Fast Computation of Equispaced Pareto Manifolds and Pareto Fronts for Multiobjective Optimization Problems

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Fast Computation of Equispaced Pareto Manifolds and Pareto Fronts for Multiobjective Optimization Problems *

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Abstract

In this paper we consider the problem of generating a well sampled discrete representation of the Pareto manifold or the Pareto front corresponding to the equilibrium points of a multi-objective optimization problem. We show how the introduction of simple additional constraints into a continuation procedure produces equispaced points in either of those two sets. Moreover, we describe in detail a novel algorithm for global continuation that requires two orders of magnitude less function evaluations than evolutionary algorithms commonly used to solve this problem. The performance of the methods is demonstrated on problems from the current literature.

1 Introduction

Multi-objective optimization is becoming a common tool in Engineering and Scientific applications. Most optimization problems in industry are multi-objective, non-linear, constrained and multi-modal, i.e., very tough.

Multi-objective optimization has a long history, especially in areas such as decision and game theory. It is only in the last twenty years that it has become more popular in Engineering and Scientific applications. Many books describe a myriad of approaches to the "solution" of these problems [4, 23, 8, 6, 22, 13]. The quotes are used to emphasize that the most common situations lead to a continuum of equilibrium points, which in the absence of additional information are all valid "solutions" to the problem. That is where many of the different approaches appear, attempting to decide which of these infinity many possibilities should be chosen. Thus, many methods use additional "expert" or subjective information to guide the search towards a particular solution [14, 21].

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In recent times, with the increase in computing power, it has become possible to attempt to generate a complete discrete approximation to the Pareto manifold (set of equilibrium points) or its image by the objectives, the Pareto front. Some of these methods use scalarization as their basic tool, i.e., minimization of a linear combination of the objectives is employed to generate Pareto points. However, since the mapping from weight space to input or output space can be very nonlinear, different strategies need to be devised to obtain a well sampled set of solutions.

Evolutionary and genetic algorithms (GA) are one class of powerful methods that has been favoured in recent times for their robustness, specially in the versions that permit to calculate a discrete representation of the Pareto manifold [3]. The negative side to these methods is the number of function evaluations required to obtain a reasonable accuracy, which grows exponentially with the dimension of the design parameter space. This is totally inadequate for realistic high fidelity design applications, where function evaluations can be very costly. They still can be useful if one replaces these expensive evaluations by surrogates. These methods do not use linear aggregation.

In this paper we show that the Pareto manifold for a convex bi-objective problem can be approximated by solving numerically a two-point boundary value problem and from this insight we mimic technics for the solution of such problems to obtain a continuation method that updates a whole discrete representation of the Pareto manifold while maintaining and even spacing between solutions. As an additional bonus this procedure is easily parallelizable. We also consider for comparison a simple continuation procedure and an enhanced version that includes a equispacing constraint. We will see that this last method is capable of obtaining equispaced points in the non-convex parts of the Pareto front and it is also useful to initialize the global method when this fails to converge from poor initial guesses, specially for highly curved Pareto fronts. The simple implementations we have devised to produce the results of this paper are not production codes but give us a quick way to show that this approach works in some test cases taken from the literature.

In [2], a method that is not based on GA is presented. This method is most akin to ours in terms of economy of function evaluations. It is pointed out in that paper that it is very difficult to guarantee an uniform spread of points in the Pareto manifold or in the Pareto front when using the popular method of solving many single optimization problem for convex combinations of the objectives. We will show that the methods of this paper do just that without any guessing and in a manner as economical as the method proposed in that reference, but with a simpler algorithm.

The full front continuation described in this paper does not suffer from the disadvantages pointed out in [2], with regards to usual initial value continuation, such as that of [18].

We discuss in this paper the unconstrained case to understand the geometry of the problem and that of the optimality conditions. In a companion paper we will extend these ideas to the constrained case. The methods extend naturally to problems with more than two objectives.
2 Optimality Conditions

The multiobjective optimization problem is defined as:

$$\min_{x \in D} F(x),$$  \hspace{1cm} (1)

where \( x \in \mathbb{R}^n \), \( F = \{ f_1(x), \ldots, f_k(x) \} \in \mathbb{R}^k \) and \( D \subset \mathbb{R}^n \) is defined by a set of constraints.

One of the main features that distinguishes multi-objective from single-objective optimization is that it is very unlikely that there will be an \( x^* \) that is a minimizer for each and every objective. In fact, these problems are characterized for the requirement of a subjective trade-off between conflicting objectives. The optimality concept here is known as Pareto (or Nash, Edgeworth) equilibrium, which in words establishes that \( x^* \) is a global Pareto equilibrium point if there is no other point that is dominated by \( x^* \).

A point \( x \) is dominated by a point \( y \) iff \( f_i(y) \leq f_i(x) \), with strict inequality for at least one of the objectives. Thus, a global Pareto equilibrium point is such that no improvement for all objectives can be achieved by moving to any other feasible point. A local version of this concept is obtained if we limit the movement to an open neighborhood around the optimal point.

For differentiable convex objectives we can use the usual geometrical concepts of single objective optimization to arrive at an analytical characterization of the local Pareto points. This is easily seen first for the unconstrained case of two objectives and two independent variables. Let \( x_1^*, x_2^* \) be local minima of \( f_1 \) and \( f_2 \) respectively. Then a segment of the Pareto manifold is a curve that joins these two points and is defined as the parametric set of solutions \( x^*(\lambda) \) of the first order necessary optimality condition:

$$G(x(\lambda); \lambda) = (1 - \lambda)\nabla f_1(x) + \lambda \nabla f_2(x) = 0, \hspace{0.5cm} 0 \leq \lambda \leq 1.$$ \hspace{1cm} (2)

Geometrically, (2) says that a point is Pareto optimal if the contours of the two objectives are tangent at it, with gradients pointing in opposite directions; i.e., the two functionals have no descent directions in common (see Figure 1 for a cartoon description). The image by \( F \) of the Pareto manifold is called the Pareto front.

3 A Differential Equations Approach to Finding a Discrete Representation of the Pareto Manifold

By differentiating \( G \) with respect to \( \lambda \) we obtain:

$$x_\lambda = -H^{-1}(x) G_\lambda(x), \hspace{0.5cm} \text{subject to } x(0) = x_1^*, \hspace{0.5cm} x(1) = x_2^*.$$ \hspace{1cm} (3)
This is a two-point boundary value problem for a set of $n$ first order ordinary differential equations in the independent variable $\lambda$. Here

$$G_\lambda = \nabla f_2(x) - \nabla f_1(x), \quad H(x) = G_x = (1 - \lambda) \; H_1(x) + \lambda \; H_2(x),$$

with $H_i$ the Hessian matrix of $f_i$. In order for this global result to hold we must have of course that $H(x(\lambda))$ is nonsingular along the Pareto manifold. For the non-convex case, segments of the Pareto manifold will join each pair of minima of the functionals $f_i$.

In order to obtain a discrete representation of a segment of the Pareto manifold we can solve problem (3) numerically. Since we also want an uniform sampling of this manifold segment, we can require a mesh refinement equidistribution of the arc length as additional constraints. An initial approximation to the trajectory can be obtained by the straight line joining the two minima:

$$x_i^{(0)} = x^{(0)}(\lambda_i) = (1 - \lambda_i) \; x_1^* + \lambda_i \; x_2^*, \quad \lambda_i = i \ast \delta \lambda, \quad i = 0, ..., \; l + 1. \quad (4)$$

Unfortunately we will need still another derivative to apply Newton’s method to solve the resulting nonlinear system of algebraic equations that arises when using global discretization methods to solve this 2PBVP.

Although this is an elegant approach that sheds some light on the potential structure of the Pareto manifold, it may not be too practical, since it requires third order derivatives, so we go back to (2) and consider standard continuation [19], combined with a direct application of Newton’s method, in case we have the required differentiability.

Starting from $x_1^*$, i.e., $\lambda = 0$, we solve successively $G(x(\lambda_i); \; \lambda_i) = 0$, with $\lambda_i$ defined as above, although an adaptive algorithm is also possible and probably
wise (see for instance [7]). Newton’s method applied to the above equation for a given $\lambda_i$ can be written as:

$$ x^{j+1} = x^{(j)} - G_{x}^{-1}(x^{(j)}(\lambda_i)) \cdot G(x^{(j)}(\lambda_i)) , \quad j = 0, \ldots $$

Of course, we write the above iterative step as a system of linear equations with matrix $G_{x}$. Singularity of $G_{x}$ indicates possible bifurcation of the solution path. Another potential problem, turning points, should not affect this parametrization (arc length). If $x^* (\lambda_i)$ is the approximate solution of the above equation, then we use it as initial value for the next one: $x^{(0)}(\lambda_{i+1}) = x^*(\lambda_i)$.

For bi-objective problems one can also use a general continuation code for solving systems of nonlinear equations, such as [19], or one that applies directly to the optimization problem [18], although we have preferred to use a simple implementation of Newton’s method with constant step continuation, just for the sake of a rapid comparison.

4 Parallel Pareto Manifold Calculation

Although the reduction to a TPBVP is not an attractive proposition, applying ideas from the numerical solution of such problems by finite differences [15, 12], we can develop a novel continuation approach for convex problems that will deform an initial discrete guess of a convex Pareto manifold into its actual target, maintaining an equidistant discretization. One of the most attractive features of such an approach is parallelization, which is not possible with regular continuation from one initial solution, an essentially sequential process. A different approach with apparently similar amount of work can be found in [2].

Going back to problem (1) and the resulting system of optimality conditions (2), we consider the initial mesh of approximate solutions $x^{(0)}_i (\lambda_i)$ defined in (4). Instead of the standard sequential continuation approach mentioned above we consider now deforming the whole set of solutions until it matches the Pareto manifold with enough accuracy.

One desirable feature, emphasized in algorithms such as those of [3, 2], is to have an equidistant representation of the Pareto manifold. Observe that we start with such a mesh, but in order to guarantee that we end with an approximately uniform representation we will also solve for the parametrization $\lambda_i$, insisting that the spacing (essentially arc length along the manifold) stays uniform, by imposing the additional conditions:

$$ L_i(x, \lambda) = |x_i - x_{i-1}|^2 \cdot P / (l + 1) = 0, $$

where $P = \sum |x_j - x_{j-1}|^2$. Some authors do not this kind of approach into the linear aggregation methods, meaning simply that the parameter $\lambda_i$ is not prescribed but is calculated along with the solution $x_i$.

Although this global approach seems to lead to a formidable system of nonlinear equations, it is a very structured one and careful crafting of the solution algorithm results in a reasonable floating point count.
Applying Newton’s linearization to the system (2-4) we obtain,

\[
E \begin{bmatrix}
\Delta x \\
\Delta \lambda
\end{bmatrix} = \begin{bmatrix}
-G_1 \\
-G_2 \\
\vdots \\
-G_n \\
-\delta x_0 + \frac{p}{(l+1)^2} \\
-\delta x_{l-1} + \frac{p}{(l+1)^2}
\end{bmatrix},
\]

where \( \delta x_i = ||x_i - x_{i-1}||^2 \), and

\[
E = \begin{bmatrix}
G_{x_1} & 0 & 0 & G_{x_2} & 0 & 0 \\
0 & G_{x_2} & 0 & 0 & G_{\lambda_2} & 0 \\
0 & G_{x_1} & G_{\lambda_1} & 0 & 0 & 0 \\
L_{1x_1} & L_{1x_2} & 0 & 0 & 0 & 0 \\
L_{lx_2} & L_{lx_1} & 0 & 0 & 0 & 0
\end{bmatrix}.
\]

Or in block matrix form:

\[
\begin{bmatrix}
A & B \\
C & 0
\end{bmatrix} \begin{bmatrix}
\Delta x \\
\Delta \lambda
\end{bmatrix} = \begin{bmatrix}
a \\
b
\end{bmatrix},
\]

where \( A = \text{diag}(G_{x_1}) \), \( B = \text{diag}(G_{\lambda}) \), and \( a \), \( b \) are the corresponding right hand side vectors from (5).

This \( 2 \times 2 \) block system has a special structure that we will exploit. The lower block is full and of size \( l \times n.l \). The upper blocks are diagonal with a total of \( l \times l \) blocks each. The first one has blocks of size \( n \times n \), while the second one has blocks of size \( n \times 1 \), for a whole matrix of size \( (n+1)l \times (n+1)l \). The partial Frechet derivatives corresponding to the \( L \)–equations are:

\[
\frac{\partial L_m}{\partial x_i} = \begin{bmatrix}
2(x_i - x_{i-1}) + \eta_i, & i = m \\
-2(x_i - x_{i-1}) + \eta_i, & i = m - 1 \\
\eta_i, & \text{all other } i \text{'s}
\end{bmatrix},
\]

where \( \eta_i = -\frac{2}{l+1}(2x_i - x_{i-1} - x_{i+1}) \).
We use block Gaussian elimination to solve this $2 \times 2$ block system.

We first eliminate $C$ by multiplying the first row by $C \text{ diag}(G_{x_1}^{-1}) = C G_{x_1}^{-1}$, and replacing the second row by its difference with the modified first row, thus obtaining a 0 matrix in the low left corner and the Schur complement in the right one:

$$
D^* = CG_{x_1}^{-1}G\lambda, \\

b^* = -CG_{x_1}^{-1}G - b.
$$

Then back-substituting:

$$
\Delta \lambda = D^{*-1}b^*,
$$

and finally:

$$
\Delta x_i = -G_{x_i}^{-1}G\lambda\Delta \lambda_i - G_{x_i}^{-1}G_\lambda.
$$

Since all these last systems are independent, they can be solved in parallel. In fact, for each $i$ we factorize $G_{x_i}$ and then solve the two systems:

$$
G_{x_i}c_i = G_\lambda, \quad G_{x_i}d_i = G_{\lambda_i},
$$

to obtain:

$$
\Delta x_i = -d_i\Delta \lambda_i + c_i.
$$

With these corrections we step the solutions:

$$
x_i^{(j+1)} = x_i^{(j)} + \Delta x_i, \\

\lambda_i^{(j+1)} = \lambda_i^{(j)} + \Delta \lambda_i.
$$

It is wise to control the Newton step to increase its convergence region. In particular we should insist in the monotonicity of the parameters $\lambda_i$, which can also be enforced by controlling the length of the Newton step.

We observe that Gaussian elimination applied directly to the full system, ignoring its structure, would require $(n+1)^3l^3/3$ operations, while the proposed algorithm has complexity $(2l+1)n^3/3 + l^3/3$, which is substantially simpler for large $l$ (the number of Pareto points). The parallel algorithm also speeds up with increasing $l$. 

7
An uniform discrete representation of the Pareto front

In the previous section we showed how to obtain a uniform discrete representation of the Pareto manifold, i.e., a set of equally spaced Pareto equilibrium points in design parameter space. However, it is usually simpler to inspect and make decisions by looking at the Pareto front, which is the image of the Pareto manifold in objective space. Unfortunately, equal spacing in design space does not guarantee equal spacing in objective space. If that is the goal, then it needs to be stated explicitly by replacing conditions (4) by:

\[ L_i = \| \mathbf{F}(\mathbf{x}_i) - \mathbf{F}(\mathbf{x}_{i-1}) \|^2 - \| \mathbf{F}(\mathbf{x}_i) - \mathbf{F}(\mathbf{x}_{i-1}) \|_2^2 \frac{(l+1)}{(l+1)} = 0, \quad i = 1, ..., l. \]

This of course, requires a re-calculation of the Jacobian of the resulting system (albeit, only its lower block):

\[
\frac{\partial L_m}{\partial x_i} = \begin{cases} 
2[f_1(x_i) - f_1(x_{i-1})]\nabla f_1(x_i) + 2[f_2(x_i) - f_2(x_{i-1})]\nabla f_2(x) + \eta_i, & i = m \\
-2[f_1(x_{i+1}) - f_1(x_i)]\nabla f_1(x_i) -2[f_2(x_{i+1}) - f_2(x_i)]\nabla f_2(x) + \eta_i, & i = m - 1 \\
\quad \eta_i \quad \text{all other } i's
\end{cases}
\]

where \( \eta_i = -\frac{2}{(l+1)}\frac{2 [f_1(x_i) - f_1(x_{i-1}) - f_1(x_{i+1})]\nabla f_1(x_i) + [2 f_2(x_i) - f_2(x_{i-1}) - 2 f_2(x_{i+1})]\nabla f_2(x_i). \)

Finally,

\[ b_m = -L_m. \]

5 The general case

We like to point out to the dual aspect of the differential equation (1), and similarly to the continuation system of the previous section, with regards to the original problem.

From Table 1 one can see that the complexity of calculating the Pareto manifold as the number of input or design variables increases implies more differential
(or algebraic) equations, while an increase in the number of objectives beyond 2 leads to partial differential equations (or continuation with multiple parameters) in spaces of increased dimensionality. Observe that for the bi-objective problem the Pareto manifold is always a curve in the space of input variables parameterized by \( \lambda \).

The continuation method can be extended to the more than 2 objectives case although, as mentioned, it will have increased complexity (see [20]). An alternative method is presented in [2].

Our approach can also be extended at the price of introducing more constraints, in order to define an uniformly spaced mesh of solutions in the \( k-1 \)–dimensional Pareto manifold or in the Pareto front. For instance, for 3 objectives and a \( l \times l \) mesh of Pareto points, we would have to solve the following set of equations:

\[
G(\mathbf{x}(\lambda), \lambda) = \lambda_1 \nabla f_1 + \lambda_2 \nabla f_2 + \lambda_3 \nabla f_3, \quad \lambda_1 + \lambda_2 + \lambda_3 = 1,
\]

\[
\left\| \mathbf{x}_{i,j} - \mathbf{x}_{i-1,j} \right\|^2 - \frac{1}{(l+1)^2} \sum_{s=0}^{l} \sum_{t=0}^{l} \left\| \mathbf{x}_{s,t} - \mathbf{x}_{s-1,t} \right\|^2 + \left\| \mathbf{x}_{s,t} - \mathbf{x}_{s,t-1} \right\|^2 = 0,
\]

\[
i = 0, ..., l, \quad j = 0, ..., l,
\]

with similar equations in the jth direction and the distances to points outside the mesh equated to zero.

For the algorithm to obtain an equispaced representation of the Pareto front, the distance constraints should be imposed in objective space.

The method also extends to problems with constraints, by using the complete Lagrangian. It is probably wiser to use well proven nonlinear programming codes to solve each of the individual constrained optimization problems, after effecting the first step of eliminating the \( \lambda \)'s. The equidistance constraints can then be naturally included in the nonlinear program. Parallelization can be achieved by decoupling the problems for different points and keeping a data base of current values of the solutions that is updated asynchronously by each optimization process. One then uses whatever values are present in the data base to calculate the couplings between equations. This will lead to a linearly convergent algorithm which will be protected from individual processor failures (see [16]).

Quasi-Newton approximations can be used if derivative information is not readily available. Direct search methods are also plausible, although we will again be moving in the undesirable direction of large number of evaluations.

In Das and Dennis [2] it is shown the equivalence between their method and those that minimize linear combinations of the objectives, such as the methods that we are presenting here. Thus, a fortiori, we can say with Das and Dennis that these methods are independent of scalings of the objectives, which essentially amount to a change in the parametrization of the Pareto set.
Examples

We will compare four different methods:

1. Global front continuation with equispacing constraints on the Pareto manifold.
2. Global front continuation with equispacing constraints on the Pareto front.

Method 1 was explained in detail in Section 4. Method 2, a modification of the distance constraints of Method 1 was explained in Section 4.

Method 3 was explained briefly in Section 3. It is used with constant step in \( \lambda \). It should be able to obtain convex segments of the Pareto manifold, but we do not expect it to sample it uniformly.

Method 4 combines Newton continuation with the front equispacing constraints to calculate simultaneously \((x, \lambda)\), so that \(||F(x_j) - F(x_{j-1})||_2^2 = cte\). The cte. should be the quotient of the length of the Pareto front divided by the number of points. Since we do not know the length of the Pareto front \( a \) \( pri \) or, we guess it to be somewhat larger than the distance between the two end points, attempting to account for the curvature of the front and the parameterization mapping.

We consider the first test problem (SCH) in [3].

\[
 \begin{align*}
 f_1(x) &= x^2, \\
 f_2(x) &= (x - 2)^2.
 \end{align*}
\]

This problem is obviously trivial. The unique minima of the goal functionals are: \( x_1^* = 0, \ x_2^* = 2 \), and the Pareto manifold is the whole interval \([0, 2]\). Thus the problem we need to solve (2-4), has \( 2 * l \) equations and \( 2 * l \) unknowns, in order to generate \( l \) points equally spaced in the Pareto manifold. We stop when the RMS is less than \( 10^{-6} / \sqrt{(2 * l)} \). Also, the mapping from \( \lambda \) to \( x(\lambda) \) is trivial and preserves equidistance: i.e., a uniform mesh in \( \lambda \) will results on a uniform spacing both on input and goal space, so all methods work well, as we will see below.

Since taking the initial values (4) would lead in this case to the solution of the problem, we perturb them so that \( x \) and \( \lambda \) are chosen at random within 0.5 of those values.

For most problems, Newton’s method takes a maximum of four iterations to reduce RMS below \( 10^{-6} \). Since each iteration requires one evaluation of the vector goal functional and its first and second derivatives (all trivial in this...
problem), we can say in the language of evolution algorithms that the method requires 4 generations of a population with l points, to converge to high precision, in contrast with evolutionary algorithms that usually require hundreds of generations to converge to the Pareto front (see [3]).

Observe that, as expected and anticipated, for the method described in Section 4, the points in the Pareto front are not equispaced, although they are equispaced in the Pareto set. However, in this simple problem the Pareto front sampling is quite satisfactory. The following Figure shows the results with the equispaced constraints on the Pareto front (30 points), which of course are perfectly spaced by construction.

It is interesting to contrast these results on a trivial problem with the evolutionary methods of [3] as shown below. We see that the evolutionary methods, specially the most recent algorithm NSGA-II, which are extremely costly in terms of function evaluations, do a good job in sampling the front, although they are not perfectly evenly spaced as we saw above and leave a number of gaps in the coverage, besides of requiring several orders of magnitude more functional evaluations.

Results for problem SCH of Deb et al [3]. Simple continuation with equispacing in the Pareto set.
Figure 2: Problem SCH with global continuation

According to [3], the results for NSGA-II in Figure 3 were obtained using a population of 100 points and a maximum of 250 generations, or 25,000 function evaluations! They are compared with an earlier method (PAES) to show its improved sampling capability.

It turns out that Method 4, although based on linear aggregates of the objectives, is capable of obtaining points in the non-convex part of the Pareto front, as we see in the next non-convex example (FON), also from [3]:

\[
\begin{align*}
  f_1(x) &= 1 - e^{-\sum_{i=1}^{3}(x_i - 1/\sqrt{3})^2} , \\
  f_2(x) &= 1 - e^{-\sum_{i=1}^{3}(x_i + 1/\sqrt{3})^2} .
\end{align*}
\]

Observe that the simple continuation not only produces irregularly spaced Pareto points, but also only covers a small part of the initial convex segment (beware of the different scales in the two graphs!), while Method 4 has been able to continue through the inflection point and into the concave part of the front, besides of producing a nicely equispaced sample. This seems to contradict the results of [11], and therefore it requires a closer inspection. We are missing one point at the end since we did not estimate the length of the Pareto front accurately enough.

A more challenging problem is POL, also from [3], since it is non-convex and disconnected:

\[
  f_1(x) = 1 + (A_1 - B_1)^2 + (A_2 - B_2)^2 ,
\]
Fig. 6. NSGA-II finds better spread of solutions than PAES on SCH.

Figure 3: Results for SCH from [3]
Problem FON: Simple Newton continuation

Problem FON: continuation with equispacing

Figure 4: Problem FON. Both methods used 30 points.
\[ f_2(x) = (x_1 + 3)^2 + (x_2 + 1)^2, \]

where,

\[ A_1 = 0.5 \sin 1 - 2 \cos 1 + 2 \sin 2 - 1.5 \cos 2, \]

\[ A_2 = 1.5 \sin 1 - \cos 1 + 2 \sin 2 - 0.5 \cos 2, \]

\[ B_1 = 0.5 \sin x_1 - 2 \cos x_1 + \sin x_2 - 1.5 \cos x_2, \]

\[ B_2 = 1.5 \sin x_1 - \cos x_1 + 2 \sin x_2 - 0.5 \cos x_2. \]

Here we see that the methods correctly stop when reaching the end of the left segment and again, plain Newton does a poor job with the spacing while Method 4 gives a nicely sampled front segment.

Now we consider a simplified version of Example 1 of [20]:

\[ f_1(x) = (x_2 - 1)^2 + (x_3 - 1)^2 + (x_1 - 1)^4, \]

\[ f_2(x) = (x_1 + 1)^2 + (x_3 + 1)^2 + (x_2 + 1)^4. \]

This is a convex problem, but as we see below, the Pareto front has high curvature, thus the global continuation procedure equispaced in input space (Pareto manifold) does not produce an equispaced Pareto front, while the shooting method with equispacing condition on the front does. However, we still have a problem in getting the last points in the front, although we have tried to adjust the step.

Unfortunately, Method 2 that would be the solution to these two difficulties, does not seem to work for this problem, when starting from the simple guess of equispaced points on the straight line joining the minima of the two objectives. However, if we use as initial guess the values obtained by the marching continuation (Method 4), we can successfully calculate an equispaced representation of the Pareto front.

Finally we consider a problem with 10 input variables to see what is the impact on performance. We chose initially problem ZDT4 from [3], but this problem is non-convex and \( f_1(x) = x_1 \) has a minimum at \(-\infty\). The problem
Problem POL: Simple Newton continuation

Problem POL: Newton continuation with equispacing constraints

Figure 5: Results for problem POL (30 points)
Problem SCHUTZE1: Newton equispaced

Problem SCHUTZE1 Global Equifront

Figure 6: Results for problem SCHUTZE1 (30 points)
Problem DZT4 with \( f_1 = x^4 + 2 \)

Simple Newton continuation with 30 points

Problem DZT4 with equispaced continuation

Using \( f_1 = x^4 + 2 \)

\( x_1 \) in \([0, 1]\), which makes it essentially a bound constrained problem, which is outside the scope of this paper. Thus we changed \( f_1 \) to \( f_1(x) = x_1^2 \), making the problem convex and with a minimum at \( x = 0 \). To complete the specification of the problem:

\[
f_2(x) = g(x)[1 - \sqrt{x_1/g(x)}], \quad g(x) = 1 + 10(n-1) + \sum_{i=2}^{n} [x_i^2 - 10\cos(4\pi x_i)], \quad n = 10.
\]

In Figure 7 we see the results for simple continuation and for continuation with equispaced front constraints and in Table 2 we see that there is no significant impact by the increase in dimensionality. Again, simple continuation does not do a good job, but continuation with equispaced constraints produces an excellent representation of the Pareto front in about 120 function evaluations (for 30 points in the front). The RMS upon convergence is never greater than \(10^{-12}\) (this combines the norm of the gradient with the residual of the equispacing constraints). Although they are not directly comparable, Deb et al take 30,000 function evaluations for a fairly poor approximation to the front in the unmodified problem ZDT4, while needing 50,000 evaluations to obtain a reasonable approximation.

In terms of performance we offer some timings in Table 2. For these simple problems the times are essentially negligible. We emphasize that since they are all based on Newton’s method they use very few function evaluations as compared to evolutionary algorithms. This will be a distinct advantage when applying these methods to problems with expensive function evaluations, provided the necessary derivatives are available, hardly the case in problems where large scale modeling is required to evaluate the goals and constraints, but it
Table 2: Performance on an Intel dual Xeon 3.2 GH machine (LINUX). Time in msec.

might be useful if those are replaced by differentiable surrogates. The time for Method 2 in problem SCHUTZE1 includes the initialization time by Method 4.

6 Conclusions

We have discussed in detail the problem of finding well sampled representations of the Pareto manifold or of its image by the objectives, the Pareto front, in a unconstrained multi-objective optimization problem. Explicit algorithms for bi-objective problems were described in detail, implemented and demonstrated on problems taken from the recent literature. Extensions to more objectives and constrained problems were indicated.

We showed that by adding simple equispacing constraints it was possible to have both simple homotopic continuation of scalarized problems or a novel global "bending" method to produce perfectly evenly spaced representations of either the Pareto manifold or the Pareto front.

The algorithms, based on the solution of the KKT first order optimality conditions by Newton’s method were shown to use two orders of magnitude less objective evaluations than commonly used evolutionary algorithms. Although we agree with the fact that evolutionary algorithms are applicable to a wider class of problems, we like to emphasize that for problems that are common in Engineering and Scientific applications involving expensive simulations in order to evaluate the objectives and constraints, the proposed methods could be a useful alternative.

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References


