A demonstration of the utility of fractional experimental design for finding optimal genetic algorithm parameter settings

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This paper demonstrates that the use of sparse experimental design in the development of the structure for genetic algorithms, and hence other computer programs, is a particularly effective and efficient strategy. Despite widespread knowledge of the existence of these systematic experimental plans, they have seen limited application in the investigation of advanced computer programs. This paper attempts to address this missed opportunity and encourage others to take advantage of the power of these plans. Using data generated from a full factorial experimental design, involving 27 experimental runs that was used to assess the optimum operating settings of the parameters of a special genetic algorithm (GA), we show that similar results could have been obtained using as few as nine runs. The GA was used to find minimum cost schedules for a complex component assembly operation with many sub-processes. *Journal of the Operational Research Society* (2004) **55**, 132–138. doi:10.1057/palgrave.jors.2601703

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Introduction

Design of experiments (DOE) techniques provide a systematic, effective and efficient approach to the investigation of a phenomenon.¹ The approach is often sequential in nature, potentially increasing in complexity as the knowledge and understanding of the application and domain evolves.² The main advantages of the strategy are the savings in time and resources expended compared to other approaches and the resulting mathematical models that help users to better understand the phenomena under investigation more fully. This investigation demonstrates the application of a number of efficient experimental designs to re-address a previous application of DOE techniques within the optimization of the structure of a GA. The particular experimental designs considered within this investigation include: a two-level 'fullfactorial' design with one centre point in nine experimental trials; a Box–Behnken design with 13 trials, and, a central composite with 15 trials. The focus is on demonstrating that these designs, in particular, the nine trial designs may be used with a fitness check of the model to assess whether there is any requirement to undertake further trials. In computer simulation work of this nature, there is no penalty from adding later trials, unlike the situation in industry where 'blocking' or 'nuisance effects' are often observed. By this, we refer to changes over time in the response that is being measured. In an industrial setting, for example, we may see different results at night, the next day, or because a shift has

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changed. Material inputs may become different, environmental conditions may change, and thus adding the results of trials run at a later time may result in wrongful conclusions unless these potential effects are considered. In simulation work this is not an issue, running a program with the same parameters and the same initialization will always produce the same results. The paper suggests that a minimum number of treatments can be investigated which need only be added to later, if the initial analysis suggests that this is required, resulting in a considerable saving in time when investigating large and complex problems like this one. The paper goes on to show how, in this case, a reduced number of experimental runs would have produced a similar result to those found in the actual investigation. A brief description of the scheduling problem is given first, then we show how experimental design techniques were used to determine the 'best' structure of the GA and then the adoption of a sequential strategy is discussed. The use of this particular study as an example should, we hope, serve as a springboard for others to use these methods in other computer program optimization work. It has been overlooked by some writers that computer programs have good and bad ways of being written, and optimum programs can often achieve better results.

Scheduling problem description

Scheduling has been defined as 'the allocation of resources over time to perform a collection of tasks'.³ These types of problems are often difficult to solve because they involve complex combinatorial optimisation and can only be solved

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by non-deterministic polynomial algorithms. These problems are well suited to solving by Heuristic means. In particular, the use of GAs has been shown to speed discovery of good solutions.

The work analysed here involved using the Genetic Algorithm (GA) to produce improved solutions for a large, computationally intensive scheduling problem,^{4,5} in which several other popular optimization techniques had been shown to produce very much inferior schedules. The GA used was based on a modification of the Goldberg algorithm⁶ that includes a repair function to avoid infeasible solutions. The capital product being assembled had six levels of product structure, 46 components, 497 machining operations and 39 assembly operations, which were performed using 24 machine tools and one assembly area. The fitness function minimized the various cost penalties that were associated with either being early or late with the various components in the assembly.

Experimental design

Pongcharoen *et al*⁵ investigated the application of genetic algorithms for scheduling the production of capital goods using existing data. Experiments were designed which varied the population size, number of generations, and, mutation and crossover probabilities. It was discovered in that work that algorithms with large population sizes and that ran for many generations tended to produce the largest improvement with the best solutions having approximately 80% lower penalty costs than the original company schedules. This work was extended using designed experiments to select the most efficient GA parameters to achieve minimum total costs and spread within a specified execution time.⁴ The number of generated chromosomes was fixed at 1200 in all cases. The population/generation combination and crossover and mutation probabilities were varied according to Table 1 with these values chosen on the basis of previous investigations—Pongcharoen *et al*⁵ and Todd.⁷ Each trial run of the algorithm took 2.5 h to complete on a PC.

A full factorial experiment was used to investigate cost schedules over these three specific levels or settings of the parameters and was replicated five times using different random number seeds to facilitate the determination of real predictors-Figure 1a. A full factorial refers to the fact that every possible combination of the specific settings is used in

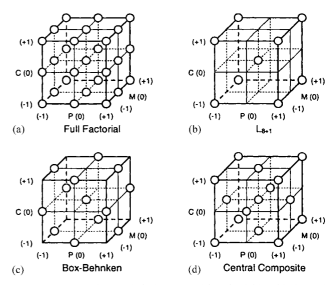


Figure 1 Experimental designs considered within this investigation.

the set of trials. Thus, for three-factors (here the GA parameters) at three levels, we get 27 trials in total. In factorial designs, it is possible to estimate N-1 terms for a plan involving N trials. A three-level factorial over three factors like this will provide us with a mathematical model or polynomial that has up to 26 terms, plus a constant that represents the effects on the response (here the cost) of the parameter settings in combination. This will include, where deemed necessary, quadratic terms as well as first-order ones, plus interaction terms that consider the changing effects on the response of one parameter as another is varied. Table 2 shows the full range of terms available from this three-level full factorial including the constant. The 'P'values shown here indicate that factors P and M are considered important if all the terms are considered together in an analysis. In the table, a '2' in the first column of terms indicates a quadratic term or quadratic component of a term.

These models can be quite comprehensive and provide a good approximation to reality over the whole range of the chosen settings, including combinations not chosen in the trials run, even in situations displaying considerable nonlinearity. The results were analysed with a multiple regression-based method known as 'best subsets'. A fuller general

Table 1	Experimental parameter settings						
Parameters with denoted coding		Parameter settings and coded values					
	(-1)	(0)	(+1)				
Population × generation combination size—P Chance of crossover— C (%)	20:60 30	40:30 60	60:20 90				
Chance of mutation— <i>M</i> (%)	2	10	18				

 Table 2
 p-values and terms for full ANOVA analysis of original full factorial

Term	p-value				
Constant	0.000				
Р	0.007				
M	0.614				
PM	0.038				
С	0.638				
PC	0.651				
P2	0.208				
M2	0.704				
C2	0.54				
CM	0.37				
PCM	0.929				
PM2	0.826				
PC2	0.324				
MP2	0.322				
MC2	0.776				
CP2	0.77				
CM2	0.574				
M2P2	0.431				
M2C2	0.393				
P2C2	0.36				
P2C2M2	0.379				
P2M2C	0.987				
P2C2M	0.515				
M2C2P	0.661				
P2MC	0.33				
M2PC	0.848				
C2MP	0.331				

description of the method is given in Draper and Smith,⁸ Montgomery⁹ but the essence is to look at all possible combinations of the potential terms within the mathematical model. Important terms are deemed to be significant if by adding them to the model there is significant improvement in the predictive capability of the mathematical model as discussed in a later section. Other methods are possible, such as ANOVA—the analysis of variance—or graphical methods that are covered in later sections. In ANOVA, the terms are given the well-known 'p'-values that display the probability of the term not being important. Traditionally, a term with a p-value of 0.05 or less is taken as important and thus retained in the model. This point is further discussed in later sections.

Parameters and associated effects found to be significant using 'best subsets' and sequential ANOVA were P, M and the interaction between P and M (PM). The results indicated that the crossover probability C is not significant within this application, across the region considered, which concords with results in Pongcharoen *et al.*^{4,5} None of the other potential model terms were found to make a significant difference to the results. This paper assesses trying small subsets of the whole factorial to determine that a smaller design with fewer runs would have been sufficient to discover the same important parameters.

Fractional experimental design

A number of different smaller experimental designs were considered within this investigation including: a two-level fractional-factorial with one centre point (denoted L_{8+1}) in nine runs as described in Grove and Davis;¹⁰ a Box–Behnken design with 13 trials (see Box and Draper¹), and, a central composite first described in Box and Wilson¹¹ with 15 trials—Figure 1. All are based on the relevant subset of the data from the full factorial using all five replicates. 'Fractional' indicates that the design is a fraction of a full factorial.

The L_{8+1} experiment uses the extreme points on the vertices of the design plus a centre point, an extra trial run with all parameters set in the middle of their range, over the five replicates giving a total of 45 trials-Figure 1b. The results from this combination were then used to produce standardized coefficients (these are weighted in relation to the standard deviation associated with their estimation) as in Grove and Davis,¹⁰ as well as *T*-values which are displayed within Figure 2. These results are interesting since they again suggest that the dominant predictor is P, however, the only other significant predictor, according to ANOVA is M, with the *PM* interaction and all other predictors being insignificant. In a screening experiment (a term used to denote that we are at the first stage of a set of trials and may add to them later) it should be borne in mind, however, that effects that appear at first sight to be too small by traditional standards (where terms with a *p*-value over 0.05 were excluded) may in fact be important, see for example Box and Liu.² In Sexton et al,¹² it is pointed out that effects with a P-value of 0.2 or less should not be excluded in these initial stages. The reason for this is that in small screening designs the statistical power of the analysis, literally our ability to detect real effects, is reduced because of the smaller number of runs conducted. Thus, a larger *p*-value, traditionally attributed to insignificant effects, has to be viewed in relation to the amount of information available. A fuller discussion of this, in relation to the structure of GAs, is given in Poncharoen et al.¹³ In the half-normal plot in Figure 2, that is basically a normal probability plot that is folded in half, the standardized coefficient¹⁰ for the curvature (denoted curve) can also be seen to be small giving no indication of curvature on main effects within the design space. Details of the procedure used to determine the 'curvature' are also given in Grove and Davis.¹⁰ This suggests that we do not need to determine the magnitude of the effect of the quadratic terms, and the current model is sufficient as a good predictor of the response. Of course, in our case we know this already from the full factorial results, but it is instructive that this smaller design gives similar results to our bigger experiment with all 27 trials. Previous investigations have, however, demonstrated that the test for curvature within a main effect may fail if the design space contains or surrounds a saddle point,¹⁴ so some care has to be taken.

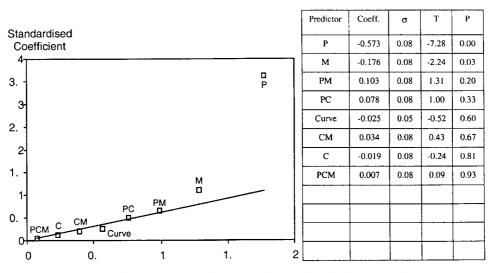


Figure 2 Half-normal plot and ANOVA for standardized L_{8+1} coefficients.

	Statistics			Predictors								
Terms	R_p^2	\bar{R}_p^2	C_p	S	Р	М	PM	С	РС	СМ	РСМ	
2	59.7	57.8	1.0	0.481	×	×						
2	56.3	54.2	4.4	0.501	×		×					
3	61.4	58.6	1.3	0.476	×	×	×					
3	60.7	57.8	2.0	0.481	×	×			×			
4	62.5	58.7	2.3	0.475	×	×	×		×			
4	61.6	57.8	3.1	0.481	×	×	×			×		
5	62.7	57.9	4.1	0.48	×	×	×		×	×		
5	62.5	57.7	4.2	0.481	×	×	×	×	×			

Table 3 Best subsets regression for L_{8+1}

The design space was tested for saddle points within the PM plane, by examination of the PM plot based on the mathematical model (see Figure 5) and was discovered to be relatively flat and free from such phenomena. Thus, we can trust the fact that the 'curve' estimate is reliable in this case.

Analysis of 'best subsets' (see Montgomery⁹ or Hines and Montgomery¹⁵ for a fuller explanation by writers from an engineering background) was also tried in order to find the best regression equation based upon a number of standard criteria as shown in Table 3. In this table, column 1 shows the number of terms in the model on that row, the specific terms are listed by \times s in the last seven columns. The criteria are:

- the coefficient of multiple determination, R_p^2 which represents the proportion of the sum of the squares deviation in the response variable *y*, about the predicted values \hat{y} , that can be attributed to the regression;
- the adjusted determination coefficient \bar{R}_p^2 , that accounts for the number of predictors used in the model and the number of treatments within the experiment;

- the square root of the mean squared error, s;
- and, Mallow's C_p statistic which is a measure of the total mean square error for the regression model, compared to the estimate of background uncertainty, but adjusted for p.¹⁶

As the R_p^2 criterion increases, so does the predictor's collective ability to predict the response, hence the largest value for R_p^2 generally represents the best model. However, the coefficient of multiple determination rises as the number of predictors grows. The adjusted coefficient of determination (\bar{R}_p^2) is independent of the number of predictors in the model and is consequently a more useful indicator than R_p^2 . Increasing values of \bar{R}_p^2 represent increasingly better models. Mallow's C_p may be used to estimate the relative amount of bias, with less biased models having lower values for C_p , which are usually close or equal to the number of predictors within the model. It can be seen from Table 3 (that gives the two best models for each number of terms in the model up to 5) for example that the model with only *PM* and *P* is not as good as the model with *P* and *M* main effects. The results

suggest that a model including just *P*, *M* and the *PM* interaction is the 'best' one. No increase in predictive power or reduction in uncertainty in the statistical model can be achieved from adding more terms. The best model with four terms has almost identical R_p^2 , \bar{R}_p^2 and *s* values and a higher C_p . We can rely on the model with three terms.

The central composite design is composed of a standard two-level factorial or fractional factorial design augmented with one or more centre points run in the middle of the design space, as in the design just discussed above, plus two extra runs per main predictor known as star points. The star points can be run in the centre of each 'face' of the cube portion of the design-Figure 1d. The idea of the star points is to enable estimates of the quadratic terms if the test for curvature suggests that this is necessary. Thus the CCD in this case is just the L_{8+1} design with the addition of star points and thus not surprisingly gave similar results to the L_{8+1} . In fact, it is clear from the results above that inclusion of the star points would not provide better models. The Box-Behnken design is another quadratic design that does not contain an embedded factorial or fractional factorial matrix—Figure 1c. In this design, the treatment combinations are at the midpoints of the edges of the design space plus one at the centre. It is mostly used in situations where the most extreme conditions of the experimental space are difficult or impossible to run. The Box-Behnken design gave similar results, as shown in Table 4. We can see that the addition of the quadratic form of parameter P does improve the model slightly. As a rule of thumb, to justify the addition of another term to a model, a fall in s should be greater than the equivalent of the inverse of 'the number of trials run minus 1 minus the number of parameters in the model' (here 74-4=70). So a fall of over 0.499/70=0.00712 is required and so the addition of P^2 does not really improve the model in this case.

Sequential experimentation

The idea of a sequential strategy is well suited to computer experimentation, there being no penalty from adding extra trial results at a later stage. In the industrial experimental

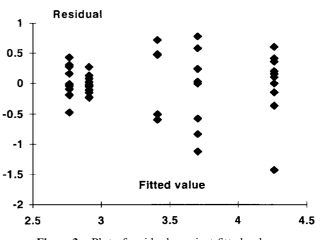


Figure 3 Plot of residuals against fitted values.

setting, we must take account of blocking or time effects and treat the results using split-plot analysis.⁹ Here there are no time effects, we can add extra results knowing that they will be identical to ones run under the same combination of factor settings at any time. Thus, the optimum strategy would be to run the minimum number of trials, observe the results, and if necessary add further trials as required. In this case, if the centre point results from the L_{8+1} had indicated the presence of curvature in a main effect, we could run the star points in order to estimate the quadratic effects that cause the curvature. Another tool used to assess regression models is the residual plot. This is simply the plot of the difference between observed values and model predictions for those observations, against various criteria. These plots would usually include; the order of experimentation, the factor settings and the fitted values of the model. In the case of computer experiments, the run plot is immaterial but if a pattern exists in the other types this may indicate the need for a missing term in the model. By the sequential use of these plots, and the model assessing criteria shown in Table 2, we can continue until a good model is selected. Figure 3 shows the fitted values against residuals for the L_{8+1} model and Figure 4 the mutation settings against the same residuals. No particular non-random patterns are visible although one possible outlier is present.

 Table 4
 Best subsets regression for Box–Behnken

Statistics					Predictors								
Р	R_p^2	$ar{R}_p^2$	C_p	S	Р	М	С	PM	PC	СМ	P^2	C^2	M^2
2	46.6	44.9	4.6	0.514	×	×							
2	44.7	42.9	6.8	0.522	×			×					
3	50.4	47.9	2.1	0.499	×	×		×					
3	48.9	46.4	3.9	0.506	×	×					×		
4	52.6	49.5	1.4	0.492	×	×		X			×		
4	51.1	47.9	3.3	0.500	×	×		×		×			
5	53.3	49.3	2.6	0.492	×	×		X		×	×		
5	52.9	48.9	3	0.494	×	×	×	×			×		

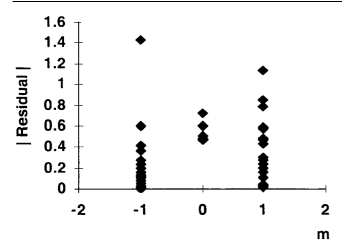
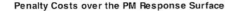


Figure 4 Plot of absolute value of residual *versus* coded probability settings for mutation.



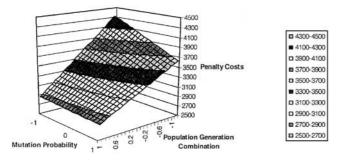


Figure 5 3D plot of penalty cost response surface.

If the residuals had indicated a clear pattern, then we could have added the star points or the edge midpoints from the Box-Behnkin design. Clearly, such a strategy will lead to savings in time and computer resource in many cases. In the present case, running just the L_{8+1} would have led to a saving of 66% in the time taken to run the original experiment. Cases with a greater number of factors can lead to many times this saving, see for example Poncharoen.¹³ A sparse design, if considered prior to the original investigation, could also have included the separate investigation of the population count and number of generations, rather than including these combined as a ratio as they were in this case. Other research suggests that mutation appears to be more useful than crossover for small population sizes;¹⁷ while crossover may be more useful than mutation for large population sizes.18

Follow up experimentation

Having discovered the best settings for the parameters, in this case setting the population generation combination at the high level (60–20) and setting the chance of mutation at the high level (18%) we can see that the experimenters had not yet found the optimum solution. Figure 5 shows a 3D response surface of the final model⁷ over the *PM* design space. The model was: *Predicted total cost* $\pounds = 3380 - 553P - 180M + 149PM$.

It can be seen that the optimum would appear to be somewhere off the plot at higher levels of the probability of mutation and higher populations. By adding a few more runs in the 'direction of steepest descent' which is easily determined,⁹ we could track smaller and smaller penalty costs until we found a design area that contained a minimum. The strategy would then be to run another small factorial experiment around the new minimum, possibly augmenting with star points if determined necessary, to find the definitive minimum. In the event, further work¹³ showed that a better GA (with differently written mutation and crossover operators) existed so further work using this particular one has ceased. It should also be noted that crossover probabilities became important in the new GA, whereas here they were not. It should be noted further that the probability range for crossover is completely different than that for Mutation. We can only say that probabilities in the range of 0.3-0.9 make little difference to results, but a choice of, say, 0.1 may well make a difference, we cannot tell from these results.

Strategy for DOE-based computer program development

- Establish the parameters or factors over which experiments will be conducted. These can be measured on categorical, ordinal or continuous scales. They should be able to be controlled (that is, can be set at particular levels) during the experiment, independently of the other factors. If this is not true, derived factors of these that can be set independently may be used, such as the ratio between two factors. Categories may be the type of coding, a program structure or a problem type. Include as many factors as possible.
- Decide on the response to be optimized. This can also be measured on categorical, ordinal or continuous scales. This could also be a derived measure such as a ratio or sum.
- Decide on the range of parameter settings. These should ideally be wide enough to cover factor levels that might show a discernable difference in the response. Typically, the highest and lowest possible, where a scale is available, are chosen.
- Decide for each factor if a quadratic term may be likely to be needed. If so this will require three levels in the design, meaning at least a centre point.
- Decide on an initial 'screening' design. This will mean choosing as few trials as possible that cover the design space over all the factors. This should be as sparse as

possible and will typically have no fewer than (2N) + 1 trials for N factors.

- Run the screening design and analyse the results to produce an initial mathematical model.
- Discard all unimportant factors and run further trials if the results are not clear or if the need for more complicated terms in the model is indicated.
- Now search for the most optimum combination of factors, possibly moving outside the original range of factor settings and running more experiments.
- Report the optimum model and use to demonstrate the best computer program.
- Use the model as a benchmark against which to compare new programs and developments in future as these occur. Add new runs as necessary to include new factors as they surface.

Conclusions

This paper demonstrates how a sequential strategy using experimental design for investigating a GA would have resulted with significantly fewer experimental treatments than that performed within the actual investigations. The L_{8+1} design demonstrated how similar conclusions would have been drawn with a more parsimonious initial design. Identical results to those obtained from the full design were also obtained using the central composite design. The overall conclusion is that a sparse experimental design would have sufficed in this case, however, additional experimental runs could have been conducted had the analysis indicated the need to do this or if the designer suspected the need to do so. Further experiments could be used to add to the experimental area once the direction of the optimum solution has been established. We have argued that computer experiments are an ideal use of DOE due to the lack of any experimental error that is usually experienced in real-life applications because of unplanned differences between experimental conditions. We have given a brief roadmap for the use of these designs and highly recommend a sequential strategy based on these designs in all such investigative work whatever the nature of the computer program being written.

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