and the procedure can now be continued with only one new evaluation per stage. It remains to be shown that this can be continued indefinitely with a constant reduction in step size, and to calculate what that reduction will be. Clearly we would get the maximum reduction on the first step if $x_{1}$ and $x_{2}$ were very close together, but we must not forget that $x_{1}$ (or $x_{2}$ ) will then be used for the next stage and should therefore be close to the middle of this new interval as well. The situation is illustrated in the figs. 2 and 3 , where the distances indicated are imposed by the symmetry of the intervals and the condition that the reduction in range must be a factor of $t$ in each stage. The new range after evaluation of $F\left(x_{3}\right)$ will be $x_{3}<x<x_{2}$ and its length must be $t^{2}$.

This will be possible since there is a real root to the equation:

$$
\begin{aligned}
t^{2} & =1-t \\
t & =\frac{\sqrt{5}-1}{2} \approx 0.616
\end{aligned}
$$

Since this ratio $t$ is known as the golden section, the minimization technique is called a golden section search. If the number of stages to be taken is known in advance, it is possible to improve very slightly on this technique by using a Fibonacci search, as described for example in Kowalik and Osborne [1]. Although Fibonacci can be shown to be optimal (in a sense described below), the slight improvement is probably not worth the added complication. The golden section search is optimal among algorithms where the stopping point is not decided in advance.

The above techniques are optimal only in the minimax sense, that is they minimize the maximum number of function evaluations necessary to obtain a given accuracy. It might be called the pessimist's optimality, since in game theory it is the best strategy against an intelligent opponent who is trying to make you lose. It should therefore be effective in minimizing pathological functions, but in more normal cases we should expect other methods to be better. Such methods are described in the following sections.

### 2.4 Quadratic interpolation and extrapolation

A more optimistic approach consists in studying the expected behaviour of the function and then hoping that the deviations of the real function from this behaviour are not too great. From the Taylor's series analysis of Section 1.5, it would be reasonable to proceed by assuming that the function is nearly quadratic.

Since a parabola is determined by three points, this method requires the function to have been evaluated for three different values $x_{1}, x_{2}$ and $x_{3}$. It then predicts the minimum to be at the minimum of the parabola passing through these points. If the three function values are $F_{1}, F_{2}$, and $F_{3}$, the predicted minimum is at $x_{4}$ given by

$$
x_{4}=\frac{\frac{\left(x_{2}+x_{3}\right) F_{1}}{\left(x_{1}-x_{2}\right)\left(x_{1}-x_{3}\right)}+\frac{\left(x_{1}+x_{3}\right) F_{2}}{\left(x_{2}-x_{1}\right)\left(x_{2}-x_{3}\right)}+\frac{\left(x_{1}+x_{2}\right) F_{3}}{\left(x_{3}-x_{1}\right)\left(x_{3}-x_{2}\right)}}{2\left[\frac{F_{1}}{\left(x_{1}+x_{2}\right)\left(x_{1}-x_{3}\right)}+\frac{F_{2}}{\left(x_{2}+x_{1}\right)\left(x_{2}-x_{3}\right)}+\frac{F_{3}}{\left(x_{3}+x_{1}\right)\left(x_{3}-x_{2}\right)}\right]}
$$

Considerable simplification results when the three points are equally spaced, a distance $d$ apart, in which case:

$$
x_{4}=x_{2}+\frac{d}{2} \frac{\left(F_{1}-F_{2}\right)}{\left(F_{1}+F_{3}-2 F_{2}\right)}
$$

The function is then evaluated at $x_{4}$, this point replaces one of the first three, and a new point is predicted, again by quadratic interpolation using the new set of three
points. The method terminates when the predicted function value at some new point agrees with the actual value within a specified tolerance.

This algorithm usually performs quite well when applied to easy (nearly quadratic) functions, but suffers from a number of instabilities which can be quite serious, as follows:
i) At any step the three points may determine a parabola with a maximum rather than a minimum, in which case the method diverges.
ii) If the three points lie nearly in a straight line, the algorithm takes an enormous step which may cause numerical difficulties as well as diverging.
iii) After each step there is a choice of which two of the three previous points to retain for the next step. It is usually more convenient and logical to retain the most recent points, but this may also lead to instabilities by throwing away the best points.
iv) Even without any of the above difficulties, the method may oscillate about the minimum instead of converging toward it.

All the problems can be fixed by including checks and safeguards in the algorithm, but the remedies always involve abandoning, at least temporarily, the quadratic interpolation step. The best remedy is probably to reserve the method for wellbehaved functions and to abandon it entirely as soon as trouble arises. It is most often used as the last step in algorithms which depend principally on other methods, since physical functions are usually quite parabolic in the immediate vicinity of the minimum.

When derivatives of the function are available, variations of quadratic interpolation are possible, using instead of three points to determine the parabola, either two function values and one first derivative, or the function value and the first two derivatives at one point. These variations tend to be even more unstable than the basic method, since they use information from fewer points.

### 2.5 The success-failure method

A good compromise between the stability of the grid search and the rapid convergence of quadratic interpolation is found with the success-failure technique of Rosenbrock [2]. A start point $x_{0}$ and initial step size $d$ are required, and the function is evaluated at $x_{0}$ and $x_{0}+d$. The first step is termed a success if $F\left(x_{0}+d\right)<F\left(x_{0}\right)$, otherwise it is a failure. If it is a failure, $d$ is replaced by $-\beta d$, where $\beta$ is a contraction factor less than one, and the test is repeated. If it is a success, $x_{0}$ is replaced by $x_{0}+d, d$ is replaced by $\alpha d$, where $\alpha$ is an expansion factor greater than one, and the test is repeated. The process continues in this way until the function values change by less than a specified amount, The numerical values usually used for the expansion and contraction parameters are $\alpha \approx 3.0$ and $\beta \approx 0.4$.

An interesting feature of this method is that a local minimum is always bracketed
whenever a success is followed by a failure. When this happens, the middle one of the last three points is always lower than the outer two, so that one is in a favourable position for trying a quadratic interpolation step. The success-failure method, with one quadratic interpolation step each time a success is followed by a failure, is probably the most effective one-dimensional technique for use on general functions although in special cases other methods may be superior.


Fig. 4


Fig. 5

## 3 Stepping methods in many variables

### 3.1 Grid search and random searches

An excellent illustration of the enormous increase in complexity in going to spaces of high dimensionality is afforded by the grid search technique in many variables. In order to localize a minimum to $1 \%$ of the range of one variable by this technique requires 100 function evaluations; in ten variables the number of points required is $10^{20}$. Clearly we can forget about this method when more than one or two parameters are involved.

In fact it is a general rule in function minimization, as in function integration, that one should not expect good one-dimensional techniques to be good when extended to higher dimensionality. Experience with integration suggests that a Monte Carlo search is more efficient than a grid search in many dimensions. The Monte Carlo technique consists in choosing points randomly according to some distribution (usually uniform or normal).

But even when these methods are refined by using variable search ranges, they prove far too slow for general use and we must turn to more efficient techniques.

### 3.2 Single-parameter variation

Since the condition for a minimum which is a stationary point in $n$ variables $x_{i}$ is the vanishing of all $n$ first derivatives $\partial F / \partial x_{i}$, it is natural to try to make each derivative vanish separately, one after the other. This is the old method of single parameter variation, where one seeks a minimum with respect to one variable at a time using one of the techniques described earlier. Of course when you have finished minimizing with respect to $x_{2}$ you may no longer be at a minimum with respect to $x_{1}$, so you generally have to start all over again, but the process usually does converge, as illustrated for two variables in fig. 4. Here the curves represent contours of equal function value, and the straight lines show the steps taken
in minimizing $F$ with respect to $x_{1}$, then $x_{2}$, then $x_{1}$, etc. In this case the method converges nicely after only four single-parameter minimizations.

Consider now the function represented by the contours shown in fig. 5. Here the method proceeds much more slowly because of the narrow valley. It still converges, but as the valley becomes narrower, the convergence becomes arbitrarily slow.

Such behaviour in many dimensions causes this method to be generally considered as unacceptably slow.

Two of the more successful improvements aimed at avoiding such behaviour are due to Hooke and Jeeves [3] and Rosenbrock [2]. We discuss the latter below.

### 3.3 The Rosenbrock's method

Rosenbrock's algorithm [2] starts by performing single-parameter minimizations as above. Then when one full cycle of all parameters has been completed, a new set of orthogonal axes is defined with one axis taken as the vector from the start point to end point of the cycle. This vector points in the direction of previous over-all improvement and is expected to be a good direction for future improvement. In the case of the narrow valley seen above, it should point more or less along the valley and avoid the zig-zag behaviour. The next cycle of single-variable minimizations is performed using multiples of the newly defined axes as variables.

The Rosenbrock method generally performs well, being quite stable and capable of following narrow valleys, but as the number of variables increases, the efficiency drops, probably because the new axis defined by past improvement is the only 'intelligent direction' used in the next cycle. All the other minimization directions are simply chosen orthogonal to the first one. Also, its terminal convergence is slow compared with the more 'quadratic' methods described in Section 4.

Another technique, that of Davies, Swann, and Campey [4] (unpublished, see Ref. 4) is similar to Rosenbrock's and will not be described here.

### 3.4 The simplex method

One of the most successful stepping methods in many variables is that of Nelder and Mead [5], based on the simplex. A simplex is an $n$-dimensional figure specified by giving its $n+1$ vertices. It is a triangle in two dimensions, a tetrahedron in three, etc. The algorithm takes the name simplex because at each step the information it carries about the function consists of its values at $n+1$ points. One can easily visualize how the method works by considering the two-dimensional case as in fig. 6 . The three starting simplex points are somehow chosen (perhaps randomly) and the function is evaluated at each point. Let the point $P_{H}$ be that at which the function value is highest (worst) and $P_{L}$ that at which it is lowest. Let $\bar{P}$ be the centre-of-mass


Fig. 6
of all points in the simplex except $P_{H}$; that is:

$$
\bar{P}=\frac{1}{n}\left\{\sum_{i=1}^{n+1} P_{i}-P_{H}\right\}
$$

From the original simplex, a new simplex is formed by replacing $P_{H}$ by a better point if possible. The first attempt to find a better point is made by reflecting $P_{H}$ with respect to $\bar{P}$, producing $P^{*}=\bar{P}+\left(\bar{P}-P_{H}\right)$. If $F\left(P^{*}\right)<F\left(P_{L}\right)$, a new point is tried at $P^{* *}=\bar{P}+2\left(\bar{P}-P_{H}\right)$. If $F\left(P^{*}\right)>F\left(P_{H}\right)$, a new point is tried at $P^{* *}=\bar{P}-1 / 2\left(\bar{P}-P_{H}\right)$. The best of the new points then replaces $P_{H}$ in the simplex for the next step, unless none of them is better than $P_{H}$. In the latter case, a whole new simplex is formed around $P_{L}$, with dimensions reduced by a factor of 0.5 .

Variations on the method are possible by using different contraction or expansion factors when searching along the line from $P_{H}$ through $\bar{P}$ (dotted in diagram). Another interesting possibility is to attempt a quadratic interpolation step along the dotted line whenever three points have been determined $\left(P_{H}, P^{*}, P^{* *}\right)$. However, one must be careful not to accept a point too close to $\bar{P}$, for then the simplex collapses into a line (or in general a hyperplane of $n-1$ dimensions) from which it can never recover.

The simplex algorithm, being designed always to take as big steps as possible, is rather insensitive to shallow local minima or fine structure in the function caused by rounding errors, statistical errors (Monte Carlo output), etc. Another of its virtues is that of requiring few function evaluations, usually one or two per iteration. In addition, each search is in an 'intelligent' direction, pointing from the highest value to the average of the lowest values. Compare this with Rosenbrock's method, where really only the principal axis is an 'intelligent' direction, and all other searches are for exploring along orthogonal axes to determine a new principal axis.

A convenient convergence criterion for the simplex method is based on the difference $F\left(P_{H}\right)-F\left(P_{L}\right)$. The iterations are stopped when this difference is less than a preset value. As a final step, the function is evaluated at $\bar{P}$, which is often slightly better than $F\left(P_{L}\right)$.

In view of the danger mentioned above - of the simplex collapsing into a hyperplane of dimension $n-1$-it has been suggested to use $n+2$ or more points rather than $n+1$ at each step. I have tested this idea, which is equivalent to introducing a dummy parameter of which the function is independent, and have always found the efficiency of the algorithm to decrease under these conditions.

### 3.5 The conjugate directions method

This method does not require information about the derivatives of the function, but the exploration requires the material developed in Chapter 4, so it is discussed in Section 4.5.

## 4 Gradient methods

### 4.1 Calculating derivatives

I will call a gradient method any technique which uses information from a very small range of the variables (i.e. essentially derivatives) to predict good trial points relatively far away. This does not necessarily mean that they follow the gradient, but only that the gradient, and perhaps higher derivatives, are used or estimated.

It is of course possible in most cases to calculate analytically the numerical values of the derivatives of a function, just as it is possible to calculate the value of the function itseif. However, it is often inconvenient and dangerous if the algebra is complicated, so that very often we are faced with minimizing a function for which no derivatives are provided. Since the most powerful algorithms discussed below require derivatives, a general minimization program must be able to estimate the derivatives of the function by finite differences.

A first derivative may be estimated from

$$
\left.\frac{\partial F}{\partial x}\right|_{x_{0}} \approx \frac{F\left(x_{0}+d\right)-F\left(x_{0}\right)}{d}
$$

where $d$ is a 'small' displacement. The error will be, to lowest order in the Taylor's expansion,

$$
\left.\delta \approx \frac{d}{2} \cdot \frac{\partial^{2} F}{\partial x^{2}}\right|_{x_{0}}
$$

It is therefore advantageous to make $d$ as small as possible, but still large enough so that the rounding error in the computation of $F$ does not become larger than the error introduced by $\delta$. Since the second derivatives may not be known, it may not be possible to find an optimum step-size $d$, so we may just have to close our eyes and guess.

A much safer method would be to use points chosen symmetrically on either side of $x_{0}$ giving

$$
\left.\frac{\partial F}{\partial x}\right|_{x_{0}} \approx \frac{F\left(x_{0}+d\right)-F\left(x_{0}-d\right)}{2 d}
$$

for in this case the error $\delta$ vanishes to second order and the lowest order term is proportional to the third derivative. A disadvantage of this method is that it requires $2 n$ function calls to estimate the $n$ first derivatives, whereas the asymmetric steps require only $n+1$ [or only $n$ if $F\left(x_{0}\right)$ has to be evaluated anyway]. An advantage of the symmetric steps method, however, is that it gives the second derivatives as a by-product [assuming $F\left(x_{0}\right)$ known]:

$$
\frac{\partial^{2} F}{\partial x^{2}} \approx \frac{F\left(x_{0}-d\right)+F\left(x_{0}+d\right)-2 F\left(x_{0}\right)}{d^{2}}
$$



Fig. 7


Fig. 8
and from the relationship for the error $\delta$ in the asymmetric method, a conservative upper limit of the uncertainty in the first derivative results assuming at least that the symmetric formula gives a smaller error than the asymmetric one. A complete treatment of step sizes is beyond the scope of these lectures but can be found in a paper by Stewart [6].
The numerical evaluation of second derivatives is facilitated by the fact that they should be approximately constant over small regions, so that symmetrical steps are usually not necessary. Unfortunately, however, there are a lot of second derivatives to evaluate; since they form a symmetric $n \times n$ matrix, there are $n(n+1) / 2$ independent components, requiring at least $n(n-1) / 2$ points in addition to those required for the symmetric derivatives. For two parameters, a minimum point pattern is shown fig. 7. The odd point (for the mixed second derivative) could have been chosen in any corner. The two-dimensional diagram is somewhat misleading since for large $n$, the number of 'odd points' is $n$ times larger than the number of 'symmetric' points.

### 4.2 Steepest descent

As soon as the function's first derivatives are known, it is natural to follow the direction of the negative gradient vector in seeking a minimum, since this is the direction in which the function is decreasing the fastest. Such a technique was used by Cauchy more than a century ago, and is the basis of what is now known as the method of steepest descent.

This method consists of a series of one dimensional minimizations, each one along the direction of local steepest descent (gradient) at the point where each search begins. Of course the direction of the gradient is not constant along a line even for a general quadratic function, so we expect many iterations to be necessary, but the method can be shown to converge for a quadratic function. Let us follow its progress for a typical function whose contours are shown in fig. 8. We immediately see an unfortunate property of the successive search directions: if each linear minimization is exact,


Fig. 9
successive searches must be in orthogonal directions. In two dimensions, this yields steps which look just like the single parameter variation method (fig. 5) with the axes rotated to line up with the gradient at the start point. In many dimensions the situation is not quite so bad, but successive directions are still orthogonal and the algorithm cannot be considered acceptable. It is in fact easy to draw contours for a reasonably well-behaved hypothetical function (fig. 9) where the direction to the minimum is just perpendicular to the gradient.

### 4.3 Newton's method

It is clear that since a general quadratic function is determined by specifying its value, first derivatives, and second derivatives at a point, it can be minimized in one step if and only if all this information (or its equivalent) is taken into account. Let us write a quadratic function as

$$
F(\underline{x})=F\left(\underline{x}_{0}\right)+\underline{g}^{T}\left(\underline{x}-\underline{x}_{0}\right)+\frac{1}{2}\left(\underline{x}-\underline{x}_{0}\right)^{T} \mathbf{G}\left(\underline{x}-\underline{x}_{0}\right),
$$

where the gradient $\underline{g}$ is evaluated at $\underline{x}_{0}$ and the second derivative matirx

## G

is a constant. Then the minimum is given directly by

$$
\underline{x}_{m}=\underline{x}_{0}-\mathbf{G}^{-1} \underline{g}=\underline{x}_{0}-\mathbf{V} \underline{g},
$$

where the inverse of the second derivative matrix is the covariance matrix $\mathbf{V}$.
This is then the many-dimensional equivalent of quadratic interpolation discussed earlier, and it is subject to the same sort of difficulties when applied as an iterative
technique to general non-quadratic functions. But let us first point out its good features:
i) the step size is no longer arbitrary, but is prescribed precisely by the method;
ii) the step directions are no longer necessarily along the gradient vector but take account of parameter correlations (narrow valleys or ridges) through the mixed second derivative terms.

In practice, however, the method is unstable, essentially for the reasons given in Section 2.4. In particular, it diverges whenever the matrix $\mathbf{G}$ (or $\mathbf{V}$ ) is not positivedefinite (see next section). In its unmodified form the method is used only when the minimum is known to be very close or when the function is known to be positive quadratic (for linear least squares). However, it is clearly a powerful technique and is worth studying in some detail since all the most successful algorithms are based on Newton-like steps, as discussed below.

### 4.4 Positive-definite quadratic forms

We pause here briefly to consider the properties of quadratic forms useful for understanding the more powerful gradient methods. In one dimension the description is simple; a general quadratic form can be written

$$
F(x)=a+g x+\frac{1}{2} G x^{2}
$$

where $g=\partial F / \partial x$ at $x=0$, and $G=\partial^{2} F / \partial x^{2}$ also at $x=0$. This function has a minimum if and only if $G \geq 0$. If $\mathbf{G}=0$, the minimum is at infinity, The minimum (if it exists) is at $x=-g / G$. When using a quadratic approximation to minimize a general non-linear function, it makes sense to take a step to $x=-g / G$ only if $G>0$ since otherwise we step to a predicted maximum or to infinity. A possible remedy if $G<0$ is to take a step $x=-g$; that is, to set $\mathbf{G}$ arbitrarily equal to unity so that the step will at least be in the right direction although it will now have arbitrary length. Consideration of fig. 10 shows that this is the only thing we can do unless more information is available, since the quadratic part of the function is not convex or positive-definite at the point $x_{0}$.

These arguments may now be extended to many dimensions where $g$ becomes the gradient vector $g$, and $\mathbf{G}$ becomes the second derivative matrix $\mathbf{G}$. Then the Newton step to $\underline{x}=-\overline{\mathbf{G}}^{-1} \underline{g}$ makes sense only if $\mathbf{G}$ (hence $\mathbf{G}^{-1}$ ) is a positive-definite matrix, since only then does the quadratic form

$$
F(\underline{x})=a+\underline{g}^{T} \cdot \underline{x}+\frac{1}{2} \underline{x}^{T} \mathbf{V} \underline{x}
$$

have a minimum. If $\mathbf{G}$ is singular, the predicted minimum (or maximum) is not unique.


Fig. 10


Fig. 11

Unfortunately there is no simple way of telling, in general, if a matrix is positivedefinite by inspecting individual components, but we can at least state some of the many useful properties of such matrices. Two necessary (but not sufficient) conditions for a (square, symmetric) matrix to be positive-definite are:
i) the diagonal elements must be positive (this is in fact sufficient for a $1 \times 1$ matrix);
ii) the off-diagonal elements must obey $G_{i j}^{2}<G_{i i} G j j$.
[Properties (i) and (ii) together are sufficient for a $2 \times 2$ matrix.] While the above conditions are easy to check, they are not in general sufficient. Some necessary and sufficient conditions are the following:
iii) All the eigenvalues of the matrix are positive. This is generally a rather difficult calculation and is usually approximate.
iv) The determinants of all the upper left square submatrices (formed as indicated in the diagram in fig. 11) are positive. This is probably the easiest method.
v) The scalar $\underline{e}^{T} \mathbf{G} \underline{e}$ is positive for all vectors $\underline{e}$ This is usually taken as the definition of a positive-definite matrix, and explains why a positive-definite matrix yields a quadratic form with a minimum: the function increases in all directions from $\underline{e}=0$.
vi) The inverse $\mathbf{G}^{-1}=\mathbf{V}$ is positive-definite.

Now suppose that $\mathbf{G}^{-1}$ is calculated for a Newton step and turns out to be non-positive-definite. In analogy to the one dimensional case we would simply take $\mathbf{G}=\mathbf{I}$, the unit matrix, and the Newton step would become a steepest-descent step of arbitrary length, which is probably not so bad an idea and is in fact often done. But we can do better by trying to make a positive-definite matrix which is as 'close' as possible to the unacceptable $\mathbf{G}$. This is done as follows: The matrix $(\mathbf{G}+\lambda \mathbf{I})^{-1}$ is used instead of $\mathbf{G}^{-1}$, where $\lambda$ is greater than the largest negative eigenvalue of $\mathbf{G}$. This requires a fair amount of calculation and so is not very convenient, but it is quite appealing since it amounts to taking a step which is intermediate between a Newton step and a steepest-descent step (for large values of $\lambda$ the step becomes short and in the direction of the gradient).

If we are willing to calculate eigenvectors as well as eigenvalues, the non-positivedefiniteness can be turned into an advantage, since the eigenvector corresponding to a negative eigenvalue indicates a direction (or directions) in which the negative first derivative is increasing in magnitude rather than decreasing. This suggests an especially fruitful direction for a single-parameter-variation step which should not only lead to a good decrease of the function value but should also lead more quickly to a region of positive-definiteness.

Minimization methods based on variations of Newton's method as suggested by the above considerations are usually called quasi-Newton methods. Many such algorithms have been published and some are quite successful, but the field is still open for new ideas.

The principal drawback of such techniques is the repeated evaluation and inversion of the second-derivative matrix. The calculation of the second derivatives usually requires a rather long time, proportional to $n^{2}$, and the matrix inversion, although usually faster, increases with $n$ like $n^{3}$.

One of the most interesting results concerning quadratic forms is the basis of a collection of related techniques described in the next sections, which do not require explicit repeated evaluations of G.

### 4.5 Conjugate directions

The vectors $\underline{d}_{i}$ and $\underline{d}_{j}$ are said to be conjugate with respect to a positive-definite symmetric matrix $\mathbf{A}$ if

$$
\underline{d}_{i}^{T} \mathbf{A} \underline{d}_{j}=0 \quad \text { for } \quad i \neq j
$$

If $\mathbf{A}$ is the unit matrix $\mathbf{I}$, the conjugate vectors $\mathbf{d}$ would be orthogonal, so conjugacy can be thought of as a generalization of orthogonality. A set of $n$ conjugate vectors span an $n$-dimensional space, and any point in the space can therefore be expressed as a linear combination of $n$ conjugate vectors.

Although the matrix A does not uniquely define a set of conjugate vectors, such a set can always be constructed by a procedure similar to the Gram-Schmidt orthogonalization method. Let us start for example with an arbitrary vector $\underline{d}_{1}$. Then the vector

$$
\underline{d}_{2}=\mathbf{A} \underline{d}_{1}-\frac{\underline{d}_{1}^{T} \mathbf{A} \mathbf{A} \underline{d}_{1}}{\underline{d}_{1}^{T} \mathbf{A} \underline{d}_{1}} \underline{d}_{1}
$$

can be seen to be conjugate to $\underline{d}_{1}$ since the product $\underline{d}_{1}^{T} \mathbf{A} \underline{d}_{2}$ vanishes identically. The process can then be continued in the same way to construct a $\underline{d}_{3}$ which will be conjugate to both $\underline{d}_{1}$ and $\underline{d}_{2}$, and so forth up to $\underline{d}_{n}$.

Such vectors become interesting for minimization problems when they are conjugate with respect to the hessian (second derivative) matrix G. In this case a theorem of Fletcher and Reeves [7] states that a sequence of linear minimizations in each of the $n$ conjugate directions will minimize a general quadratic function of $n$ variables. That this is true can be seen quite easily as follows. Let the quadratic function be

$$
F(\underline{x})=F(\underline{0})+\underline{g}^{T} \underline{x}+\frac{1}{2} \underline{x}^{T} \mathbf{G} \underline{x}
$$

and the $n$ directions $d_{i}$ be conjugate with respect to $\mathbf{G}$ :

$$
\underline{d}_{i}^{T} \mathbf{G} \underline{d}_{j}=0, \quad i \neq j .
$$

Then the vectors $\underline{x}$ and $\underline{g}$ can be expressed as linear combinations

$$
\begin{aligned}
& \underline{x}=\sum_{i} y_{i} \underline{d}_{i} \\
& \underline{g}=\sum_{i} c_{i} \underline{d}_{i}
\end{aligned}
$$

so that the general quadratic becomes

$$
F(\underline{x})=F(\underline{0})+\left(\sum_{i} c_{i} \underline{d}_{i}^{T}\right)\left(\sum_{j} y_{j} \underline{d}_{j}\right)+\frac{1}{2}\left(\sum_{i} y_{i} \underline{d}_{i}^{T}\right) \mathbf{V}\left(\sum_{j} y_{j} \underline{d}_{j}\right) .
$$

Now if the last term above is regrouped as a double sum, the terms with $i \neq j$ drop out because of the conjugacy condition, so that the whole expression can be simplified as

$$
\begin{aligned}
F(\underline{x}) & =F(\underline{0})+\sum_{i} \sum_{j} c_{i} \underline{d}_{i}^{T} \underline{d}_{j} y_{j}+\frac{1}{2} \sum_{j} y_{j}^{2} \underline{d}_{j}^{T} \mathbf{G} \underline{d}_{j} \\
& =F(\underline{0})+\sum_{j}\left(b_{j} y_{j}+b_{j}^{\prime} y_{j}^{2}\right)
\end{aligned}
$$

where

$$
b_{j}=\sum_{i} c_{i} \underline{d}_{i}^{T} \underline{d}_{j}
$$

and

$$
b_{j}^{\prime}=\underline{d}_{j}^{T} \mathbf{G} \underline{d}_{j}
$$

are constants. By expressing the quadratic in terms of $y$ instead of $x$ we have separated it into a sum of independent one-parameter quadratic functions. A minimization with respect to $y_{i}$ (a linear minimization along the direction $\underline{d}_{i}$ ) will therefore be independent of the minimizations along the other conjugate directions, which demonstrates the validity of the theorem.

The above theorem tells us what is 'wrong' with the single-parameter-variation method: we should be using conjugate directions rather than simply orthogonal


Fig. 12
axes. However, since the construction of conjugate vectors seems to require knowledge of the hessian $\mathbf{G}$, this does not yet help very much in practice, for if we knew $\mathbf{G}$ (and $\underline{g}$ ) we could minimize a quadratic immediately by means of Newton's method, and would not need to use $n$ linear minimizations.

The usefulness of conjugate directions comes from the fact that there are ways of determining such directions implicitly, without first evaluating the entire hessian matrix $\mathbf{G}$. Of course, by the time all $n$ conjugate directions are determined, by whatever method, information equivalent to the matrix $\mathbf{G}$ must have been determined. However, by that time considerable minimization may already have been performed, as in the method implied by the following theorem.

If $\underline{x}_{0}$ and $\underline{x}_{1}$ are minimum points in two parallel subspaces, then the direction $\underline{x}_{1}-\underline{x}_{0}$ is conjugate to any vector which lies in either subspace. This can easily be seen in two dimensions as illustrated in fig. 12. Since $\underline{x}_{0}$ is a minimum along the direction $\underline{d}_{1}$ the gradient of $F$ at $\underline{x}_{0}$ must be orthogonal to $\underline{d}_{1}$ :

$$
\underline{d}_{1}^{T}\left(\underline{g}+\mathbf{G} \underline{x}_{0}\right)=0
$$

where $\underline{g}$ is the gradient at $\underline{x}=\underline{0}$. Similarly at $\underline{x}_{1}$ :

$$
\underline{d}_{1}^{T}\left(\underline{g}+\mathbf{G} \underline{x}_{1}\right)=0 .
$$

Subtracting the above equations, the first terms drop out and we have:

$$
\underline{d}_{1}^{T} \mathbf{G}\left(\underline{x}_{1}-\underline{x}_{0}\right)=0
$$

showing that $\left(\underline{x}_{1}-\underline{x}_{0}\right)$ is conjugate to $\underline{d}_{1}$.
Unfortunately, extending this algorithm to three dimensions requires three additional minimizations in order that the third direction be conjugate to both of the first two, so that convergence for a general quadratic in $n$ variables is obtained only after $n$
iterations involving in all $n(n+1) / 2$ linear minimizations. Since this is just the number of independent elements in the second derivative matrix, we would be better off for quadratic functions to calculate this matrix directly and avoid the linear searches. On the other hand, for non-quadratic functions the conjugate directions method should be much more stable since it proceeds by a series of linear searches in independent directions and still guarantees convergence in a finite number of steps once a quadratic region is entered. In addition, this method has the advantage of requiring neither first nor second derivatives of the function. (Strictly speaking, then, it should have been discussed in Section 3 rather than in this section.)

A disadvantage of the algorithm described above is that for each iteration, $n$ minimizations are performed in direction $\underline{d}_{1}$, whilst only one is performed in direction $\underline{d}_{n}$. This undesirable asymmetry is largely avoided in a variation due to Powell [8].

### 4.6 Conjugate gradients

When the first derivatives of the function are calculated, a somewhat more elegant method can be used, known as the method of conjugate gradients [7]. Suppose that the function and its gradient are evaluated at two points $\underline{x}_{0}$ and $\underline{x}_{1}$, giving differences:

$$
\begin{aligned}
& \underline{\Delta x}=\underline{x}_{1}-\underline{x}_{0} \\
& \underline{\Delta g}=\underline{g}_{1}-\underline{g}_{0} .
\end{aligned}
$$

Then if the function were quadratic with hessian $\mathbf{V}$ we would have

$$
\underline{\Delta g} \underline{g}=\mathbf{G} \underline{\Delta x} .
$$

Any vector $\underline{d}_{1}$ orthogonal to $\underline{\Delta g}$ would then be conjugate to $\underline{\Delta x}$ :

$$
\underline{d}_{1}^{T} \underline{\Delta} \underline{g}=\underline{d}_{1}^{T} \mathbf{G} \underline{\Delta x}=0
$$

which immediately suggests a method for obtaining conjugate directions without knowing $\mathbf{G}$, based on the change in gradient along a previous direction.

In the method of conjugate gradients, successive one-dimensional minimizations are performed along conjugate directions with each direction being used only once per iteration. The first direction is taken as $\underline{d}_{0}=-\underline{g}_{0}$, the steepest descent vector at $\underline{x}_{0}$. Let the minimum along this direction be at $\underline{x}_{1}$ where the gradient is $\underline{g}_{1}$. Then the next search direction $\underline{d}_{1}$, which we want to be conjugate to $\underline{d}_{0}$ must be a linear combination of the only vectors we have at hand, namely:

$$
\underline{d}_{1}=-\underline{g}_{1}+b \underline{d}_{0} .
$$

The conjugacy condition is

$$
\underline{d}_{1}^{T} \mathbf{G} \underline{d}_{0}=\underline{d}_{1}^{T} \mathbf{G}\left(\underline{x}_{1}-\underline{x}_{0}\right)=0
$$

or

$$
\left(-\underline{g}_{1}^{T}+b \underline{d}_{0}^{T}\right) \mathbf{G} \underline{d}_{0}=\left(-\underline{g}_{1}^{T}-b \underline{g}_{0}^{T}\right)\left(\underline{g}_{1}-\underline{g}_{0}\right)=0 .
$$

Since $\underline{x}_{1}$ is a minimum along direction $\underline{d}_{0}=-\underline{g}_{0}$, the direction $\underline{g}_{0}$ is orthogonal to the gradient at $\underline{x}_{1}$, so that $\underline{g}_{1}^{T} \underline{g}_{0}=0$. We are then left with

$$
b=\frac{\underline{g}_{1}^{T} \underline{g}_{1}}{\underline{g}_{0}^{T} \underline{g}_{0}}
$$

so that the new conjugate direction is

$$
\underline{d}_{1}=-\underline{g}_{1}+\left(\frac{\underline{g}_{1}^{T} \underline{g}_{1}}{\underline{g}_{0}^{T} \underline{g}_{0}}\right) \underline{d}_{0} .
$$

This process can be continued to generate $n$ directions, each one conjugate to all the others. It turns out that the same simple formula holds for all the successive conjugate directions

$$
\underline{d}_{i+1}=-\underline{g}_{i+1}+\left(\frac{\underline{g}_{i+1}^{T} \underline{g}_{i+1}}{\underline{g}_{i}^{T} \underline{g}_{i}}\right) \underline{d}_{i} .
$$

### 4.7 Variable metric methods (VMM)

In analogy with the methods of differential geometry and general relativity, it is convenient to consider the properties of the function $F(\mathbf{x})$ as being in fact properties of the space of the variables $\mathbf{x}$. This was already used rudimentarily when generalizing from the usual orthogonal coordinate axes to a system defined by axes pointing in conjugate directions. One now wishes to go further and be able to express the properties of the function $F$ geometrically as the properties of the non-Euclidean space of its variables $\mathbf{x}$.

The fundamental invariant in a non-Euclidean space is the squared distance element

$$
\Delta s^{2}=\Delta \mathbf{x}^{\mathrm{T}} \mathbf{A} \Delta \mathbf{x}
$$

where $\Delta \mathbf{x}$ is a differential coordinate displacement and $\mathbf{A}$ is a covariant metric tensor which determines all the properties of the space under consideration. When $\mathbf{A}$ is just the unit matrix $\mathbf{I}$, the above formula for $\Delta s^{2}$ just expresses the Pythagorean theorem for an $n$-dimensional Euclidean space. When off-diagonal elements of $\mathbf{A}$ are nonzero and when the elements are allowed to vary as functions of $\mathbf{x}$, a generalized non-Euclidean space is generated.

It is easily verified that the second derivative (Hessian) matrix $\mathbf{G}$ behaves under coordinate transformations like a covariant tensor and will be identified with the metric tensor of our space. The inverse $\mathbf{V}=\mathbf{G}^{-1}$ is a contravariant tensor and becomes the contravariant metric tensor. (For a discussion of covariant and contravariant tensors,
see for example chapter 10 of [4].) This immediately enables to construct two scalar (invariant under coordinate transformations) quantities:

$$
\begin{equation*}
\Delta s^{2}=\Delta \mathbf{x}^{\mathrm{T}} \mathbf{G} \Delta \mathbf{x} \tag{4.1}
\end{equation*}
$$

is the square of the generalized distance between the point $\mathbf{x}$ and the point $\mathbf{x}+\Delta \mathbf{x}$. When $F$ is a $\chi^{2}$ function which is minimized to determine some best parameters $\mathbf{x}$, the physical meaning of the generalized distance $\Delta s$ is just the number of "standard deviations" $\mathbf{x}+\Delta \mathbf{x}$ is away from $\mathbf{x}$. That is, the use of the metric tensor $\mathbf{V}$ enables one to scale the distance $\Delta \mathrm{x}$ so that it comes out as a physically (or statistically) meaningful invariant quantity instead of being expressed in arbitrary units (or a mixture of arbitrary units!).

And

$$
\begin{equation*}
\rho=\mathbf{g}^{\mathrm{T}} \mathbf{V} \mathbf{g} \tag{4.2}
\end{equation*}
$$

is twice the difference between the function value at the point where $\mathbf{V}$ and the gradient $\mathbf{g}$ are calculated and the minimum of a quadratic form with hessian matrix $\mathbf{G}=\mathbf{V}^{-1}$. That is, $\rho / 2$ is the expected (vertical) distance to the minimum if the function $F$ were quadratic. This provides an important scale-free convergence criterion for any method which provides approximations to $\mathbf{V}$ and $\mathbf{g}$.

When the function $F$ is quadratic, $\mathbf{G}$ is constant everywhere and, in the sense outlined above, this is equivalent to working in a space with constant metric. For real non-linear functions higher-order terms are expected to be small but not negligible, so that one can think of working in a space with a slowly-varying metric tensor. Minimization methods based on this approach are known as variable metric methods. They differ from the basic Newton-Raphson method in the sense that the matrix G is not completely re-evaluated at each iteration, but is assumed to be well approximated by taking the $\mathbf{G}$ of the previous iteration and apply a correction based on new information from the current iteration. This correction is known as the matrix updating formula, which in general differs from method to method.

Variable metric methods therefore proceed generally by the following steps:

1. A starting point $\mathbf{x}_{0}$ is given, the gradient $\mathbf{g}_{0}$ at that point is calculated, and some approximation to $\mathbf{G}^{-1}$, say $\mathbf{V}_{0}$, is constructed. The starting $\mathbf{V}_{0}$ may be only the unit matrix, or it may actually be the inverse of the full second derivative matrix.
2. A step is taken to $\mathbf{x}_{1}=\mathbf{x}_{0}-\mathbf{V}_{0} \mathbf{g}_{0}$, which would be the minimum if $F$ were quadratic and if $\mathbf{V}_{0}$ were the true covariance matrix. Since $\mathbf{x}_{1}$ is not the position
of the true minimum in the general case, it is usual to perform a linear search along this direction, finding the $\alpha$ which minimizes $F\left(\mathbf{x}_{0}-\alpha \mathbf{V}_{0} \mathbf{g}_{0}\right)$. In either case let the new point be called $\mathbf{x}_{1}$ and let the gradient calculated at $\mathbf{x}_{1}$ be $\mathbf{g}_{1}$.
3. The matrix $\mathbf{V}$ is corrected using an updating formula of the form

$$
\mathbf{V}_{1}=\mathbf{V}_{0}+\mathbf{f}\left(\mathbf{V}_{0}, \mathbf{x}_{0}, \mathbf{x}_{1}, \mathbf{g}_{0}, \mathbf{g}_{1}\right)
$$

Then $\mathbf{g}_{0}$ is replaced by $\mathbf{g}_{1}, \mathbf{x}_{0}$ by $\mathbf{x}_{1}$, and $\mathbf{V}_{0}$ by $\mathbf{V}_{1}$, and steps (2) and (3) are repeated until some convergence criteria are satisfied.

The different methods differ chiefly in the choice of updating function $\mathbf{f}$, as described in the following sections, and in the extent to which linear minimizations are necessary. Less important variations involve the starting approximation $\mathbf{V}_{0}$ and various safeguards against "unreasonable" steps and non-positive-definiteness as for the Newton techniques.

### 4.8 Davidon's rank-two formular

Probably the first - and perhaps still the best - variable metric method was developed in 1959 by Davidon and later published in simplified form 1963 by Fletcher and Powell [9]. Davidon's updating formula for the covariance matrix is the following:

$$
\mathbf{V}_{1}=\mathbf{V}_{0}+\frac{\delta \delta^{\mathrm{T}}}{\delta^{\mathrm{T}} \gamma}-\frac{\mathbf{V}_{0} \gamma \gamma^{\mathrm{T}} \mathbf{V}_{0}}{\gamma^{\mathrm{T}} \mathbf{V}_{0} \gamma}
$$

where the changes in position and gradient on the last step were

$$
\delta=\mathbf{x}_{1}-\mathbf{x}_{0}
$$

and

$$
\gamma=\mathbf{g}_{1}-\mathbf{g}_{0}
$$

and $\mathbf{V}_{0}$ was the previous estimate of the covariance matrix. This is called the ranktwo formula since the correction $\mathbf{V}_{1}-\mathbf{V}_{2}$ is a matrix of rank two in the space of $\delta$ and $\mathbf{V}_{0} \gamma$ as can be seen directly by inspection of the formula.

One fundamental requirement of an updating formula is that the new matrix satisfies the relationship

$$
\mathbf{V}_{1} \gamma=\delta
$$

since $\gamma=\mathbf{G} \delta$ for a quadratic function with hessian $\mathbf{G}$. It is easily seen that Davidon's formula satisfies this requirement:

$$
\begin{align*}
\mathbf{V}_{1} \gamma & =\left[\mathbf{V}_{0}+\frac{\delta \delta^{\mathrm{T}}}{\delta^{\mathrm{T}} \gamma}-\frac{\mathbf{V}_{0} \gamma \gamma^{\mathrm{T}} \mathbf{V}_{0}}{\gamma^{\mathrm{T}} \mathbf{V}_{0} \gamma}\right] \gamma  \tag{4.3}\\
& =\mathbf{V}_{0} \gamma+\frac{\delta \delta^{\mathrm{T}} \gamma}{\delta^{\mathrm{T}} \gamma}-\frac{\mathbf{V}_{0} \gamma \gamma^{\mathrm{T}} \mathbf{V}_{0} \gamma}{\gamma^{\mathrm{T}} \mathbf{V}_{0} \gamma}  \tag{4.4}\\
& =\mathbf{V}_{0} \gamma+\delta-\mathbf{V}_{0} \gamma  \tag{4.5}\\
& =\delta \tag{4.6}
\end{align*}
$$

An unfortunate feature of the Davidon algorithm is the need to perform at each iteration a linear minimization along the direction given by a Newton step, $\mathbf{- V g}$. This linear search step is, however, necessary in order to assure convergence for general functions. Fletcher and Powell show in [9] that if the starting approximation to $\mathbf{V}$ is positive-definite, then $\mathbf{V}$ will remain positive-definite after all updatings, but they have to use the fact that each iteration is a linear minimization, that is

$$
\mathbf{g}_{1}^{\mathrm{T}} \mathbf{V}_{0} \mathbf{g}_{0}=0
$$

It can be shown that this method is quadratically convergent, at most $n$ iterations ( $n$ line searches and $n$ gradient calculations) being required for an $n$-dimensional quadratic form.

### 4.9 The rank-one formula

In an effort to avoid the linear minimizations required by Davidon's algorithm, several workers have independently developed an interesting updating formula of rank one. In this case Davidon in 1968 was the first to publish an algorithm [10] based on the formula, and Powell [11] has summarized the properties of this formula and of algorithms based on it.

The rank-one updating is:

$$
\mathbf{V}_{1}=\mathbf{V}_{0}+\frac{\left(\delta-\mathbf{V}_{0} \gamma\right)\left(\delta-\mathbf{V}_{0} \gamma\right)^{\mathrm{T}}}{\gamma^{\mathrm{T}}\left(\delta-\mathbf{V}_{0} \gamma\right)}
$$

It can be shown [11] that this is the only formula of rank two (or less) for which not only $\mathbf{V}_{1} \gamma=\delta$ but:

$$
\mathbf{V}_{1} \gamma_{i}=\delta_{i}
$$

where $\delta_{i}$ and $\gamma_{i}$ are the step and the gradient changes at any previous iteration. This is known as the hereditary property, sine $\mathbf{V}_{1}$ can be said to inherit the fundamental property $\mathbf{V} \gamma=\delta$ with respect to all previous iterations (up to $n$ ).

The hereditary property assures that after $n$ iterations, $\mathbf{V}_{1}$ will be the true covariance matrix if $F$ is quadratic, no matter what steps were taken (almost), so that if Newton
steps are taken, convergence for a quadratic function is assured after $n$ iterations, without the need for linear minimizations.

In addition, the rank-one formula is symmetric, in the sense that the expression for $\mathbf{V}_{1}^{-1}$ in terms of $\mathbf{V}_{0}^{-1}$ is the same as that for $\mathbf{V}_{1}$ in terms of $\mathbf{V}_{0}$ provided $\delta$ and $\gamma$ are interchanged. The meaning of this symmetry property will be discussed in the next section.

But, as nothing is perfect, so the elegance and mathematical beauty of the ran-one formula hide a number of numerical and practical difficulties which can make it highly unstable when applied to a general function. In particular, if the vector $\gamma$ happens to be orthogonal to the vector $\left(\delta-\mathbf{V}_{0} \gamma\right)$, the denominator goes to zero in the updating formula, and an unbounded correction is possible. Since these vectors may be orthogonal, even for a quadratic function, the problem of numerical instability is a serious one.

Moreover, the matrices $\mathbf{V}_{1}$ do not really converge to the true covariance matrix in the usual meaning of the term convergence. Although it is true that $\mathbf{V}_{1}$ will be equal to the true covariance matrix at the $n^{\text {th }}$ step for a quadratic function (barring numerical difficulties), the intermediate matrices $\mathbf{V}$ may vary wildly from step to step, so that on any particular iteration $\mathbf{V}_{1}$ may be a rather poor approximation. This is especially dangerous when the function is not quadratic, since the large corrections necessary in later iterations will generally not compensate properly the fluctuations in early steps. Also, there is no guarantee that intermediate matrices will remain positivedefinite, and hence no guarantee of a reduction in the value of $F$ at each step, even for a quadratic F .

All these difficulties can, of course, be overcome by programming enough safeguards into the algorithm, but this can only be done at the expense of efficiency and sometimes only by abandoning temporarily the updating formula itself, which makes it lose some of its appeal.

Different approaches are possible depending on whether it is considered important to maintain positive definiteness as in the Davidon algorithm [10], or important not to abandon the exact rank-one formula as in Powell's method [11].

### 4.10 Fletcher's unified approach to VMM

The existence of two different updating formulas with very different properties generated a lot of interest in variable metric methods (VMM) during the years 1967-1971, since it showed VMM to be very promising and left many questions unanswered, such as:

1. How can it be that the rank-one and rank-two formulas have such different properties? What is the relationship between them?
2. Is there a way to combine the best properties of both formulas?
3. Are there other good formulas? Is it posible to define a class of "admissible" formulas?

A certain understanding of the above problems was made possible by the work of a number of people. In particular, a paper by Fletcher [12] presents a unified approach to VMM, which will be given here.
Recall that the rank-one equation is symmetrical (in a sense defined in ??? 4.9), but as we shall now see, the rank-two formula is not. Indeed the asymmetry suggests a way to construct a possible third formula by taking the "mirror image" of the ranktwo formula. The basic idea is that the new formula should satisfy the fundamental relationship

$$
\mathbf{V}_{1} \gamma=\delta
$$

and therefore its inverse should satisfy

$$
\gamma=\mathbf{V}_{1}^{-1} \delta
$$

One can indeed write down the updating formula for $\mathbf{V}_{1}^{-1}$ which corresponds to the rank-two formula for $\mathbf{V}_{1}$ :

$$
\mathbf{V}_{1}^{-1}=\left(\mathbf{I}-\frac{\gamma \delta^{\mathrm{T}}}{\delta^{\mathrm{T}} \gamma}\right) \mathbf{V}_{0}^{-1}\left(\mathbf{I}-\frac{\gamma \delta^{\mathrm{T}}}{\delta^{\mathrm{T}} \gamma}\right)^{\mathrm{T}}+\frac{\gamma \gamma^{\mathrm{T}}}{\delta^{\mathrm{T}} \gamma}
$$

This matrix $\mathbf{V}_{1}^{-1}$ can now be thought of as a mapping from $\delta \rightarrow \gamma$, since $\gamma=\mathbf{V}_{1}^{-1} \delta$. If $\gamma$ and $\delta$ are interchanged in the formula, it will then give a mapping from $\gamma \rightarrow \delta$, thereby producing a new updating formula where $\mathbf{V}_{1} \gamma=\delta$. The new dual formula will be just

$$
\mathbf{V}_{1}=\left(\mathbf{I}-\frac{\delta \gamma^{\mathrm{T}}}{\delta^{\mathrm{T}} \gamma}\right) \mathbf{V}_{0}\left(\mathbf{I}-\frac{\delta \gamma^{\mathrm{T}}}{\delta^{\mathrm{T}} \gamma}\right)^{\mathrm{T}}+\frac{\delta \delta^{\mathrm{T}}}{\delta^{\mathrm{T}} \gamma}
$$

If one tries this trick with the rank-one formula, one just gets the same rank-one formula back again, since it is symmetric in this sense, or dual to itself. But with the rank-two formula, the process of inverting and interchanging yields a new formula, also of rank-two, which is also a valid updating formula in the sense that it gives rise to a quadratically convergent VMM algorithm.

In a further step consider the class of formulas which includes both rank-two and dual formulas as special cases. Introducing the notation

$$
\mathbf{V}_{1}=\mathbf{T}\left(\mathbf{V}_{0}\right) \quad \text { for the rank-two formula }
$$

and

$$
\mathbf{V}_{1}=\mathbf{D}\left(\mathbf{V}_{0}\right) \quad \text { for the dual formula }
$$

and consider the class of updating expressions as introduced by Fletcher [12]:

$$
\mathbf{V}_{\phi}=(1-\phi) \mathbf{T}+\phi \mathbf{D}
$$

where $\phi$ is some parameter which determines the exact formula. (Broyden [13], using a somewhat different notation, has also considered the same class of formulas.)

It then turns out that the rank-one formula is also in this class, with

$$
\phi(\text { rank }- \text { one })=\frac{\delta^{\mathrm{T}} \gamma}{\delta^{\mathrm{T}} \gamma-\gamma^{\mathrm{T}} \mathbf{V}_{0} \gamma}
$$

Having now constructed a wide class of updating formulas, which in fact include all formulas known to the author, it will prove interesting to consider their properties as a function of the generating parameter $\phi$. Probably the most important property, and the only one considered here, is that of monotonic convergence of $\mathbf{V}$ toward the true covariance matrix for a quadratic function. (this is called Property 1 in Fletcher's paper [12] which should be consulted for details of the definition and for theorems concerning it.) The use of an updating formula with this property will guarantee an improvement in the approximation $\mathbf{V}$ at each iteration (for a quadratic function).
Any formula $\mathbf{V}_{\phi}$ with $\phi$ in the interval $[0,1]$ possesses the monotonic convergence property. Such a formula is said to belong to the convex class of formulas. For any $\mathbf{V}_{\phi}$ with outside range $[0,1]$, there exists some quadratic function for which $\mathbf{V}$ diverges from the true covariance matrix.

From what has already been seen from the rank-one formula, it is not surprising to find that it does not belong to the convex class. Since $\delta^{\mathrm{T}} \gamma>0$ for any step which is an improvement, and since $\gamma^{\mathrm{T}} \mathbf{V}_{0} \gamma>0$ if $\mathbf{V}_{0}$ is positive-definite, it can be seen immediately from inspection of the equation for $\phi$ (rank-one) that it must either be less than zero or greater than one.

The above considerations lead Fletcher to propose an new algorithm [12] which is probably the most elegant and powerful of any VMM algorithm. Basically, he uses the general updating formula $\mathbf{V}_{\phi}$, with the value of $\phi$ chosen according to the following scheme: If $\phi$ (rank-one) $<0$, set $\phi=0$, corresponding to the usual ranktwo formula. If $\phi($ rank-one $)>1$, set $\phi=1$, corresponding to the dual formula. In this way, one always uses a formula in the convex class, and chooses that one which is "closest" to the rank-one formula. It seems that the linear searches can then be eliminated and replaced simply by Newton's steps, unless the function is highly non-quadratic. The latter condition can easily be detected by comparing the actual improvement with the expected improvement at each iteration.

## 5 Specialized techniques

All the methods outlined so far here are of rather general applicability, the only assumption being - for some methods - a predominantly quadratic behaviour in the immediate vicinity of the minimum. In order to develop more powerful methods than those already presented, one will have to give up some of this generality and exploit particular features of the functions to be minimized. In this section a few specialized techniques are discussed which are still of rather wide applicability in the sense that most functions of physical interest fall in one or more of these classes.

## $5.1 \chi^{2}$ minimization

Probably the most common application of minimization in scientific research is in least square fitting, where the function to be minimized is the sum of squares of deviations, between measured values and predictions of a model containing variable parameters:

$$
F(\mathbf{x})=\sum_{k=1}^{K} f_{k}^{2}(\mathbf{x})=\sum_{k=1}^{K}\left(\frac{Y_{k}-T_{k}(\mathbf{x})}{\sigma_{k}}\right)^{2}
$$

where $Y_{k}$ and $\sigma_{k}$ are measured values and errors, and $T_{k}(\mathbf{x})$ are the values predicted by the model, depending on some parameters $\mathbf{x}$. Minimizing $F$ then yields best values (estimates) of the $n$ parameters $\mathbf{x}$, based on $K$ measurements $Y_{k}$ with random errors $\sigma$, where $K$ must be greater than or equal to $n$, and it is usually much greater than $n$.

Now consider the second derivative matrix for $F(\mathbf{x})$, expressed in terms of the individual $f_{k}(\mathbf{x})$ :

$$
\begin{aligned}
\frac{\partial^{2} F}{\partial x_{i} \partial x_{j}} & =\frac{\partial}{\partial x_{i}} \frac{\partial}{\partial x_{j}} \sum_{k} f_{k}^{2} \\
& =\frac{\partial}{\partial x_{i}} \sum_{k} 2 f_{k} \frac{\partial f_{k}}{\partial x_{j}} \\
& =\sum_{k} 2 \frac{\partial f_{k}}{\partial x_{i}} \frac{\partial f_{k}}{\partial x_{j}}+\sum_{k} 2 f_{k} \frac{\partial^{2} f_{k}}{\partial x_{i} \partial x_{j}} .
\end{aligned}
$$

In the above r.h.s., it is usual to make the approximation that the second sum, involving second derivatives, is small compared with the first term involving products of first derivatives. This is called linearization. (Note that it is the model $\mathbf{T}(\mathbf{x})$ that is linearized, not the function $F(\mathbf{x})$.) In the important special case of linear least squares, the second sum is exactly zero, so that $F(\mathbf{x})$ is quadratic, and the whole minimization problem reduces to the inversion of the above matrix $\partial^{2} F / \partial x_{i} \partial x_{j}$ (i.e. the taking of one Newton step).

In the more general case of non-linear least squares, the linearization approximation consists in taking

$$
\frac{\partial^{2} F}{\partial x_{i} \partial x_{j}} \approx \sum_{k} 2 \frac{\partial f_{k}}{\partial x_{i}} \frac{\partial f_{k}}{\partial x_{j}} .
$$

This has the advantage of being easy to calculate and, moreover, it is always positivedefinite (under rather weak conditions such as the existence of the derivatives, and provided non-singularity). In fact in many cases the use of the above approximation in computing Newton steps is actually more effective than using the exact second derivative matrix because of the positive definiteness. Of course it must be remembered that the covariance matrix obtained by inverting this approximate matrix does not in general converge to the true covariance matrix even though the minimization based on it may converge to the true minimum.

### 5.2 Likelihood maximization

An increasingly important alternative to the least squares method in data fitting is the method of maximum likelihood. In this case the function to be minimized is of the form

$$
F(\mathbf{x})=-\sum_{k=1}^{K} \ln f_{k}(\mathbf{x})
$$

that is, a sum of logarithms. Here again, an approximation for the second derivative matrix can be found which envolves only products of first derivatives:

$$
\begin{aligned}
\frac{\partial^{2} F}{\partial x_{i} \partial x_{j}} & =-\frac{\partial}{\partial x_{i}} \frac{\partial}{\partial x_{j}} \sum_{k} \ln f_{k} \\
& =-\frac{\partial}{\partial x_{i}} \sum_{k} \frac{1}{f_{k}} \frac{\partial f_{k}}{\partial x_{j}} \\
& =-\sum_{k} \frac{1}{f_{k}^{2}} \frac{\partial f_{k}}{\partial x_{i}} \frac{\partial f_{k}}{\partial x_{j}}-\sum_{k} \frac{1}{f_{k}} \frac{\partial^{2} f_{k}}{\partial x_{i} \partial x_{j}} .
\end{aligned}
$$

As with least squares, one can neglect the second sum, involving second derivatives. In the case of the likelihood function, the second derivatives of $f$ are never exactly zero over any finite range (exactly linear maximum likelihood does not exist, essentially because the likelihood function must be normalized so that its integral over the space of measurements is independent of the parameters $\mathbf{x}$ ). However, the approximation

$$
\frac{\partial^{2} F}{\partial x_{i} \partial x_{j}} \approx \sum_{k} \frac{1}{k^{2}} \frac{\partial f_{k}}{\partial x_{i}} \frac{\partial f_{k}}{\partial x_{j}}
$$

has the same advantages as in the non-linear least square case, namely speed of calculation and assured positive-definiteness.

## 6 Local and global minima

### 6.1 The problem of multiple minima

All the methods presented so far have been designed to find a local minimum, without any consideration of whether or not other local minima exist, or whether the minimum found is actually the global minimum. If the function has more than one local minimum, there is not even any guarantee that these methods will find the minimum closest to the starting point, let alone the global minimum. In fact, it is usually assumed, when using these algorithms, that the function is unimodal (has one minimum) in the region of interest likely to be explored during the minimization.

Whenever the function may have more than one local minimum, new problems arise in addition to the problem of local minimization. First of all, the user must decide what he wants to know about the function. The following four possibilities are the most common and will be discussed here:
i) it is sufficient to know the location of any one local minimum;
ii) only the global minimum is of interest;
iii) only one minimum is of interest (the 'physical solution'), but it need not be the global minimum; or
iv) all local minima, including the global one, must be found and catalogued.

The first possibility, (i), is quite rare, but is easy to deal with, since any local minimization routine is sufficient.

Possibility (ii) is much more common, particularly in system optimization where the cost must be the smallest possible, not just small compared with other nearby solutions. Several methods exist for finding global minima, of which two will be discussed in the next sections. All such methods suffer from the absence of a stopping rule: even if the global minimum is found there is no way of recognizing it unless the function is known to be bounded and has reached its lower bound.

Possibility (iii) often arises in scientific research where the approximate values of some parameters are known in advance and one seeks a solution not too far from these values, corresponding to 'the right valley' where the function may have several faraway valleys which may be deeper. The usual technique for making sure of staying in the right valley is first to fix the approximately known parameters at their assumed values and minimize with respect to all other variables, then starting from this point minimize in the entire variable space.

Possibility (iv), of having to find and record all local minima, is the most difficult of all. It arises, for example, in energy-dependent phase-shift analyses where all 'solutions' are recorded at each energy, and a continuous set of solutions is sought, one at each energy, which have a smooth energy dependence. Although the techniques


Fig. 13
described below may help in this problem, no exhaustive method is known to the author except for the prohibitive one of using many starting points equally spaced on an n-dimensional grid.

### 6.2 The Gelfand algorithm

Relatively few minimization methods are specifically designed for non-local search in many parameters. Probably the most successful of the ad hoc stepping methods is that of Gelfand [14]. It is non-local because it provides a natural way to allow for function increases as well as decreases in any one step, while tending generally to decrease the function value.

The procedure is as follows. From the starting point $\underline{x}_{0}$, a local minimization is begun (for example along the gradient) until the function differences between steps become small (at the point $\underline{a}_{0}$ ). Then, going back to the starting point, a 'long' random step is taken to the point $\underline{x}_{1}$, and another rough local minimization is performed to reach the point $\underline{a}_{1}$ (see figure above). Then the so-called 'precipitous step' is taken along a line from $\underline{a}_{0}$ to $\underline{a}_{1}$, some distance past $\underline{a}_{1}$ to $\underline{x}_{2}$. Then from $\underline{x}_{2}$ another rough local minimization is performed, yielding $\underline{a}_{2}$, and another precipitous step is taken from $\underline{a}_{1}$ past $\underline{a}_{2}$ to $\underline{x}_{3}$ and the search continues in this way.

The choice of the 'precipitous step' length is important in determining whether the method will 'roll over small ridges, but skirt a high mountain', as its authors say it should. But no precise way is given, except that 'the choice of the length of the precipitous step is carried out experimentally (by trials) and it constitutes an important charactistic of the function'.

Moreover, there is no stopping rule, since the method is essentially searching rather than converging. In practice one usually stops after a given length of computer time, but one would also stop if the program went around in circles repeating itself (which is very possible but not so easy to detect) or if a predetermined 'acceptably small' function value was attained. This problem of stopping seems to be common to all non-local minimization methods.

### 6.3 The Goldstein-Price method

Goldstein and Price [15] have proposed an elegant yet simple method for seeking other local minima after one local minimum has been found It is based on a consideration of the analytic (Taylor series) properties of the function. Let us assume that the function can be represented as a Taylor series about a local minimum $\underline{x}_{1}$, where the first derivatives vanish:

$$
F(\underline{x})=F(\underline{x})_{1}+\frac{1}{2}\left(\underline{x}-\underline{x}_{1}\right)^{T} \mathbf{G}\left(\underline{x}-\underline{x}_{1}\right)+\text { h.t. . }
$$

Now the higher terms (h.t.), involving third and higher derivatives, are important since these are the terms that will give rise to other local minima. In fact, we seek a way of transforming the function so that only the higher terms remain. Such a transformed function is $F_{1}$ such that:

$$
F_{1}\left(\underline{x}_{1}, \underline{x}\right)=\frac{2\left(F(\underline{x})-F\left(\underline{x}_{1}\right)\right)}{\left(\underline{x}-\underline{x}_{1}\right)^{T} \mathbf{G}\left(\underline{x}-\underline{x}_{1}\right)}=1+\text { h.t. }
$$

By means of this transformation, we have 'removed' the minimum at $\underline{x}_{1}$, and the way is cleared to search for other minima generated by the higher terms of the expansion about $\underline{x}_{1}$. The method therefore consists of seeking a local minimum of the function $F_{1}$ (It is required to know the second derivative matrix $\mathbf{G}$ at the local minimum $\underline{x}_{1}$.) Since the quadratic form $\left(\underline{x}-\underline{x}_{1}\right)^{T} \mathbf{G}\left(\underline{x}-\underline{x}_{1}\right)$ is always positive for positive-definite G, th efunction $F_{1}$ will become negative as soon as an improvement on $\underline{x}_{1}$ is found. Then starting from this improved point, the original function $F$ can be minimized locally to yield a new, improved local minimum of $F$.

If the minimum value found for $F_{1}$ is positive, then it may correspond to a new local minimum of $F$, but not an improvement over $\underline{x}_{1}$.

In this case the procedure may be continued from this new point, forming a new function $F_{2}$, related to $F_{1}$ just as $F_{1}$ was related to $F$. As usual, no stopping rule is given by the theory.

The method seems to work in practice, although experience with it is limited and no conditions are known under which it is guaranteed to work. It is appealing for reasons of its elegance and simplicity, and could prove to be an important tool in global minimization.

## 7 Some sample problems for minimization routines

Here a collection of test problems is assembled which were found to be useful in verifying and comparing minimization routines. Many of these are standard functions upon which it has become conventional to try all new methods, quoting the performance in the publication of the algorithm.

### 7.1 Rosenbrock's curved valley

start point:

$$
F(x, y)=100\left(y-x^{2}\right)^{2}+(1-x)^{2}
$$

$$
F(-1.2,1.0)=24.20
$$

minimum:

$$
F(1.0,1.0)=0
$$

This narrow, parabolic valley is probably the best known of all test cases. The floor of the valley follows approximately the parabola $y=x^{2}+1 / 200$, indicated by the dashed line in fig. 14. In the cross-hatched area above the dashed line, the covariance matrix is not positive-definite. On the dashed line it is singular. Stepping methods tend to perform at least as well as gradient methods for this function.
[Reference: Comput. J. 3, 175 (1960).]

### 7.2 Wood's function in four parameters

$$
\begin{aligned}
& \qquad \qquad(w, x, y, z)=\begin{array}{l}
100\left(x-w^{2}\right)^{2}+(w-1)^{2}+90\left(z-y^{2}\right)^{2} \\
+(1-y)^{2}+10.1\left[(x-1)^{2}+(z-1)^{2}\right] \\
+19.8(x-1)(z-1)
\end{array} \\
& \text { start point: } \quad F(-3,-1,-3,-1)=19192 \\
& \text { minimum: } \\
& \\
& \quad F(1,1,1,1)=0 .
\end{aligned}
$$

This is a fourth-degree polynomial which is reasonably well-behaved near the minimum, but in order to get there one must cross a rather flat, four-dimensional 'plateau' which often causes minimization algorithm to get 'stuck' far from the minimum. As such it is a particularly good test of convergence criteria and simulates quite well a feature of many physical problems in many variables where no good starting approximation is known.
[Reference: Unpublished. See IBM Technical Report No. 320-2949.]


Fig. 14

### 7.3 Powell's quartic function

$$
F(w, x, y, z)=(w+10 x)^{2}+5(y-Z)^{2}+(x-2 y)^{4}+10(w-z)^{4}
$$

start point:
$F(3,-1,0,1)=215$
minimum:

$$
F(0,0,0,0)=0
$$

This function is difficult because its matrix of second derivatives becomes singular at the minimum. Near the minimum the function is given by $(w+10 x)^{2}+5(y-5)^{2}$ which does not determine the minimum uniquely.
[Reference: Comput. J. 5, 147 (1962).]

### 7.4 Fletcher and Powell's helical valley

$$
F(x, y, z)=100\left\{[z-10 \Psi(x, y)]^{2}+\left(\sqrt{x^{2}+y^{2}}-1\right)^{2}\right\}+z^{2}
$$

where

$$
\begin{aligned}
2 \pi \Psi(x, y) & =\arctan (y / x) & \text { for } x>0 \\
& =\pi+\arctan (y / x) & \text { for } x<0
\end{aligned}
$$

start point:

$$
F(-1,0,0)=2500
$$

minimum:

$$
F(1,0,0)=0 .
$$

$F$ is defined only for $-0.25<\Psi<0.75$.
This is a curved valley problem, similar to Rosenbrock's, but in three dimensions. [Reference: Comput. J. 6, 163 (1963).]

### 7.5 Goldstein and Price function with four minima

$$
\begin{aligned}
F(x, y)= & \left(1+(x+y+1)^{2} *\left(19-14 x+3 x^{2}-14 y+6 x y+3 y^{2}\right)\right) \\
& *\left(30+(2 x-3 y)^{2} *\left(18-32 x+12 x^{2}+48 y-36 x y+27 y^{2}\right)\right)
\end{aligned}
$$

local minima: $\quad F(1.2,0.8)=840$

$$
F(1.8,0.2)=84
$$

$$
F(-0.6,-0.4)=30
$$

global minimum:

$$
F(0,-1.0) \quad=3 .
$$

This is an eighth-order polynomial in two variables which is well behaved near each minimum, but has four local minima and is of course non-positive-definite in many regions. The saddle point between the two lowest minima occurs at $F(-0.4,-0.6)$ $=35$, making this an interesting start point.
[Reference: Math. Comp. 25, 571 (1971).]

### 7.6 Goldstein and Price function with many minima

$F(x, y)=\exp \left\{\frac{1}{2}\left(x^{2}+y^{2}-25\right)^{2}\right\}+\sin ^{4}(4 x-3 y)+\frac{1}{2}(2 x+y-10)^{2}$
global minimum:
$F(3,4)=1$.
This function has 'many' local minima.
[Reference: Math. Comp. 25, 571 (1971).]

### 7.7 Quadratic function in four parameters

$$
F(x, y, z, w)=\frac{1}{70}\left(21 x^{2}+20 y^{2}+19 z^{2}-14 x z-20 y z\right)+w^{2}
$$

minimum:

$$
F(0,0,0,0)=0
$$

covariance matrix:

$$
\left(\begin{array}{llll}
4 & 1 & 2 & 0 \\
1 & 5 & 3 & 0 \\
2 & 3 & 6 & 0 \\
0 & 0 & 0 & 0
\end{array}\right)
$$

Except for the reasonably strong parameter correlations, this function poses no special problem to any minimization routine. It was found useful in debugging programs based on quadratically convergent methods, since these programs should minimize the function exactly in one iteration. It is also used to check the calculation of the covariance matrix.

A variation consists of adding $|x|^{3}-1$ whenever $|x|>1$, and similarly with the other variables. This introduces in a reasonably smooth way terms which alter the quadratic behaviour far from the minimum while leaving it unchanged inside the unit cube, thus providing a test for those methods which are supposed to converge to the correct covariance matrix by updating.

### 7.8 Chebyquad

$$
F(\vec{x})=\sum_{i=1}^{n}\left\{\int_{0}^{1} T_{i}\left(x^{\prime}\right) d x^{\prime}-\frac{1}{n} \sum_{j=1}^{n} T_{i}\left(x_{j}\right)\right\}^{2}
$$

where $T_{i}(x)$ are shifted Chebyshev polynomials of degree i;
start point: $\quad x_{j}=j /(n+1)$.
This function is designed to have a variable and possibly large number of parameters, and to resemble functions encountered in actual practice rather than being contrived to be especially difficult. Each term of $F$ represents the squared difference between the true integral of a polynomial of degree $i$ and the integral estimated by Chebyshev (equal-weight) quadrature on $n$ points:

$$
\int_{0}^{1} P(x) d x \approx \frac{1}{n} \sum_{j=1}^{n} P\left(x_{j}\right)
$$

The starting values correspond to equally spaced points $x_{j}$ which is not too far away from the solution. Fletcher gives a complete Algol-coded, procedure for this function
in the reference quoted below.
[Reference: Comput. J. 8, 33 (1965).]

### 7.9 Trigonometric functions of Fletcher and Powell

$$
F(\vec{x})=\sum_{i=1}^{n}\left\{E_{i}-\sum_{j=1}^{n}\left(A_{i j} \sin x_{j}+B_{i j} \cos x_{j}\right)\right\}^{2}
$$

where

$$
E_{i}=\sum_{j=1}^{n}\left(A_{i j} \sin x_{0 j}+B_{i j} \cos x_{0 j}\right)
$$

$B_{i j}$ and $A_{i j}$ are random matrices composed of integers between -100 and 100; for $j$ $=1, \ldots, n: x_{0 j}$ are any random numbers, $-\pi<x_{0 j}<\pi$;
start point:

$$
\begin{aligned}
& x_{j}=x_{0 j}+0.1 \delta_{j},-\pi<\delta j<\pi \\
& F\left(\vec{x}=\vec{x}_{0}\right)=0
\end{aligned}
$$

minimum:
This is a set of functions of any number of variables $n$, where the minimum is always known in advance, but where the problem can be changed by choosing different (random) values of the constants $A_{i j}, B_{i j}$, and $x_{0 j}$. The difficulty can be varied by choosing larger starting deviations $\delta_{j}$. In practice, most methods find the 'right' minimum, corresponding to $\vec{x}=\vec{x}_{0}$, but there are usually many subsidiary minima. [Reference: Comput. J. 6163 (1963).]

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