# Scaling Effects in the CSP Phase Transition<sup>\*</sup>

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Abstract. Phase transitions in constraint satisfaction problems (CSP's) are the subject of intense study. We identify a control parameter for random binary CSP's. There is a rapid transition in the probability of a CSP having a solution at a critical value of this parameter. This parameter allows different phase transition behaviour to be compared in an uniform manner, for example CSP's generated under different regimes. We then show that within classes, the scaling of behaviour can be modelled by a technique called "finite size scaling". This applies not only to probability of solubility, as has been observed before in other NP-problems, but also to search cost. Furthermore, the technique applies with equal validity to several different methods of varying problem size. As well as contributing to the understanding of phase transitions, we contribute by allowing much finer grained comparison of algorithms, and for accurate empirical extrapolations of behaviour.

#### 1 Introduction

A phase transition in random CSP problems has recently been the subject of intensive theoretical and empirical study [30, 13, 26, 27, 21, 22, 5]. Theory predicts approximately where the phase transition can be expected, but otherwise very little information is available as to what behaviour can be expected at different problem sizes and at different points with respect to the phase transition.

Compared to SAT, the model for generation of random CSP problems is complicated, and no control parameter is used in presenting data. Yet the identification of a control parameter for random SAT problems [2, 19] is fundamental to current research on phase transitions in SAT. In this paper we introduce a control parameter for CSP. The phase transition is always expected at the same value of this parameter. Using extant data, we show that it can be much more easily understood by plotting with respect to this parameter than by raw plots as previously presented.

In the rest of the paper, we show that the control parameter can be used to direct detailed and meaningful comparisons of significantly different methods of

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generating random binary CSP's. We show that our proposed parameter gives meaningful results as we change the number of variables in our problems, change the number of values in the domain of variables, or vary both simultaneously. In each case the phase transition in probability seems to occur at similar values of the parameter with changing problem size. Furthermore, median search cost seems to peak over the same range of the parameter.

Using our results on these problems, we are able to show that the technique of "finite size scaling" can be applied to problems in CSP phase transitions. The result is an empirical prediction of how probability of solubility varies with problem size, and this prediction might be used to help derive experimental parameters for future experiments. We show this for each different method of varying problem size that we investigate. This is the first time finite size scaling has been applied to a computational phase transition on changing domain size.

The interest in how the probability of solubility varies with problem size is largely due to the correlation with peak in search cost at that phase transition. We show how remarkable a correlation this is by demonstrating that finite size scaling, applied with parameters derived only from examination of probability data, seems to apply to search cost also. The implications for the experimental analysis of algorithms are very significant.

# 2 Binary Constraint Satisfaction Problems

In the binary constraint satisfaction problem (CSP) we have a set of variables, where each variable has a domain of values, and a set of constraints acting between pairs of variables. The problem is then to assign values to variables, from their respective domains, such that the constraints are satisfied [3, 17, 28]. One way of addressing this problem is via systematic search using backtracking, the objectives being to find a solution, or determine that none exists, with minimal search effort, where effort is measured as the number of compatibility checks performed between pairs of variables. Given a CSP with n variables with uniform domain size of m, there will be  $m^n$  possible assignments of variables to values. The best known complete algorithms for CSP'sare exponential in the worst case.

Numerous studies have been performed on random CSP's, in order to measure the performance of algorithms [4, 24, 29] and to investigate the nature of problems [21, 26]. Random CSP's re typically categorised using four parameters, namely  $\langle n, m, p_1, p_2 \rangle$ , where n is the number of variables, m is the uniform domain size,  $p_1$  is the proportion of edges in the constraint graph (ie. the density of the constraint graph), and  $p_2$  is the proportion of pairs of instantiations over a constraint that are disallowed (ie. the tightness of the constraints) [21, 26, 5]<sup>3</sup> It has been observed that if n, m, and  $p_1$  are held constant, there is a small

<sup>&</sup>lt;sup>3</sup> That is, in a random CSP  $\langle n, m, p_1, p_2 \rangle$  as defined in [21, 26] there will be exactly  $p_1.n.(n-1)/2$  constraints, and each constraint will have exactly  $p_2.m^2$  conflicts. Lisp and Scheme versions of such a problem generator, and supporting search algorithms etc., are available via anonymous FTP at site ftp.cs.strath.ac.uk in directories local/pat/csp-lab/ for Lisp and local/pat/csp-lab.scm/ for Scheme.

range of values of  $p_2$  where average search effort rapidly increases to a peak and then falls away, while at the same time the proportion of soluble problems drops to zero. That is, there is a phase transition [26, 21, 5].



**Fig. 1.** Gaschnig's experiments on random 10-queens, and ours on (10, 10, 1.0)

Probably the earliest report of the complexity peak in CSP'sis by Gaschnig [7]. One of the studies in his thesis was on random 10-queens. In the n-queens problem n non-attacking queens have to be placed on an  $n \times n$  chess board, and in the random 10-queens problem a solution (or proof that none exists) has to be found for  $\langle 10, 10, 1.0, p_2 \rangle$ . Figure 1(a) shows a plot of the results from Gaschnig's experiments for random 10-queens. Constraint tightness  $p_2$  is varied in steps of 0.1 (with the exception of the point  $p_2 = 0.35$ ) and 150 problems are generated at each point. Three curves are plotted, one for chronological backtracking (BT), one for backjumping (BJ), and one for backmarking (BM) [11, 7, 6].<sup>4</sup> The y-axis is the average number of consistency checks and the x-axis is  $p_2$ . Figure 1(a) clearly shows a peak in average search effort at  $p_2 = 0.4$  for the three algorithms.<sup>5</sup> In Figure 1(b) the experiments are repeated, but  $p_2$  is varied in steps of 0.01, and this confirms that the peak in average search effort does indeed occur at  $p_2 = 0.4$ . Furthermore, 52% of the  $\langle 10, 10, 1.0, 0.4 \rangle$  problems are soluble. It appears that Gaschnig failed to notice this phenomenon.

### **3** A Control Parameter for Binary CSP's

Given the random CSP  $(n, m, p_1, p_2)$  the expected number of solutions is given by (1)

$$E(N) = m^{n} (1 - p_{2})^{\frac{p_{1} n(n-1)}{2}}$$
(1)

In [26] it is conjectured that average search effort will be greatest when an ensemble of problems have on average one solution, i.e. E(N) = 1, and this

<sup>&</sup>lt;sup>4</sup> Note that no variable or value ordering heuristics were used.

<sup>&</sup>lt;sup>5</sup> Gaschnig referred to the CSP as a SAP (satisficing assignment problem) and L as the degree of a constraint (the fraction of distinct pair tests that have the value true, ie.  $L = 1 - p_2$ ). The plot of 1(a) uses the data in Figure 4.4.3-1, page 301 of [7]. Gaschnig noted the existence of a sharp peak at  $L \approx 0.6$ , pages 179 and 180.

will correspond to the crossover point where half the problems are soluble. An equivalent theory was independently developed by Williams and Hogg [30]. For given values of n, m, and  $p_1$  the critical value of constraint tightness  $p_{2crit}$ , where average search effort will be a maximum, may be predicted via (2)

$$p_{2crit} = 1 - m^{-2/((n-1)p_1)} \tag{2}$$

For example, using (2) we can predict the critical value of constraint tightness for (10, 10, 1.0), ie. Gaschnig's random 10-queens experiments, and that is  $p_{2crit} = 0.400$ , in full agreement with his observations.



**Fig. 2.**  $(20, 10, p_1)$  (a) Median search effort against  $p_2$ , and (b) Percentage solubility against  $p_2$ 

Figure 2 shows the median search effort for the CSP's( $20, 10, p_1, p_2$ ) (ie. 20 variables, uniform domain size of 10,  $p_1$  varying from 0.1 to 1.0 in steps of 0.1,  $p_2$  varying from 0.01 to 0.99 in steps of 0.01). In Figure 2(a) 10 contours are given, the leftmost is for  $p_1 = 1.0$  and the rightmost for  $p_1 = 0.1$ . The x-axis is constraint tightness,  $p_2$ , varying in steps of 0.01, and the y-axis is the log of the median search effort. At each value of  $p_1$  and  $p_2$  one hundred problems were sampled using the algorithm forward checking with conflict-directed backjumping (FC-CBJ) [20] allied to the fail first heuristic (FF) [23, 12]. What we see is that as the density of the constraint graph increases (ie.  $p_1$  increases) the critical value of constraint tightness falls (ie.  $p_2$  falls). Figure 2(b) shows how the solubility of problems varies with  $p_1$  and  $p_2$ . Again 10 contours are given, the leftmost for  $p_1 = 1.0$  and the rightmost for  $p_1 = 0.1$ .

In some respects Figure 2 suggests that it might be difficult to compare CSP's of different size (ie. varying n and m) or structure (ie. varying  $p_1$  and  $p_2$ ) because their graphs translate along the x-axis, changing shape as they go.<sup>6</sup> What we would like to find is some parameter that characterises CSP regardless of size or structure, ie. a control parameter. Control parameters have been identified for 3-SAT, namely  $\frac{clauses}{variables}$ , and in 3-COL the average degree  $\gamma$  [2]. We

<sup>&</sup>lt;sup>6</sup> Although all the curves in Figure 2(a) have the same *signature*, at high values of  $p_1$  they are more defined.

can derive a control parameter  $\tau$  for CSP's as follows. First, we rearrange (2) to get

$$\frac{n-1}{2}p_1\log_m(\frac{1}{1-p_{2\,crit}}) = 1 \tag{3}$$

This gives us a prediction for the location of the crossover point expressed by a function of the random generation parameters taking a certain *constant* value, and that function no longer has a first order dependency on n, i.e. it does not have the exponential behaviour of (1). This immediately suggests that the LHS of (3), may be a suitable control parameter for CSP's. Accordingly, we *define* the parameter, which we call  $\tau$ , by

$$\tau =_{def} \frac{n-1}{2} p_1 \log_m(\frac{1}{1-p_2})$$

Note that we define  $\tau$  in terms of  $p_2$  instead of  $p_{2crit}$  in (3). This means that  $\tau$  is defined for *all* values of n, m,  $p_1$ , and  $p_2$ , and so can be used to compare CSP'sgenerated with different parameters. If the theory of equation (2) were exactly correct, then values of  $\tau$  less than 1 would lead to soluble problems, while values more than 1 would give insoluble problems. We will see in the rest of this paper that this is a reasonable but not completely accurate prediction.



**Fig. 3.**  $(20, 10, p_1)$  (a) Median search effort against the control parameter  $\tau$ , and (b) Percentage solubility against  $\tau$ .

Figure 3 shows the same data as in Figure 2 but with  $\tau$  on the x-axis. In Figure 3(a) the contours of median search effort peak when  $0.75 \leq \tau \leq 1.0$ , close to the expected value of the control parameter. However, for increasing  $p_1$ the phase transition occurs more sharply and at values of  $\tau$  nearer 1. That is, for denser constraint graphs, the prediction of Smith, Williams and Hogg for the location of the phase transition becomes more accurate.<sup>7</sup> Finally, it is clear that as  $p_1$  increases towards 1, so the peak mean search effort increases considerably. Only the last of these points could have been seen clearly from the graphs as presented in Figure 2, even though in this case we have presented *less* data – we

<sup>&</sup>lt;sup>7</sup> This change appears to be related to a decreasing variance in the number of solutions at the phase transition [27].

cut off values of  $p_2$  where  $\tau > 2.0$ . These points were made in [21], but this had to be done by further analysis, and could not be read off directly from a simple plot as we did by looking at Figure 3.

We hope that our graphs argue for us the case that data should be presented with respect to the proposed parameter,  $\tau$ . We show in the rest of the paper that there are further advantages in studying this parameter. Indeed we will be able to make detailed numerical predictions based on it.

#### 4 Changing Number of Variables



**Fig. 4.**  $\langle n, 3, p_1, \frac{2}{9} \rangle$  plotted using  $p_1$ 

A set of experiments was carried out to investigate the effect of increasing the number of variables, i.e. n, on the parameter  $\tau$ . In these experiments we vary the number of variables and the density of the constraint graph, while holding the domain size and tightness of constraints constant. We chose a domain size m = 3 and tightness of constraint  $p_2 = \frac{2}{9}$ , corresponding to the set of experiments reported in [5], and we will refer to them as  $\langle n, 3, p_1, \frac{2}{9} \rangle$ .<sup>8</sup> The search algorithm used for this set of experiments, and all subsequent experiments reported here, was FC-CBJ-FF, i.e. forward checking with conflict-directed backjumping using the fail-first heuristic. A report on the implementation of this procedure is given in [18]. For each n from 10 to 110 in steps of 10 we tested problems from  $p_1 = \frac{1}{n-1}$ to  $p_1 = \frac{10}{n-1}$  in steps of  $\frac{1}{5(n-1)}$ . (An exception is n = 10 where the maximum value of  $p_1$  is  $\frac{9}{n-1}$ .) These parameters are equivalent to varying the average degree of nodes in the constraint graph from 1 to 10 in steps of 0.2. For each nfrom 10 to 70 inclusive we tested 10,000 randomly generated problems at each value of  $p_1$ , while for n from 80 to 110 we tested 1,000 problems for each  $p_1$ .

<sup>&</sup>lt;sup>8</sup> This corresponds to the experiments by Frost and Dechter with N and C varying, K = 3, and  $T = \frac{2}{9}$ . See Figure 1. in [5]. A problem with N variables and C constraints is exactly equivalent to a problem generated using our model and parameters  $\langle N, 3, 2C/N(N-1), \frac{2}{9} \rangle$ .



**Fig. 5.**  $\langle n, 3, p_1, \frac{2}{9} \rangle$  plotted using control parameter  $\tau$ 

In Figure 4(a) we show how probability of solubility varies as n changes. Because the parameter that varies for each n is  $p_1$ , we plot probability of solubility (y-axis) against  $p_1$  (x-axis). With increasing n, the phase transition occurs at smaller values of  $p_1$ , i.e. the left most contour is for n = 110 and the right most contour is for n = 10. In Figure 4(b) we show how the median search cost changes with increasing n, on a logarithmic scale. As n increases the peak in median cost increases greatly and occurs at smaller values of  $p_1$  and appears to coincide approximately with the transition in probability of solubility.

Just as we saw in §3, our data is much more easily understood in terms of the parameter  $\tau$  than in terms of the raw parameter  $p_1$ . In Figure 5(a) we show the probability of solubility for each problem size tested, plotted against  $\tau$  (xaxis). Our data covers the range of  $\tau$  from 0.115 to 1.15. The phase transition in probability always starts at a value of  $\tau$  slightly larger than 0.5. Comparison with equation (3) shows that this is considerably smaller than the value of 1 predicted by the theory of Smith, Williams and Hogg. Nevertheless, using their theory we have derived a parameter at a fixed value of which the phase transition seems to cluster. It is also clear that the sharpness of the phase transition tends to increase with increasing n. We show in §5 that this increasing sharpness can be characterised *precisely*.

Figure 5(b) plots median search cost against  $\tau$  for each value of n tested. The peak in cost covers a similar range to the phase transition in solubility. However, as n increases, the peaks in median search cost become more sharply defined and appear at smaller values of  $\tau$ .

# 5 Scaling of Probability

In this section we show that the probability of solubility for a given problem class scales in an astonishingly simple way. The same technique that we use has been used in other NP complete problem classes and so seems to be of very general validity. However, since the technique is borrowed from statistical physics we briefly review some analogies between phase transitions in physical and computational problems.

Similar phase transition phenomena occur in many physical systems [31]. For example, in a ferromagnet (a permanent magnet) a phase transition occurs at a critical temperature, the Curie temperature. Above this temperature, and in the absence of an external magnetic field, the ferromagnet has no magnetization. If, however, the ferromagnet is cooled then it becomes abruptly magnetized at the Curie temperature. Several other macroscopic properties like the magnetic susceptibility (the change in magnetization induced by an external field) also undergo a phase transition at the Curie temperature.

A simple model of a ferromagnet is the Ising model. This has N atoms arranged on a regular lattice. Each atom has a magnetic spin which can be either "up" or "down". The ferromagnet can therefore be in one of  $2^{\rm N}$  possible states. Magnetism is a short-range force promoting neighbouring spins to line up together. Correlations can, however, occur between more distant spins. At a high temperature, thermal fluctuations are large and spins are independent of each other. The ferromagnet therefore has no net magnetization. As the temperature is lowered towards the Curie temperature, spins become correlated over increasingly large distances. At the Curie temperature, spins are totally correlated – changing the spin of a single atom changes all other spins.

Several analogies can be made with binary CSP's. A CSP has n variables taking one of m values, so there are  $m^n$  possible variable-value pairs. Although interactions between variables are restricted to binary constraints, correlations can occur between the values of variables not directly connected via a binary constraint. Our control parameter, which is related to the expected number of solutions to the CSP, serves as a proxy for the temperature. If this parameter is small then, as there are many models, variables can take values largely independently of each other. As this parameter is increased, the values of variables become increasingly correlated. If there is only one expected model at the phase transition, the values of variables are totally correlated with each other.

Statistical mechanics describes the behaviour of a ferromagnet in the thermodynamic limit when the volume and number of atoms goes to infinity. For finite systems, a heuristical technique called "finite-size" scaling have been developed to model phase transition phenomena [1]. Finite-size scaling also appears to be useful for modelling the behaviour of the phase transition in a variety of combinatorial problems including propositional satisfiability [15, 16, 8, 9], and the traveling salesman problem [10]. Around the phase transition, finite-size scaling predicts that problems of all sizes are indistinguishable except for a change of scale. This would suggest that,

$$Prob(solution) = f(\frac{\tau - \tau_c}{\tau_c} \,.\, N^{1/\nu}) \tag{4}$$

where f is some fundamental function,  $\tau$  is the control parameter,  $\tau_c$  is the critical value of this parameter at the phase transition, and  $N^{1/\nu}$  provides the change of scale.  $\frac{\tau - \tau_c}{\tau_c}$  plays an analogous rôle to the reduced temperature,  $\frac{T - T_c}{T_c}$  in physical systems.

If the prediction of (4) holds then there must be a "fixed point", a single value,  $\tau_c$  of the control parameter at which all different problem sizes give the same percentage solubility, f(0). This may appear not to be the case in Figure 5, except of course at 0 and 100% solubility. However, examining our data more closely in the region of high percentage solubility and interpolating between points on the plot where necessary, we did observe a fixed point.<sup>9</sup> We found very similar behaviour at  $\tau = 0.625$ , where all n gave probabilities in the range (0.974,0.982) except for n = 10 which gave 0.991. Taking sample sizes into account, all probabilities were within two standard deviations of an estimate for probability of solubility of 0.976 except for n = 10 and n = 20. We take 0.625 for the fixed point and thus for the critical value  $\tau_c$ . It is interesting to note that this is considerably smaller than the value 1 predicted by (3). This is however consistent with observations in [21, 22, 27] that the prediction of theory seems to be less accurate in the case of sparse constraint graphs, such as these graphs are for n > 20 at the critical value.



**Fig. 6.**  $\langle n, 3, p_1, \frac{2}{9} \rangle$  Plotted with rescaled parameter  $\tau_{\rm N}$ 

Having chosen  $\tau_c$ , if (4) holds then there will be a single value  $\nu$  to provide a fit to (4). Another way of seeing this is to rescale our data so that instead of plotting the control parameter  $\tau$ , we define a rescaled parameter which depends on the control parameter and the problem size, N. We call this  $\tau_N$  and in line with (4), using n for the problem size, define it by

$$\tau_{\rm N} =_{def} \frac{\tau - \tau_c}{\tau_c} \,.\, n^{1/\nu} \tag{5}$$

If the conjecture of (4) holds for the correct value of the exponent  $\nu$ , we expect to see the probability curves for each *n* very closely aligned if we plot them against  $\tau_{\rm N}$ . If so, then the resulting curve gives us an empirical prediction of the function *f*. Having chosen  $\tau_c$ , one can estimate  $\nu$  empirically by assuming that

<sup>&</sup>lt;sup>9</sup> Although the probability plots are clearly curved, locally straight line interpolation seems to be acceptable.

(5) holds. Then for a given probability of solubility, we can observe the values of  $\tau$  that give that probability for different values of n. Say that for  $n_1$  and  $n_2$  we observe the same probability at  $\tau_1$  and  $\tau_2$ . Then from (5) we expect that

$$\frac{\tau_1 - \tau_c}{\tau_c} \cdot n_1^{1/\nu} = \frac{\tau_2 - \tau_c}{\tau_c} \cdot n_2^{1/\nu}$$

Rearrangement gives us

$$\nu = \frac{\log(n_2/n_1)}{\log((\tau_1 - \tau_c)/(\tau_2 - \tau_c))}$$
(6)

We first estimated  $\nu$  using this formula and the 50% solubility points, again using linear interpolation where necessary. The choice of 50% is because it is significantly different from the probability of 0.976 at the fixed point, giving sufficient range for the scaling to take effect. Using (6) for each of the 55 pairs of  $10 \leq n_1 < n_2 \leq 110$ , gave a median estimate for  $\nu$  of 2.32 with a lower quartile estimate of 2.16 and an upper quartile of 2.51. Rescaling based on 25% solubility gives a very similar result, with a median estimate of 2.33. We thus choose  $\nu = 2.3$ . The fact that this choice gives a good fit to (4) is confirmed dramatically in Figure 6(a). (The vertical line represents  $\tau_{\rm N} = 0$ .) Under this scaling all probability curves are almost identical, except the curve for n = 10which rests slightly above the rest. This suggests that probability of solubility in this model can be described by finite size scaling with parameters  $\tau_c = 0.625$ ,  $\nu = 2.3$ , and f as seen in Figure 6(a).

The implications of this result are significant. First, in this particular model it should help design future experiments. For example, should we wish a probability of solubility of 0.5, then we can interpolate the empirically predicted value of the rescaled parameter, which in this case occurs at  $\tau_N \approx 1.45$ , this being the median value interpolated from the 11 values of n. We can rearrange (5) to find what value of  $\tau$  gives a given value of the  $\tau_N$  parameter for a given n. This is given by

$$\tau = \tau_c \, . \, (1 + \frac{\tau_{\mathrm{N}}}{n^{1/\nu}})$$

This suggests that in an  $\langle n, 3, p_1, \frac{2}{9} \rangle$  problem, 50% solubility occurs when  $\tau \approx 0.625 \cdot (1 + 1.45/n^{\frac{1}{2\cdot 3}})$ . We can unpack the definition of the control parameter to give the raw parameter  $p_1$ . In this problem class  $p_1 = 2\tau/(n-1)\log_3(\frac{9}{7})$ . We thus expect to see 50% solubility at

$$p_1 \approx (5.46 + 7.92/n^{\frac{1}{2\cdot 3}})/(n-1)$$

Fortunately we are able to test this prediction with published data, as Frost and Dechter [5] report the number of constraints observed at 50% solubility for this model. The number of constraints C is  $p_1n(n-1)/2$ , so we predict that for 50% solubility,

$$C \approx \frac{n}{2} (5.46 + 7.92/n^{\frac{1}{2\cdot 3}}) \tag{7}$$

At n = 275, the largest value of n reported in [5], 50% solubility occurs at 845 constraints. Equation (7) predicts 846 constraints. The largest n used to make the extrapolation was 110 variables. For smaller n, our prediction is not quite as accurate, but it is never more than 9 constraints out, which occurs at n = 150 with a prediction of 477 constraints compared to an observation of 468. Unlike data reported by [5], our data can also be used to interpolate for any other value of percentage solubility, and to extrapolate to any problem size.

More significant still is the likelihood that we will see similar kinds of finite scaling in other randomly generated CSP's. This is likely because once similar kinds of scaling were observed in SAT problems [15] they were observed in many different classes of SAT problems [16, 8]. We expect that similar kinds of predictions made from examining only small problems should be available for large problems in many different classes of CSP's.

## 6 Finite Size Scaling of Search Cost

The main feature of CSP problems that interests us is how hard it is to solve these problems. It is natural therefore to ask if changes in behaviour of search cost can be similarly corrected using finite size scaling? The remarkable answer is that this seems to be achievable using the *identical* rescaled parameter  $\tau_{\rm N}$ .



**Fig. 7.**  $\langle n, 3, p_1, \frac{2}{9} \rangle$  plotted using rescaled parameter  $\tau_{\rm N}$ 

In Figure 6(b) we show what happens if we plot the same data previously plotted in Figure 5(b) against  $\tau_N$ . Instead of the peaks in search cost occurring at different values of the parameter as seen previously, the peak appears to be at very similar values for each different n. This strongly suggests that the same finite size scaling that is effective for probability of solubility also models accurately the behaviour of search cost. Selman and Kirkpatrick have also shown that finite size scaling can be applied to search cost, in satisfiability problems [25]. It seems likely that it can be applied more generally.

It seems that not only median, but other measures of search cost scale in exactly the same way. Figure 7(a) shows how the 99 percentile behaves against

 $\tau_{\rm N}$ . That is, the graph plots at each point the cost that was exceeded by only 1% of problems. Just as with median behaviour, these contours line up very closely. Of course the 99 percentile is considerably worse than median behaviour, but we also note that the peaks in these curves occur at smaller values, peaking at  $\tau_{\rm N} \approx 0.8$  compared with  $\tau_{\rm N} \approx 1.6$  for median. Figure 7(b) shows behaviour of the 10 percentile, i.e. the cost exceeded by all but 10% of problems. Yet again the contours line up closely. This time the peaks are at a larger value,  $\tau_{\rm N} \approx 2$ .

It is particularly significant that we were able to use exactly the rescaled parameter  $\tau_{\rm N}$  with the *same* parameters  $\tau_c$  and  $\nu$  as used in §5. The values  $\tau_c = 0.625$  and  $\nu = 2.3$  were chosen to model scaling of probability of solubility, and this is an entirely algorithm independent feature of a problem. Yet the same parameters also accurately describe the scaling of search cost in a particular algorithm, FC-CBJ-FF. This would suggest that the finite size scaling of search cost behaviour that we have observed may be algorithm independent. While the details of contours seen with different algorithms will vary, the scaling parameters may be identical in each case. Of course at this stage this is only speculation since we have only observed scaling with a single algorithm, but the implications for understanding the scaling of search cost are enormous.

# 7 Changing Domain Size

To test our conjecture that very similar kinds of scaling would be seen with different random CSP classes, we tested a completely different model by generating problems with parameters  $\langle 10, m, 1.0, p_2 \rangle$ . Notice that in §4 we fixed m and  $p_2$ while varying n and  $p_1$ : we now fix n and  $p_1$  while varying m and  $p_2$ . Since we have fixed n = 10 and  $p_1 = 1$ , all constraint graphs we consider are simply 10-cliques, while before we typically looked at sparse constraint graphs.

From this problem class, we tested problems for m = 5, 10, 15, 20, 30, 40, and 50. Except for m = 5, where  $p_2$  can only vary in steps of  $1/m^2 = 0.04$  we varied  $p_2$  in steps of at most 0.01, covering at least a region of  $\tau$  from 0.5 to 2. We tested 1000 problems at each value of  $p_2$ . Figure 8 shows how probability of solubility and median search effort varies with m. We give one contour in each figure for each m, plotted against  $p_2$ . As m increases, the phase transition occurs at larger values of  $p_2$ , and the peak in problem difficulty grows.

In §3 we proposed a control parameter  $\tau$  for binary CSP's. If it is to be useful, it should aid comparison of our data for this problem class both with changing m and with our earlier data for  $\langle n, 3, p_1, \frac{2}{9} \rangle$ . This is confirmed by Figure 9 which shows our data replotted against the parameter  $\tau$ . It can be seen that the probability phase transition and worst case median behaviour always occurs at similar values of  $\tau$ . We observed a fixed point in probability of solubility at  $\tau_c = 1.02$  where solubility was always  $0.30 \pm 0.01$ . This is much closer to the expected critical value of  $\tau = 1$  than we saw in the previous problem class. Both the location of the fixed point  $\tau_c$  and the probability of solubility at that point are significantly different from the values  $\tau_c \approx 0.625$  and 0.976 solubility that we saw in  $\langle n, 3, p_1, \frac{2}{9} \rangle$ . However, just as in that class, both the probability transition and the peaks in median behaviour become sharper with increasing m.



Fig.8.  $\langle 10, m, 1.0, p_2 \rangle$  plotted using  $p_2$ 



Fig. 9.  $\langle 10, m, 1.0, p_2 \rangle$  plotted using control parameter  $\tau$ 



Fig. 10.  $\langle 10, m, 1.0, p_2 \rangle$  plotted using rescaled parameter  $\tau_{\rm N}$ 

Finite size scaling may seem to be an inappropriate techique, because we are no longer changing the number of variables, n. However, we are certainly varying the problem size N by changing the number of values each variable takes, m. It is important to take account of m when considering the problem size. For example, to specify a *solution* to a binary CSP requires  $n \log_2(m)$  bits, as the value each variable takes can be specified in  $\log_2(m)$  bits. So, properly, this is the measure we should have used in §5 when considering finite size scaling. However, m was constant at 3 and so does not affect the parameter  $\nu$  that we derived there. Here, we are varying m, so we redefine the rescaled control parameter as

$$\tau_{\rm N} =_{def} \frac{\tau - \tau_c}{\tau_c} \cdot (n \log_2(m))^{1/\nu} \tag{8}$$

Having done this we can proceed as before, equation (6) becoming

$$\nu = \frac{\log(n_2 \log_2(m_2)/n_1 \log_2(m_1))}{\log((\tau_1 - \tau_c)/(\tau_2 - \tau_c))}$$
(9)

We estimated  $\nu$  from equation (9) using the 50% probability point. We simply chose this as it is significantly different from the probability of 0.3 at the fixed point. The median estimate for  $\nu$  was 0.63 with lower and upper quartiles of 0.55 and 0.68 respectively. These estimates of  $\tau_c \approx 1.02$ ,  $\nu \approx 0.63$  give an very good fit to a prediction of finite size scaling. This is seen in Figure 10 (a) which shows our data for probability of solubility plotted against the rescaled parameter  $\tau_N$ .

Exactly as we saw in §6, we can use the identical parameters  $\tau_c$  and  $\nu$  to rescale contours of median cost. This is seen in Figure 10(b). As before the contours line up very closely, suggesting that finite size scaling can be applied to search cost in this problem class.

### 8 Changing Number of Variables and Domain Size

We have established that  $n \log_2(m)$  provides a good measure of problem size when varying m. Finally, we ask if it also provides a good measure of problem size when varying n and m together? To test this, we return to our starting point in this paper, namely Gaschnig's random n-queens model. In the terms of §2 these are  $\langle n, n, 1.0, p_2 \rangle$ . Thus we vary both the number of variables and domain size, in this case keeping them identical. In our experiments we tested n = 6, 8, 10, 12, 14, and 15. We varied  $p_2$  in steps depending on n, and tested 1000 problems at each value of  $p_2$ . Figure 11 shows probability of solubility and median search cost against  $p_2$ , one contour being given for each n. As n increases the phase transition occurs at smaller values of  $p_2$ , and search cost increases.

Once again, the use of our control parameter enables us to compare our data as n increases, and to contrast our data for this problem class with data from previous problem classes. Figure 12 shows the same data replotted against the parameter  $\tau$ . As in previous cases, we see the probability phase transition occurring over a similar range of  $\tau$ , as do the peaks in median search cost. The curves become sharper with increasing n.



Fig. 11.  $\langle n, n, 1.0, p_2 \rangle$  plotted using  $p_2$ 



Fig. 12.  $\langle n, n, 1.0, p_2 \rangle$  plotted using control parameter  $\tau$ 



Fig. 13.  $\langle n, n, 1.0, p_2 \rangle$  plotted using rescaled parameter  $\tau_N$ 

The fixed point in probability can be seen particularly clearly in Figure 12(a) at  $\tau \approx 1$ . Examining this data more closely we observed a fixed point in probability of solubility at  $\tau_c = 0.99$  where it was always  $0.51 \pm 0.03$ . Note that the standard deviation at probability 0.5 in a sample of 1000 is 0.015, so all results were within 2 standard deviations of 0.51. As in §7, this is very close to the predicted critical value of  $\tau = 1$ , and again we note that we are looking at constraint graphs which are cliques. The critical value  $\tau_c$  and the fixed point in probability of solubility are different to the previous two cases.

As in §7 we define the rescaled parameter  $\tau_{\rm N}$  by equation (8), using (9) to estimate  $\nu$ . In this case we could not estimate  $\nu$  using the 50% solubility point as it is too close to the fixed point probability of 0.51. Using 90% probability for estimation we obtained a median estimate of 1.02 with upper and lower quartiles of 0.78 and 1.09, while using 10% probability for estimation these values were 1.09, 1.03 and 1.45 respectively. Estimates of  $\tau_c \approx 0.99$ ,  $\nu \approx 1.0$  give an extremely good fit to a prediction of finite size scaling. This is seen in Figure 13 which shows our data for probability of solubility and median search cost plotted against the rescaled parameter  $\tau_{\rm N}$ . Again we point out the remarkable fact that finite size scaling can be applied to search cost using parameters derived solely from examination of probability data.

One of the standard beenchmark for CSP algorithms has been the *n*-queens, classified by Smith and Dyer as the problem class  $\langle n, n, 1.0, (7n-2)/3n^2 \rangle$ . Using the rescaled control parameter above  $\tau_N$  for the *n*-queens problem, we see that as *n* increases  $\tau_N$  decreases; for 10-queens  $\tau = 0.5$  and  $\tau_N = -32.3$ , for 100-queens  $\tau = 0.25$  and  $\tau_N = -989$ , and for 1000-queens  $\tau = 0.17$  and  $\tau_N = -16, 500$ . Therefore, as *n* increases the *n*-queens problem should become an easier instance of the class of problems  $\langle n, n, 1.0 \rangle$ . This is in full agreement with [27].

It is quite remarkable that the same kind of finite size scaling should be so accurate for three entirely different methods of varying problem size considered in this paper. We have varied n only in  $\langle n, 3, p_1, \frac{2}{9} \rangle$ , we have varied m only  $\langle 10, m, 1.0, p_2 \rangle$ , and we have varied n and m together in  $\langle n, n, 1.0, p_2 \rangle$ . In each case, the same equation (4) has been shown to be directly applicable, with only the parameters  $\tau_c$  and  $\nu$  and the function f varying between problem classes.

## 9 Conclusions

When presenting the results of experiments on random CSP's generated from the model  $\langle n, m, p_1, p_2 \rangle$  graphs have typically been plotted with either  $p_1$  or  $p_2$ on one axis. This tends to give a distorted view of the data, as contours rarely line up. We have proposed a control parameter  $\tau$  for randomly generated CSP's, where  $\tau$  characterises CSP's regardless of size. The parameter  $\tau$  is derived from a theory that predicts that on average the hardest problems will occur when the expected number of solutions E(N) = 1.

Analysing the empirical data for experiments with number of variables n varying, we observed a single value of  $\tau$  where problems of different sizes have the same percentage solubility, i.e. a fixed point  $\tau_c$ . Finite-sized scaling was then

applied to give a rescaled parameter  $\tau_{\rm N}$ . Replotting the data with respect to  $\tau_{\rm N}$  brought the picture into sharp focus; the solubility contours lie one on top of the other, and the peaks in median search effort coincide. Furthermore, we were able to use the rescaled parameter to estimate the critical number of constraints at the crossover point, i.e. 50% solubility, for larger values of n, and these were in close agreements with results reported elsewhere; i.e. we have given some evidence of the predictive power of  $\tau_{\rm N}$ . The same rescaling technique was then applied to data from experiments with domain size m varying, and experiments with domain size m and number of variables n varying together. In both cases problem size was taken as  $N = n \log_2(m)$ , and in both cases the data was again brought into sharp focus. This suggests that the technique may be quite general.

One of the surprises of this investigation is that a finite scaling of the control parameter based on the solubility of problems has carried over to a scaling of search cost. The rescaled parameter  $\tau_N$  models the solubility of the problem (a problem-dependent property) and the behaviour of search cost (something that we might expect to be an algorithm-dependent property). The other surprise has been that finite size scaling has been so accurate for three very different classes of problems (ie. *n* varying, *m* varying, *n* and *m* varying together).

Obviously this work represents a starting point. In the future, we would like to know the detailed scaling parameters as problems are varied in more ways than we could consider in this paper. It would be very valuable if we could *size* problems with respect to graph density  $p_1$  in order to rescale the data in §3. Finally, we note that the techniques applied in this paper effectively repair a theory which we showed to be inaccurate to a slight degree. However this repair is empirical. If our results could be used to help develop a more refined and accurate theory, it would be a pleasing validation of the empirical science of algorithms, as called for by Hooker [14].

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