

# Chapter 4

## Neighborhood Combination for Unconstrained Binary Quadratic Problems

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**Abstract** We present an experimental analysis of neighborhood combinations for local search based metaheuristics, using the Unconstrained Binary Quadratic Programming (UBQP) problem as a case study. The goal of the analysis is to help understand why, when, and how some neighborhoods can be favorably combined to increase their search power. Our study investigates combined neighborhoods with two types of moves for the UBQP problem within a tabu search algorithm to determine which strategies for combining neighborhoods prove most valuable.

### 4.1 Introduction

Neighborhood search or local search is known to be a highly effective metaheuristic framework for solving a large number of constraint satisfaction and optimization problems. By defining a neighborhood and starting from an initial solution, local search progressively explores the neighborhood of the present solution for improvement. In this way, the current solution is iteratively replaced by one of its neighbors (often improving) until a specific stopping criterion is satisfied.

One of the most important features of local search is the definition of its neighborhood. In general, good neighborhoods offer a high search capability and consequently lead to good results largely independent of the initial solution while the

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search performance induced by weak neighborhoods is often highly correlated to the initial solution [37]. Generally, a local optimum for one neighborhood is not necessarily a local optimum for another. Therefore, it is possible and interesting to create more powerful combined neighborhoods.

Using the Unconstrained Binary Quadratic Programming (UBQP) problem as a case study, we present in this work several combinations of neighborhoods, using one-flip and two-flip moves. The two-flip move proposed in this paper is new for the UBQP problem. To evaluate their performance, we carried out extensive experiments with a tabu search algorithm run on a large set of benchmark instances. Computational results show that certain combinations are superior to others.

The remaining part of this paper is organized as follows. Section 4.2 gives the description of the UBQP problem together with its recent advances. In Section 4.3, the one-flip and two-flip moves and their fast evaluation techniques are fully described. Section 4.4 is dedicated to several neighborhood combinations and our tabu search algorithm. In Section 4.5, we present our computational comparison on these neighborhoods and their combinations, and draw inferences from these findings about the factors that cause certain neighborhood combinations to be effective or ineffective. Finally in Section 4.6, we provide some conclusions and discuss some important issues related to this work.

## 4.2 Unconstrained Binary Quadratic Programming

The unconstrained binary quadratic programming problem may be written as:

$$\begin{aligned} \text{UBQP: Maximize } x_o &= xQx' \\ x &\text{ binary} \end{aligned}$$

where  $Q$  is an  $n \times n$  matrix of constants and  $x$  is an  $n$ -vector of binary (zero-one) variables.

In recent decades, the UBQP formulation has attracted wide attention for its ability to represent a wide range of important problems, including those from social psychology [20], financial analysis [27, 31], computer aided design [26], traffic management [11, 41], machine scheduling [1], cellular radio channel allocation [9], and molecular conformation [40]. Moreover, the potential application of UBQP is much greater than might be imagined, due to the possibilities of imposing quadratic infeasibility constraints into the objective function in an explicit manner. For instance, many combinatorial optimization problems pertaining to graphs such as determining maximum cliques, maximum cuts, maximum vertex packing, minimum coverings, maximum independent sets, maximum independent weighted sets are known to be capable of being formulated by the UBQP problem as documented in [38, 39]. A review of additional applications and formulations can be found in [2, 24, 25, 28].

For the UBQP problem, many exact algorithms have been proposed. The most successful approaches include those of [5, 21, 38]. However, due to its computational complexity, exact algorithms can only solve instances of small size (with 100 variables). Therefore, a large number of heuristic and metaheuristic solution proce-

dures have been reported in the literature to handle large instances. Some representative examples include local search based approaches such as simulated annealing [3, 7, 22] and tabu search [7, 17, 18, 35, 36], population-based approaches such as evolutionary algorithms [8, 23, 29, 32], scatter search [4] and memetic algorithms [33].

### 4.3 Neighborhood Moves and Fast Evaluation

In a local search procedure, applying a move  $mv$  to a candidate solution  $x$  leads to a new solution denoted by  $x \oplus mv$ . Let  $M(x)$  be the set of all possible moves which can be applied to  $x$ , then the neighborhood  $NB$  of  $x$  is defined by:  $NB(x) = \{x \oplus mv | mv \in M(x)\}$ . For the UBQP problem, we use two distinct moves denoted by one-flip and two-flip moves. In the following, we denote the neighborhoods with one-flip and two-flip moves  $N_1$  and  $N_2$ , respectively.

#### 4.3.1 One-flip Move

The one-flip move defining neighborhood  $N_1$  complements (flips) a chosen binary variable  $x_i$  by subtracting its current value from 1, i.e., the value of variable  $x_i$  becomes  $1 - x_i$  after a one-flip move. One-flip is widely used in local search algorithms for binary problems such as UBQP, multi-dimensional knapsack and satisfiability problems.

Let  $N = \{1, \dots, n\}$  denote the index set for components of the  $x$  vector. We preprocess the matrix  $Q$  to put it in lower triangular form by redefining (if necessary)  $q_{ij} = q_{ij} + q_{ji}$  for  $i > j$ , which is implicitly accompanied by setting  $q_{ji} = 0$  (though these 0 entries above the main diagonal are not stored or accessed). Let  $\Delta_i$  be the move value of flipping the variable  $x_i$ , and let  $q_{(i,j)}$  be a shorthand for denoting  $q_{ij}$  if  $i > j$  and  $q_{ji}$  if  $j > i$ . Then each move value can be calculated in linear time using the formula:

$$\Delta_i = (1 - 2x_i)(q_{ii} + \sum_{j \in N, j \neq i, x_j = 1} q_{(i,j)}) \quad (4.1)$$

For large problem instances, it is imperative to be able to rapidly determine the effect of a move on the objective function  $x_o$ . For this purpose, we employ a fast incremental evaluation technique first introduced by [17] and enhanced by [13] to exploit an improved representation and to take advantage of sparse data - a characteristic of many real world problems. The procedure maintains a data structure that stores the move value (change in  $x_o$ ) for each possible move, and employs a streamlined calculation for updating this data structure after each iteration.

Moreover, it is not necessary to recalculate all the move values after a move. Instead, one just needs to update a subset of move values affected by the move.

More precisely, it is possible to update the move values upon flipping a variable  $x_i$  by performing the following abbreviated calculation:

1.  $\Delta_i = -\Delta_i$
2. For each  $j \in N - \{i\}$ ,  $\Delta_j = \Delta_j + \sigma_{ij} q(i,j)$   
where  $\sigma_{ij} = 1$  if  $x_j = x_i$ ,  $\sigma_{ij} = -1$  otherwise.

We employ the convention that  $x_i$  represents  $x_i$ 's value before being flipped.

### 4.3.2 Two-flip Move

In the case of a two-flip neighborhood  $N_2$ , we are interested in the change in  $x_o$  that results by flipping two variables,  $x_i$  and  $x_j$ , and will refer to this change by  $\delta_{ij}$ . It is convenient to think of the two-flip process as a combination of two single one-flip moves, and we can derive  $\delta_{ij}$  using the one-flip move values  $\Delta_i$  and  $\Delta_j$  as follows (supposing  $i > j$ ):

$$\delta_{ij} = \Delta_i + \Delta_j + \lambda_{ij} q(i,j) \quad (4.2)$$

where  $\lambda_{ij} = 1$  if  $x_i = x_j$  and  $\lambda_{ij} = -1$  otherwise.

It is easy to observe that the size of neighborhood  $N_2$  is bounded by  $O(n^2)$ . After a two-flip move is performed (suppose variables  $x_i$  and  $x_j$  are flipped), we need only update the one-flip delta array  $\Delta$  that is affected by this move. Specifically, the following abbreviated calculation can be performed:

1.  $\Delta_i = -(\Delta_i + \sigma_{ij} q(i,j))$
2.  $\Delta_j = -(\Delta_j + \sigma_{ij} q(i,j))$
3. For each  $k \in N - \{i, j\}$ ,  $\Delta_k = \Delta_k + \sigma_{ik} q(i,k) + \sigma_{jk} q(j,k)$   
where  $\sigma_{uv} = 1$  if  $x_u = x_v$  ( $u, v = \{i, j, k\}$ ),  $\sigma_{uv} = -1$  otherwise.

Here  $x_i$  and  $x_j$  represent  $x_i$  and  $x_j$ 's values before being flipped.

One finds that the complexity of this updating rule is  $O(n)$ , i.e., at most  $n$   $\Delta$  values are recalculated each time. Accompanying this updating rule, it is possible to introduce additional data structures to speed up the process of identifying the best two-flip move for the next iteration. Interested readers are referred to [14] for more details.

In spite of the linear time complexity of the updating rule after a move is performed, it is still too time-consuming to examine all the two-flip moves using formula (4.2) since the two-flip neighborhood  $N_2$  has  $n(n-1)/2$  neighbors at each iteration. To overcome this obstacle, we employ a *candidate list strategy* to reduce the number of candidates in the neighborhood by examining only a small subset of all the possible two-flip moves. Specifically, at each iteration, we sort all the one-flip  $\Delta$  values in a decreasing order. Then, the two-flip move that flips  $x_i$  and  $x_j$  will be considered only if the values of both  $\Delta_i$  and  $\Delta_j$  ranks the first  $\beta$  best. In this paper, we empirically set  $\beta = 3\sqrt{n}$  which gives satisfying results without sacrificing solution quality. The greater the value of  $\beta$ , the greater will be the number of two-flip

neighborhood moves examined and the amount of CPU time required. Notice that if this candidate list strategy is disabled in  $N_2$ , the computation will be greater than that required by the one-flip neighborhood  $N_1$ .

## 4.4 Neighborhood Combinations and Algorithm

### 4.4.1 Neighborhood Combinations

In order to increase the search capability of single neighborhoods, it has become a popular practice to combine two or more different neighborhoods. The advantage of such an approach was demonstrated using a tabu search strategic oscillation design in [16], and additional variants of strategic oscillation for transitioning among alternative neighborhoods are discussed in [12]. More recently, the metaheuristic approach called Variable Neighborhood Search in [34] has effectively used a transition scheme that always returns to the simplest neighborhood when improvement occurs, while the transition scheme that cycles through higher levels before returning to the simplest (also studied in [16]) was examined in [10] and elaborated more fully in the metaheuristic context in [19].

Several ways for combining different neighborhoods arises. In this paper, we focus on two of them: Neighborhood union and token-ring search [10, 30].

We define two forms of neighborhood union: *Strong neighborhood union* and *selective neighborhood union*. For strong neighborhood union, denoted by  $N_1 \sqcup N_2$ , the algorithm picks each move (according to the algorithm's selection criteria) from all the  $N_1$  and  $N_2$  moves. For selective neighborhood union, denoted by  $N_1 \cup N_2$ , the search algorithm selects one of the two neighborhoods to be used at each iteration, choosing the neighborhood  $N_1$  with a predefined probability  $p$  and choosing  $N_2$  with probability  $1 - p$ . An algorithm using only  $N_1$  or  $N_2$  is, of course, a special case of an algorithm using  $N_1 \cup N_2$  where  $p$  is set to be 1 and 0, respectively.

In token-ring search, the neighborhoods are alternated, applying the currently selected neighborhood without interruption, starting from the local optimum of the previous neighborhood, until no improvement is possible. More precisely, the search procedure uses one neighborhood until a *best* local optimum is determined, subject to time or iteration limits imposed on the search (for metaheuristic searches, this may not be the first local optimum encountered). The *best* local optimum here denotes the best solution found so far by the current search. Then the method switches to the other neighborhood, starting from this local optimum, and continues the search in the same fashion. The search comes back to the first neighborhood at the end of the second neighborhood exploration, repeating this process until no improvement is possible. The token-ring search of two neighborhoods can be denoted as  $N_1 \rightarrow N_2$  (starting from  $N_1$ ) or  $N_2 \rightarrow N_1$  (starting from  $N_2$ ). More details are given in [30].

#### 4.4.2 Tabu Search Algorithm

For the purpose of studying the different neighborhoods and their combinations, we implement a simple Tabu Search (TS) algorithm [15]. TS typically incorporates a tabu list as a “recency-based” memory structure to assure that solutions visited within a certain span of iterations, called the tabu tenure, will not be revisited. The approach is designed to introduce vigor into the search by also forbidding moves leading to related solutions that share certain attributes (values of variables) in common with the visited solutions. In the present implementation, each time a variable  $x_i$  is flipped, this variable enters into the tabu list (an  $n$ -vector *TabuTenure*) and cannot be flipped for the next *TabuTenure*( $i$ ) iterations (*TabuTenure*( $i$ ) is the “tabu tenure”). For the current study, we elected to set

$$\text{TabuTenure}(i) = C + \text{rand}(10), \quad (4.3)$$

where  $C$  is a given constant and  $\text{rand}(10)$  takes a random value between 1 and 10.

For the one-flip neighborhood, our TS algorithm then restricts consideration to variables not forbidden by the tabu list, and selects a variable to flip that produces the largest  $\Delta_i$  value (thus improving  $x_o$  if this value is positive). In the case that two or more moves have the same best move value, a random best move is selected. For the two-flip neighborhood, a move is declared tabu if and only if both two flipping variables are in tabu status.

However, some of those neighborhood solutions forbidden by the tabu list might be of excellent quality and might not have been visited. To mitigate this problem, a simple aspiration criterion is applied that permits a move to be selected in spite of being tabu if it leads to a solution better than the current best solution.

In the case that the TS procedure is applied to a token ring search (denoted  $N_1 \rightarrow N_2$  for our two neighborhoods case), we start the TS procedure with neighborhood  $N_1$ . Since we need to search the two neighborhoods alternately, the application of TS to a single neighborhood stops when the best solution cannot be improved within a given number  $\theta$  of moves and we call this number the *improvement cutoff* of TS, which we empirically set to be a relatively small value (50,000 for all the tested instances).

### 4.5 Experimental Results

In this section, we show computational results for our simple TS algorithm using the following neighborhoods and neighborhood combinations:  $N_1$  (one-flip),  $N_2$  (two-flip),  $N_1 \cup N_2$  (selective union) with  $p = 0.5$ ,  $N_1 \sqcup N_2$  (strong union) and  $N_1 \rightarrow N_2$  (token-ring).

### 4.5.1 Test Instances and Experimental Protocol

Two sets of test problems are considered in our experiments. The first set of benchmarks is composed of the ten largest instances of size  $n = 2500$  introduced in [7] and available in the ORLIB [6]. These instances are used in the literature by many authors (e.g., [7, 22, 33, 35, 36]). The second set of benchmarks consists of a set of 15 randomly generated large problem instances named p3000.1, ..., p5000.5 with sizes ranging from  $n=3000$  to 5000 [35, 36]. These instances are available at: [http://www.soften.ktu.lt/~gintaras/ubqop\\\_its.html](http://www.soften.ktu.lt/~gintaras/ubqop\_its.html).

Our algorithm is programmed in C and compiled using GNU GCC on a PC running Windows XP with Pentium 2.66GHz CPU and 512M RAM. For each run of the TS algorithm, the initial solution is generated randomly, i.e., each variable  $x_i$  receives a random value of 0 or 1 with equal chance. Given this stochastic nature of our TS procedure, each problem instance is independently solved 20 times. To make the comparison as fair as possible, all the experiments use the same CPU time limits: For the ten Beasley instances with 2500 variables, the CPU time limit is set to 1000 seconds while it is set to 2000 seconds for the other 15 larger instances.

### 4.5.2 Computational Comparison

#### 4.5.2.1 Average Results Comparison

Table 4.1 shows the computational statistics of the TS algorithm on the ten Beasley instances with 2500 variables. Columns 2 and 3 give the density (dens) and the best known objective values ( $f_{prev}$ ) obtained from the literature, respectively. Columns 4 to 8 give the solution gap to the best solutions for each neighborhood and neighborhood combination. For each instance, the solution gap in Table 4.1 is represented as  $f_{prev} - \bar{f}$ , where  $\bar{f}$  is the average objective value obtained by 20 independent runs and  $f_{prev}$  represents the previous best known objective value. The overall results, averaged over ten instances, are presented in the last row.

From Table 4.1, we observe that neighborhood  $N_1$  reaches the previous best known results very stably for eight of the ten instances while it performs quite poorly on the other two cases. On the other hand, neighborhood  $N_2$  can obtain the previous best known results each time only for two instances, but obtains optimal or near-optimal solutions with quite small variance for the other cases. In terms of the average gaps to the previous best solutions, neighborhood  $N_2$  slightly outperforms  $N_1$  for these ten test problems.

When comparing the three neighborhood combinations  $N_1 \cup N_2$  (with  $p = 0.5$ ),  $N_1 \sqcup N_2$  and  $N_1 \rightarrow N_2$  with each other, one finds that the selective union  $N_1 \cup N_2$  and the token-ring search  $N_1 \rightarrow N_2$  are superior to the strong union  $N_1 \sqcup N_2$ , as well as the single neighborhoods  $N_1$  and  $N_2$ . One also observes that the strong union  $N_1 \sqcup N_2$  performs much worse than the single neighborhood  $N_1$ , implying that the strong union is not an appropriate way of combination for these two neighborhoods. For each

Table 4.1: Results of the TS algorithm on the ten Beasley instances with size  $n=2500$  within 1000 seconds

instance	dens	$f_{prev}$	solution gaps to $f_{prev}$ ( $f_{prev} - \bar{f}$ )				
			$N_1$	$N_2$	$N_1 \cup N_2$	$N_1 \sqcup N_2$	$N_1 \rightarrow N_2$
b2500.1	0.1	1515944	0	4.2	0	94.0	0
b2500.2	0.1	1471392	0	12.1	0	65.1	0
b2500.3	0.1	1414192	94.1	1.4	0	301.2	0
b2500.4	0.1	1507701	0	0	0	0	0
b2500.5	0.1	1491816	0	0	0	0	0
b2500.6	0.1	1469162	0	1.3	0	82.1	0
b2500.7	0.1	1479040	35.7	1.3	0	122.5	0
b2500.8	0.1	1484199	0	8.2	0	10.2	3.5
b2500.9	0.1	1482413	0	10.9	0	1.9	0
b2500.10	0.1	1483355	0	4.0	0	0	0
average			12.98	4.34	0	67.7	0.35

pairwise comparison of these neighborhoods, we performed a 95% confidence t-test to compare their solution quality, leading to the following ranking of the neighborhoods: For single neighborhoods  $N_2 > N_1$  while  $N_1 \cup N_2 > N_1 \rightarrow N_2 > N_1 \sqcup N_2$  for neighborhood combinations.

Table 4.2: Results of the TS algorithm on the 15 large random instances with variables ranging from 3000 to 5000 within 2000 seconds

instance	dens	$f_{prev}$	solution gaps to $f_{prev}$ ( $f_{prev} - \bar{f}$ )				
			$N_1$	$N_2$	$N_1 \cup N_2$	$N_1 \sqcup N_2$	$N_1 \rightarrow N_2$
p3000.1	0.5	3931583	319.8	8.8	103.4	1866.7	435.0
p3000.2	0.8	5193073	418.6	103.5	193.2	214.7	120.7
p3000.3	0.8	5111533	482.7	637.6	488.7	508.9	607.2
p3000.4	1.0	5761822	77.0	38.6	0	630.1	57.7
p3000.5	1.0	5675625	460.4	385.3	223.8	738.0	655.0
p4000.1	0.5	6181830	0	15.2	0	1249.3	0
p4000.2	0.8	7801355	1732.1	1364.5	402.2	2683.4	1622.7
p4000.3	0.8	7741685	1427.9	474.1	445.1	1742.0	936.1
p4000.4	1.0	8711822	1516.3	276.8	438.1	1954.1	1359.4
p4000.5	1.0	8908979	2979.9	372.5	397.4	3053.2	2723.7
p5000.1	0.5	8559355	2957.9	1287.3	1153.6	4118.4	2365.4
p5000.2	0.8	10836019	3561.5	2232.6	2716.5	3839.8	3263.0
p5000.3	0.8	10489137	8451.0	4156.4	3054.0	9634.5	3124.5
p5000.4	1.0	12252318	4760.2	3261.0	2215.4	9276.1	3416.3
p5000.5	1.0	12731803	6327.0	1369.3	1472.8	4863.2	6093.6
average			2364.82	1065.57	886.95	3091.49	1785.35

Similarly, the computational results of the TS algorithm on the fifteen larger and denser random instances are shown in Table 4.2. The symbols are the same as those in Table 4.1. Once again, we observe that neighborhood  $N_2$  outperforms  $N_1$  except for two instances (p3000.4 and p4000.1) in terms of the average gaps to the previous best known objective values. In addition, the selective union  $N_1 \cup N_2$  is superior to other two neighborhood combinations. We also performed a 95% confidence t-test to compare different neighborhoods and observed that  $N_2 > N_1$  for single neighbor-



hoods while  $N_1 \cup N_2 > N_1 \rightarrow N_2 > N_1 \sqcup N_2$  for neighborhood combinations. These results coincide well with the results observed on the ten Beasley instances with 2500 variables.

#### 4.5.2.2 Best Results Comparison

We now turn our attention to the *best results* that the TS algorithm obtains in the preceding experiments. Note that there is no difficulty to obtain the previous best known results for each neighborhood or neighborhood combination for all the ten instances with 2500 variables. Thus, we only list in Table 4.3 the best results of the TS algorithm on the fifteen larger instances. Columns 2 and 3 recall the density (dens) and the best known objective values ( $f_{prev}$ ) obtained from the literature. Columns 4 to 8 give the solution gap to the best solutions for each neighborhood and neighborhood combination, where  $f_{best}$  represents the best objective value obtained over 20 independent runs. Once again, the overall results averaged over fifteen instances are presented in the last row.

Table 4.3: Best results of the TS algorithm over 20 independent runs

instance	dens	$f_{prev}$	solution gaps to $f_{prev}$ ( $f_{prev} - f_{best}$ )				
			$N_1$	$N_2$	$N_1 \cup N_2$	$N_1 \sqcup N_2$	$N_1 \rightarrow N_2$
p3000.1	0.5	3931583	0	0	0	0	0
p3000.2	0.8	5193073	0	0	0	0	0
p3000.3	0.8	5111533	0	0	0	0	0
p3000.4	1.0	5761822	0	0	0	0	0
p3000.5	1.0	5675625	0	0	0	0	0
p4000.1	0.5	6181830	0	0	0	0	0
p4000.2	0.8	7801355	0	0	0	1686	0
p4000.3	0.8	7741685	0	0	0	0	0
p4000.4	1.0	8711822	0	0	0	0	0
p4000.5	1.0	8908979	0	0	0	0	0
p5000.1	0.5	8559355	0	-325	-325	0	0
p5000.2	0.8	10836019	582	65	0	582	0
p5000.3	0.8	10489137	354	148	148	683	663
p5000.4	1.0	12252318	608	0	0	2400	0
p5000.5	1.0	12731803	1025	0	0	0	0
average			256.9	-11.2	-17.7	366.5	66.3

Table 4.3 shows that the selective union  $N_1 \cup N_2$  performs much better than the strong union  $N_1 \sqcup N_2$ , the token-ring search  $N_1 \rightarrow N_2$  and the single neighborhoods  $N_1$ , and even slightly better than  $N_2$ . One also observes that for the fifteen larger instances, the TS algorithm with  $N_1 \cup N_2$  matches the previous best results for thirteen of them, while getting a worse result only for one instance and a better result for the remaining one. It should be noticed that the selective union  $N_1 \cup N_2$  and the single neighborhood  $N_2$  both improve the best result obtained by [36] for instance p5000.1, showing the advantage of the newly introduced neighborhood  $N_2$  over  $N_1$  and the combination mechanism of selective union. According to these results, we have the

following ranking of the neighborhoods:  $N_1 \cup N_2 > N_2 > N_1 \rightarrow N_2 > N_1 > N_1 \sqcup N_2$ . The trends of the best costs perfectly match those of the average costs mentioned above for the considered instances.

#### 4.5.2.3 Results Analysis

The preceding computational results show that for the three neighborhood combinations of  $N_1$  and  $N_2$ , the selective union  $N_1 \cup N_2$  produces much better results than other combinations. These results prompt us to focus on investigating the best and worst neighborhood combinations:  $N_1 \cup N_2$  and  $N_1 \sqcup N_2$ . In this section, we attempt to explain what causes the effectiveness and weakness of these two neighborhood unions and show evidence for this phenomenon in terms of three evaluation criteria. For this purpose, we employ a steepest descent (SD) algorithm for this experiment, where we disable the tabu list of our TS algorithm and the current solution is repeatedly replaced by a *best improving* solution in its neighborhood until no improving neighbor exists. The experiment is carried out on the large instance p5000.3 (very similar results are observed for other instances).

In [30], three evaluation criteria were employed to characterize the search capacity of a neighborhood: *Percentage of improving neighbors*, *improvement strength* and *search steps*. The authors argue that good neighborhoods should have one or more of these features: High percentage of improving neighbors (for more improvement possibilities), strong improvement strength (for important improvements) and long search steps (for long term improvements).

For a candidate solution  $x$ , a given neighborhood function  $NB : \mathcal{X} \rightarrow 2^{\mathcal{X}}$  and a neighborhood solution  $x' \in NB(x)$ , define  $\Delta f = f(x') - f(x)$ . These criteria are then defined as follows.

- *Improving neighbors*  $I(x)$ : The set of the improving neighbors in the neighborhood  $NB(x)$  given by  $I(x) = \{x' \in NB(x) \mid \Delta f > 0\}$ . Therefore, the *percentage of improving neighbors* is defined as  $|I(x)| / |NB(x)| \times 100\%$ .
- *Improvement strength*  $\Delta f^*$ : The cost variation between the current solution  $x$  and a best improving neighbor given by  $\Delta f^* = \max\{|\Delta f| : \Delta f \in I(x)\}$ .
- *Search steps*: The term *search steps* is defined as the number of iterations that the SD algorithm can run to reach a local optimum.

To calculate the values of each criterion, we run the SD algorithm for fifty independent runs with  $N_1 \cup N_2$  and  $N_1 \sqcup N_2$ , respectively. For each run, data corresponding to the above three evaluation criteria are calculated; *percentage of improving neighbors* and *improvement strength* values are collected at each iteration while *search steps* is simply the iteration number when SD stops. All the reported results correspond to the averages obtained for these fifty independent runs.

Figure 4.1 presents the percentage of improving neighbors for  $N_1 \cup N_2$  and  $N_1 \sqcup N_2$ , evolving with the local search iterations. It shows that at the beginning of the local search, the percentage of improving neighbors for the strong union  $N_1 \sqcup N_2$  is greater than that of the selective union  $N_1 \cup N_2$ . However, this trend only lasts for

the first 1000 local search iterations and then the percentage of improving neighbors for  $N_1 \sqcup N_2$  decreases dramatically during the following 300 iterations. On the other hand, the percentage of improving neighbors for  $N_1 \cup N_2$  decreases quite slowly during the first 1500 iterations. In other words,  $N_1 \cup N_2$  offers more opportunities to find improving neighbors, especially after the first iterations of the search (first 1000 iterations for this particular instance). When starting from a random initial solution even poor neighborhoods can have a certain number of improving neighbors at the first iterations while only good neighborhoods offer improving neighbors when the search progresses.

On the other hand, compared with  $N_1 \sqcup N_2$ , there exist long tails for the percentage of improving neighbors for the selective union  $N_1 \cup N_2$ , meaning that it allows the descent algorithm to run a larger number of iterations. This property is another important indicator of good neighborhoods. We argue that, in applying different neighborhoods, one that permits a descent method to continue for a greater number of iterations, and to move a greater distance during these iterations, has a greater potential to improve the solution quality in the long run.

We then evaluate the two neighborhood unions using the *improvement strength* criterion. Figure 4.2 presents how the *improvement strength* of each neighborhood evolves with the local search iterations. It shows that these two neighborhood unions have quite similar evolving trends in terms of the average improvement strength. Once again, one observes that at the beginning of the search, the improvement strength of  $N_1 \sqcup N_2$  is greater than that of  $N_1 \cup N_2$ . However, it only lasts for a small

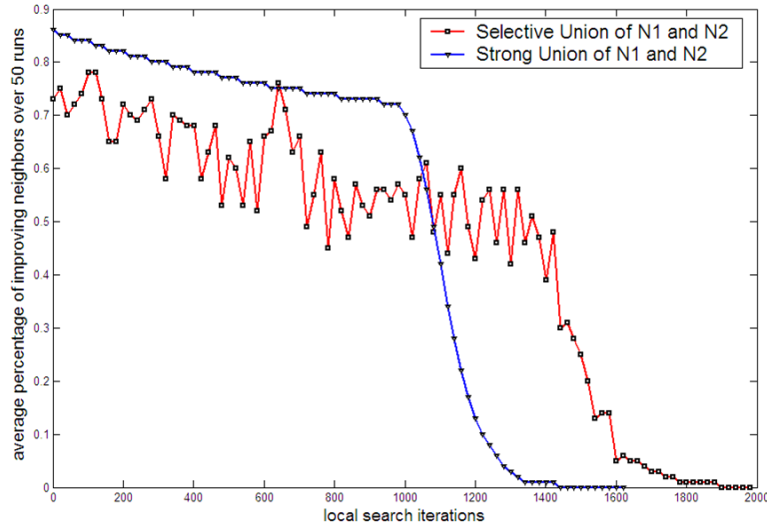


Fig. 4.1: The improving neighbors comparison between  $N_1 \cup N_2$  and  $N_1 \sqcup N_2$

number of iterations (the first 600 iteration in this particular case). This can be explained by the fact that  $N_1 \sqcup N_2$  simultaneously considers two neighborhoods  $N_1$  and  $N_2$  while  $N_1 \cup N_2$  only randomly chooses one neighborhood. Nevertheless, as the algorithm progresses (after the first 600 iterations), the  $N_1 \cup N_2$  neighborhood offers much greater improvement strength than  $N_1 \sqcup N_2$ . This phenomenon correlates well with the trend of the percentage of improving neighbors.

Based on these observations, we formulate the following conclusions.

1. Neighborhood union  $N_1 \cup N_2$  induces a higher percentage of improving neighbors and greater improvement strength than  $N_1 \sqcup N_2$  after the first iterations of the search. As a result,  $N_1 \cup N_2$  offers more choices for the search algorithm to improve the current solution at each iteration once the initial iterations are completed.
2. Neighborhood  $N_1 \cup N_2$  offers improving neighbors for a larger number of iterations than  $N_1 \sqcup N_2$ . Consequently, local search can continue for a larger number of iterations with  $N_1 \cup N_2$ .
3. Although neighborhood  $N_1 \sqcup N_2$  offers a higher percentage of improving neighbors and greater improvement strength during the first iterations, its improvements quickly disappear, limiting its search capability.

We also repeated this experiment with the TS algorithm described in Section 4.4.2 and reached similar conclusions.

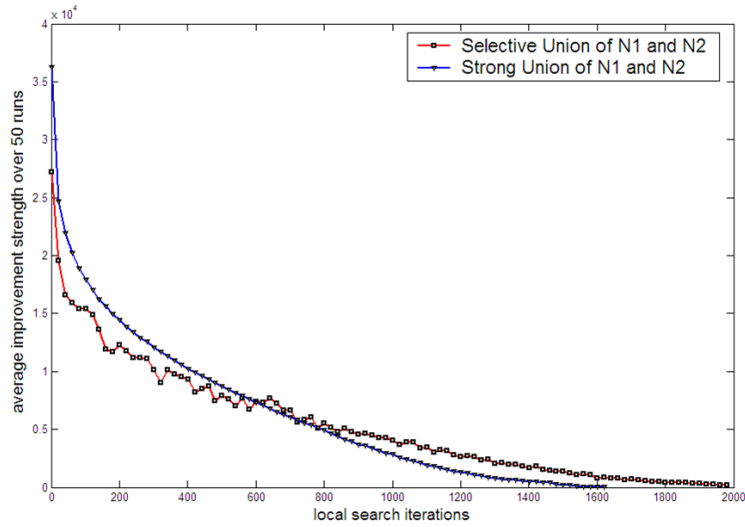


Fig. 4.2: The improvement strength comparison between  $N_1 \cup N_2$  and  $N_1 \sqcup N_2$

## 4.6 Conclusions and Discussion

In this paper, we compare and analyze two basic neighborhoods (one-flip and two-flip) and three neighborhood combinations (*selective union*, *strong union* and *token-ring search*) for the UBQP problem. The computational results show that the best outcomes are achieved with the selective union  $N_1 \cup N_2$ , followed by using the new  $N_2$  neighborhood by itself.

We employ three evaluation criteria to explain why the selective union  $N_1 \cup N_2$  performs much better than the strong union  $N_1 \sqcup N_2$ , yielding an experimental analysis that sheds light on the relative advantages and weaknesses of the neighborhoods  $N_1$  and  $N_2$  and various possibilities for combining them. Our findings are anticipated to have useful implications for combining neighborhoods in other applications, particularly when presented a choice between the use of selective unions and strong unions.

Some important questions remain.

1. These results are based on *random* instances. It would be interesting to know whether these results would be confirmed for problems that exhibit special structures of various types. To this end, a sequel to this study will carry out additional experiments using more diverse instances transformed from other problems.
2. It would be useful to identify the conditions under which a particular neighborhood or a neighborhood combination is preferable.
3. More importantly, it would be valuable to explore higher order neighborhood moves (e.g., three-flip or even higher flip moves). As observed in [14], there exists a natural way to extend the above mentioned fast two-flip move evaluation techniques to these higher order moves.
4. It would be worthwhile to investigate other ways of combining the neighborhoods, particularly with the inclusion of higher order flip moves. For example, we may consider “conditional” combinations where moves from a lower order neighborhood pass certain screening criteria as a foundation for becoming components of moves in higher order neighborhoods.

We anticipate that answers to these issues will provide information that will be valuable for the design of improved algorithms. Finally, given that the neighborhood combination strategies and the neighborhood evaluation criteria discussed in this paper are independent of the UBQP problem, they can be used to evaluate neighborhood relations of other combinatorial optimization problems.

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