
Leveraging TSP Solver Complementarity through Machine Learning

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Abstract

The Travelling Salesperson Problem (TSP) is one of the best-studied NP-hard problems. Over the years, many different solution approaches and solvers have been developed. For the first time, we directly compare five state-of-the-art inexact solvers – namely, LKH, EAX, restart variants of those, and MAOS – on a large set of well-known benchmark instances and demonstrate complementary performance, in that different instances may be solved most effectively by different algorithms. We leverage this complementarity to build an algorithm selector, which selects the best TSP solver on a per-instance basis and thus achieves significantly improved performance compared to the single best solver, representing an advance in the state of the art in solving the Euclidean TSP. Our in-depth analysis of the selectors provides insight into what drives this performance improvement.

Keywords

Travelling Salesperson Problem, automated algorithm selection, performance modeling, machine learning

1 Introduction

The Travelling Salesperson Problem (TSP) is arguably the most prominent NP-hard combinatorial optimisation problem. Given a set of n locations – which, by convention, are called *cities* – and pairwise distances between those cities, the objective in the TSP is to find the shortest round-trip or *tour* through all cities, i.e., a sequence in which every city is visited exactly once, the start and end cities are identical, and the total length of the tour is minimal. Here, we consider the *2D Euclidean TSP*, in which the cities correspond to points in the Euclidean plane and the distances are Euclidean distances. This is the most commonly studied special case of the TSP, and, like the general TSP, it is known to be NP-hard. The Euclidean TSP has important applications, e.g., in the fabrication of printed circuit boards as well as in transportation and logistics.

There are two types of TSP algorithms: exact algorithms, which are guaranteed to find an optimal solution to any TSP instance when run to completion and produce a proof of optimality; and inexact algorithms, which cannot guarantee or prove the optimality of the solutions found. Intriguingly, the state of the art for both types of algorithms has been defined by a single solver each for many years: the exact solver Concorde (Applegate et al., 2007) and the inexact solver LKH (Helsgaun, 2009). Furthermore, LKH typically finds high-quality and even optimal solutions much more quickly than Concorde, and therefore, for the purpose of finding such solutions, per-instance algorithm selection techniques, (see, e.g., Kotthoff (2014)) were previously inapplicable to the TSP.

More recently, improvements in the state of the art in inexact TSP solving in the form of a new evolutionary algorithm, EAX (Nagata and Kobayashi, 2013), and a new multi-agent optimization approach, MAOS (Xie and Liu, 2009), have made per-instance algorithm selection approaches feasible.

In our previous work (Kotthoff et al., 2015), we showed for the first time that per-instance algorithm selection techniques can be used to improve the state of the art in inexact TSP solving. Here, we expand this preliminary study by including the MAOS solver, additional instance features, and structured instance sets. We furthermore perform cost-sensitive feature selection for the algorithm selection models. We will show that this step is crucial to achieve good performance, as otherwise the single best solver (the restart variant of EAX) dominates. The best selector we achieve substantially improves over this single best solver – by a factor of two when aggregated over our instance set, and up to an order of magnitude on individual instances.

After providing background information on algorithm selection in Section 2, we describe the solvers, the TSP instance sets, and the features we use as the basis for per-instance selection in Section 3. We then define our benchmark in Section 4 and describe the experimental setup in Section 5. We report the performance improvements we have achieved (Section 6), before concluding with some general observations and directions for future work (Section 7).

2 Background on Automated Algorithm Selection

The per-instance algorithm selection problem (Rice, 1976) involves selecting from a set of candidate algorithms the one expected to perform best on a given problem instance. Similar to parallel algorithm portfolios (Huberman et al., 1997; Gomes and Selman, 2001), per-instance algorithm selection leverages complementarity within a set of solvers.

Algorithm selection systems typically use machine learning to forecast how well the algorithms in a given set will perform on a given problem instance based on features of that instance. Some approaches predict the algorithm to use directly, while others aggregate performance predictions to select one or more algorithms from the given set to be run sequentially or in parallel. Here, we consider the case where exactly one algorithm is selected per instance.

One of the most prominent and successful systems that employs this approach is SATzilla (Xu et al., 2008, 2012), which for many years defined the state of the art in solving the propositional satisfiability problem (SAT), one of the widely studied NP-complete problems in artificial intelligence with important applications in hard- and software verification. Following the initial success of early versions of SATzilla, other algorithm selection systems have been developed and proven their worth in the annual SAT competition, such as CSHC (Malitsky et al., 2013), which has also been applied to

the closely related MAXSAT problem.

Algorithm selection systems have also been applied with great success to other computationally challenging problems, including constraint programming (O'Mahony et al., 2008), continuous black-box optimization (Mersmann et al., 2011; Bischl et al., 2012), mixed integer programming (Xu et al., 2011), and AI planning (Seipp et al., 2012). There have been several prior studies on algorithm selection for the TSP, e.g., Fukunaga (2000); Pihera and Musliu (2014); however, to the best of our knowledge, only one of these – our own preliminary study for the work presented here – can claim to have achieved improvements in the state of the art (Kotthoff et al., 2015). For additional background on algorithm selection and its application, we refer the interested reader to the comprehensive survey by Kotthoff (2014).

3 Methodology

We first describe the TSP algorithms (Section 3.1), instance feature sets (Section 3.2), and benchmark instances (Section 3.3) considered in our study. We then provide an overview of the supervised learning methods that we utilized as algorithm selectors (Section 3.4), as well as the statistical tests (Section 3.5) we used to show whether observed performance differences between models are statistically significant.

3.1 Optimization Algorithms

3.1.1 LKH: Helsgaun's Lin-Kernighan heuristic

The Helsgaun variant of the Lin-Kernighan heuristic (LKH) represented a major advance in inexact TSP solving, and has been the uncontested state-of-the-art method for finding high-quality solutions to a large variety of TSP instances for many years (Helsgaun, 2000, 2009). The Lin-Kernighan heuristic (LK) is a variable-depth search method that generates complex local search moves by heuristically constructing a sequence of edge exchanges. LKH, Helsgaun's variant of LK, is based on exchange sequences using five (Helsgaun, 2000) or more edge exchanges (Helsgaun, 2009). The iterated version of the LKH algorithm restarts the local search process from new solutions that are obtained by solution perturbations, which are performed using either a random k -exchange move or a special walk strategy. An approximation of the Held-Karp lower bound is used to obtain small candidate sets for the local search steps.

We use the most recent version (2.0.7) of LKH, denoted simply as LKH in the following. After observing stagnation behaviour for that solver, Dubois-Lacoste et al. (2015) enhanced it with a restart mechanism that triggers if during n iterations, where n is the size of the TSP instance, no improving solution was found. We denote this modified version LKH+restart.

3.1.2 EAX: GA with edge assembly crossover

A successful line of research into high-performing evolutionary algorithms for the TSP are evolutionary algorithms that integrate variants of edge assembly crossover – a recombination operator that combines the edges of two parent solutions trying to add only few, short edges not found in any of the two parents (Nagata and Kobayashi, 1997). The first such algorithm known to have matched the performance of LKH in finding very high quality solutions to a broad range of Euclidean TSP instances (as considered in our work) is EAX, the evolutionary algorithm by Nagata and Kobayashi (2013). In a nutshell, EAX exploits improved local and global variants of the edge assembly crossover operator, specific diversity preservation techniques that uses edge entropy measures in the population replacement scheme, and initialization of the population

by local optimization. For a detailed description of the algorithm and the operators, we refer the reader to the original publication.

The original version of EAX does not support termination upon reaching a given solution quality and used a complex termination criterion to end runs. Therefore, Dubois-Lacoste et al. (2015) modified this version to terminate upon reaching a target solution quality (in our experiments always set to the known optimum for the given instance) or when a given time limit is exhausted. A second variant restarts whenever the original termination criterion is met (EAX+restart). Both versions of EAX were also modified to permit setting a random seed.

We consider the same versions of LKH and EAX as Dubois-Lacoste et al. (2015), where more details on the algorithms and the restart behaviour can be found.

3.1.3 MAOS: Multi-agent optimization

In contrast to LKH and EAX, MAOS is based on a complex, multi-agent framework-based optimization approach, where the agents have only limited knowledge of the TSP instance and explore possible solutions in parallel (Xie and Liu, 2009). The agents update the shared environment and thus communicate their acquired knowledge to the other agents to improve the efficiency of the search process. Specifically, each agent starts from a set of reference structures, such as disjoint graph elements that can be combined into valid tours, and candidate sets, such as nearest neighbour subgraphs. It then uses a Markov-chain approach to generate a set of states by applying search operators, such as local search steps, genetic algorithm crossovers, or simple completion heuristics. There is some evidence in the literature that at least on part of the instance sets we consider here, MAOS reaches and possibly exceeds the performance of the original versions of LKH and EAX (Pihera and Musliu, 2014; Xie and Liu, 2009). Similarly to EAX, we modified MAOS to terminate when reaching a given solution quality; following Xie and Liu (2009), we set the number of agents to 300.

3.2 Feature Sets

There are several approaches in the literature that attempt to characterise TSP instances by computing features. We focus on the three presented by Mersmann et al. (2013)¹, Hutter et al. (2014)², and Pihera and Musliu (2014), as they comprise a large and diverse set of features, and we consider the respective feature sets in isolation as well as combined with each other. Obviously, the cost of computing the feature values for a given instance can play a major part in the efficacy of an algorithm selection system. We therefore added a subset of the features of Hutter et al. (2014) that are cheaper to compute than the full set. Overall, we used the following five sets of features in our experiments:

UBC (cheap) The feature set by Hutter et al. (2014) without the more expensive features, in particular, the local search, branch and cut, and clustering distance features (13 features). The mean time of computing this set of features was 0.875 seconds per instance, with the median at 0.6 seconds and a standard deviation of 11.6 seconds.

UBC The set of all 50 features from the work by Hutter et al. (2014). The mean time of computing this feature set was 19.87 seconds per instance, with the median at 12.72 seconds and a standard deviation of 96.587 seconds.

¹<https://CRAN.R-project.org/package=tspmeta>

²http://www.cs.ubc.ca/labs/beta/Projects/EPMS/TSP_features_UBC2012.tar.gz

TSPmeta The feature set (68 features) by Mersmann et al. (2013). The mean time for computing this feature set was 11.66 seconds per instance, with the median at 10.5 seconds and a standard deviation of 19.519 seconds.

Pihera The set of all 287 features from the study by Pihera and Musliu (2014). The mean time of computing this set of features was 0.255 seconds per instance, with the median at 0.24 seconds and a standard deviation of 0.41 seconds.

$UBC \cup TSPmeta \cup Pihera$ The union of *UBC*, *TSPmeta*, and *Pihera* (405 features). Note that some of the features in the constituent sets contain the same information.

3.3 TSP Instances

Consistent with other work in this area, we use instances from multiple TSP benchmarks and generators. The optimal tour length for each of the instances we used was obtained using Concorde (Applegate et al., 2007). Note that we limited our study to instances for which the optimal tour length can be obtained within reasonable time in this way or has been published previously. This permitted us to determine with certainty whether the solvers we studied found an optimal solution³ and how long they had to be run to achieve this. This is the most ambitious goal for any TSP solver, and even though inexact solvers, such as the ones we consider here, cannot prove optimality, they are typically able to find solutions whose optimality is later proven using exact methods, and knowledge of optimal or near-optimal solutions is known to greatly facilitate this process. In the following, we give a brief overview of the types of TSP instances we used, while details of the benchmark set will be discussed later, in Section 4.

RUE instances The widely studied *random uniform Euclidean (RUE)* instances are obtained by placing n points uniformly at random in a square, with integer coordinates between 1 and 1 000 000, where each point corresponds to a city to be visited. Distances between these cities are defined as Euclidean distances between the respective points, rounded to the nearest integer. The RUE instances used in our experiments were generated using the `portgen` generator from the 8th DIMACS Implementation Challenge.

TSPLIB instances *TSPLIB* is a widely-used collection of TSP instances with different characteristics, including instances from various applications of the TSP. In our experiments, we used instances with edge weight types EUC 2D, CEIL 2D and ATT.

National instances The *national* instances are one of two sets of instances we obtained from the TSP webpage⁴ and are based on the real-world locations of cities in different countries.

VLSI instances The *VLSI* instances are the second set of instances from the TSP webpage and originate from an application in VLSI circuit design. These instances are known to be particularly hard for many TSP solvers, including Concorde and EAX.

An example instance for each of the aforementioned four benchmark sets is depicted in Figure 1. While instances from these four sets were already used in our initial work on algorithm selection for the TSP (Kotthoff et al., 2015), clustered instances were not considered then. These types of instances play an important role in many TSP applications (e.g., vehicle routing), and we added two sets of clustered TSP instances to our overall benchmark.

³There may be multiple tours with the same optimal tour length.

⁴<http://www.math.uwaterloo.ca/tsp/index.html>

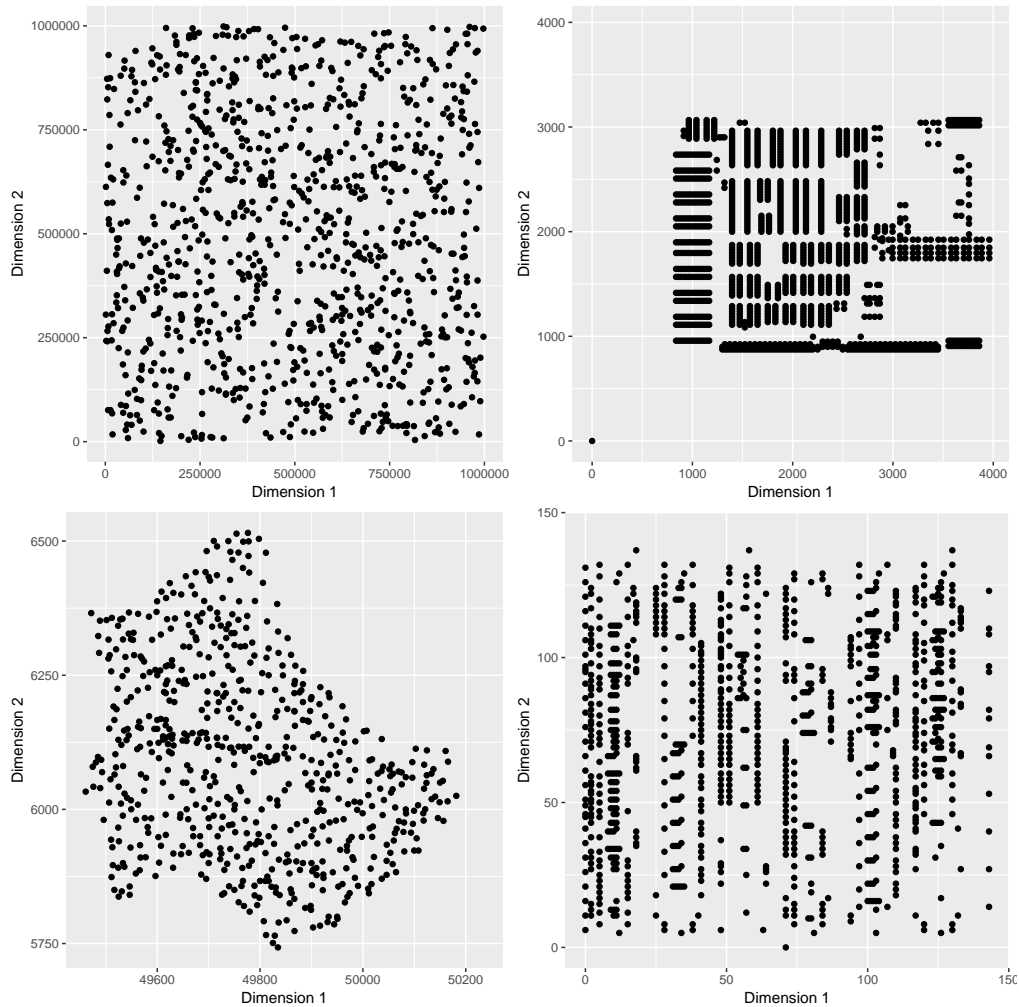


Figure 1: Examples of TSP benchmark instances: 1000-2 (RUE; top left), d1291 (drilling problem; TSPLIB; top right), 1u980 (980 cities in Luxembourg; National; bottom left) and pbd984 (VLSI; bottom right).

Netgen instances This set of clustered instances, generated using the function `generateClusteredNetwork` from the R-package `netgen` (Bossek, 2015), consists of multiple instances for each combination of the number of cities $n \in \{500, 1000, 1500, 2000\}$ and clusters $n_c \in \{2, 5, 10\}$.

The instance generation process was introduced in Meisel et al. (2015) and works as follows: First a maximin Latin hypercube sample (LHS, M. D. McKay (1979)) of size n_c is generated in two dimensions, i.e., n_c points are placed in the two-dimensional space maximizing the minimal distance between design points. The space-filling property of LHS designs ensures that the cluster centres are evenly distributed across the space. Next, random points are sampled from a multivariate normal distribution per cluster using the cluster centres as the mean value vector and the distance to the nearest cluster

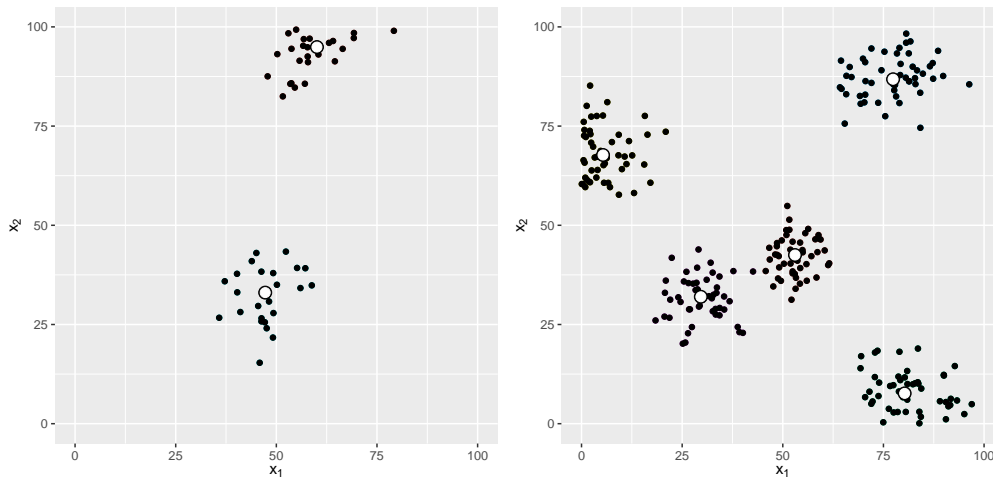


Figure 2: Examples of netgen TSP instances with parameters $n_c = 2$ and $n = 50$ (left), and $n_c = 5$ and $n = 250$ (right) on a $[0, 100] \times [0, 100]$ grid. The white points indicate the locations of the n_c cluster centers (determined by means of maximin Latin hypercube sampling), and the black points indicate the actual cities to be visited.

centre as the variance in each dimension. The number of points is equally distributed across the clusters. Figure 2 shows two examples of such instances: one with $n_c = 2$ and $n = 50$ (left) and one with $n_c = 5$ and $n = 250$ (right).

Morphed instances In addition, we created *morphed* instances, which combine properties of RUE and clustered netgen instances, by using the morphing algorithm introduced by Mersmann et al. (2012) and improved by Meisel et al. (2015). It combines two instances with the same number of cities n using the following two-phase approach.

First, corresponding points in the two instances are matched by minimizing the sum of Euclidean distances between matched points across all points through a linear programming formulation. Second, a convex combination $z_{ij} = \alpha \cdot x_{ij} + (1 - \alpha) \cdot y_{ij}$ of the node coordinates of paired points x_{ij} and y_{ij} , where $i = 1, \dots, n$ and $j = 1, 2$, determines the node coordinates in the corresponding morphed instance z_{ij} for a given morphing coefficient $\alpha \in [0, 1]$. Extreme values of $\alpha \in \{0, 1\}$ result in one of the input instances, while other values of α generate intermediate instances, as illustrated in Figure 3.

The morphed instances used in our experiments were generated by applying the function `morphInstances`, available in the R package `netgen`, with morphing factor $\alpha = 0.5$ to randomly selected instance pairs from the netgen and RUE sets.

3.4 Supervised Learning Methods for Algorithm Selection

A key aspect of this work is finding a suitable – i.e., high-performance – algorithm selection model. We evaluated three supervised learning strategies, namely *classification*, *regression* and *paired regression*, with multiple machine learning techniques for each.

In the classification approach, we label each instance with the best-performing TSP solver. In the regression approach, we predict the runtime for each solver on an instance

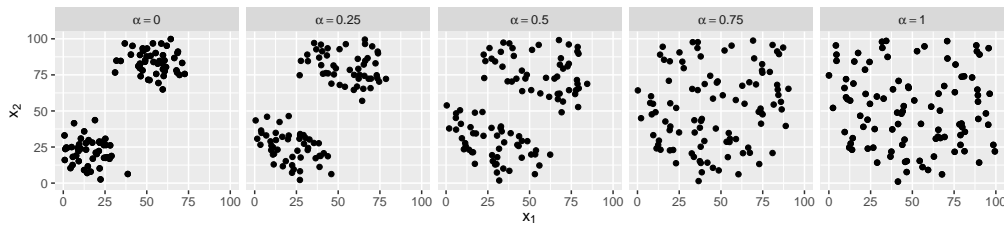


Figure 3: Example of instance morphing of a 2-cluster instance into a RUE instance with $n = 100$ nodes. A sequence of increasing morphing coefficients $\alpha \in \{0, 0.25, 0.5, 0.75, 1\}$ illustrates the transition between both instances.

separately and choose the solver with the lowest predicted runtime. The pairwise regression approach predicts the performance difference between two solvers for each pair of solvers. The solver with the best predicted performance difference to all other solvers is chosen. Further details regarding the setup and implementation of the three strategies can be found in Section 5.1.

The algorithm selectors itself were built using the R-package `mlr` (Bischl et al., 2016b). Using this machine learning toolbox, we trained recursive partitioning and regression trees (`rpart`, Therneau et al. (2015)), random forests (`randomForest`, Liaw and Wiener (2002)) and kernel-based support vector machines (`ksvm`, Karatzoglou et al. (2004)) for each strategy. In addition, multivariate adaptive regression splines (`mars`, Leisch et al. (2016)) were trained for the two regression approaches. Further details on MARS models can e.g. be found within Friedman (1991) or our Appendix.

Each of the methods was used with its default settings, i.e., without additional hyperparameter optimisation. Specifically, the random forests were built with 500 trees, using a uniform random sample of $\lfloor \sqrt{p} \rfloor$ (classification) or $\max\{\lfloor p/3 \rfloor, 1\}$ (regression / paired regression) of the p features for each data set as candidates for each split point. The support vector machines used a Gaussian kernel with an *a priori* estimate of the inverse kernel width parameter, `sigma`, obtained using the `sigest` function from the R-package `kernlab` (Karatzoglou et al., 2004).

After training our algorithm selection systems based on these models, we assess their performance on a given TSP instance by determining the running time of the solver selected on that instance plus the running time required for computing the instance features used by the selector. To get reliable generalization performance estimates on a given training set of instances, we use 10-fold cross-validation (further details will be given in Section 5).

3.5 Assessing the significance of performance differences

In general, when comparing two TSP solvers, S_1 and S_2 (one or both of which may be algorithm selectors) on the same instance set I , we want to assess whether any observed difference in performance can be considered statistically significant. Since performance is assessed using PAR10 scores for S_1 and S_2 on I , our analysis addresses the question to which extent an observed difference in PAR10 scores depends on the particular given instance set, I . We therefore used bootstrap sampling to simulate drawing new instance sets from the distribution of problem instances that gave rise to I (Fawcett et al., 2017); following common practice, each sample was constructed using uniform random sam-

pling with replacement and contains the same number of instances as I (Hastie et al., 2009). For each of the $B = 10\,000$ bootstrap samples I_1, \dots, I_B thus obtained, we computed the PAR10 scores for S_1 and S_2 .

As a first measure of statistical stability, we determined the fraction of samples in which S_1 has a better (i.e., lower) PAR10 score than S_2 . The higher this fraction, the more confident we can be that a performance advantage of S_1 over S_2 observed on benchmark set I also holds for similar sets I' (where technically, similarity means that both sets stem from the same underlying instance distribution).

To sharpen this analysis, we assessed the statistical significance of the performance difference between S_1 and S_2 on I by performing a one-sided Wilcoxon signed-rank test on the pairs of performance data ($\text{PAR10}(S_1, I_b), \text{PAR10}(S_2, I_b)$) for $b = 1, \dots, B$ setting the null hypothesis to the " \geq " relation comparing S_1 and S_2 . This standard non-parametric test for paired samples was chosen in order to avoid unrealistic assumptions regarding the nature of the distribution of the performance data (Hollander et al., 2013). We note that performing the same test on the pairs of performance values on individual instances $i \in I$ would be inappropriate, since we are interested in the difference in aggregate performance on the entire set I , rather than the performance difference on individual instances. To see why this could give misleading results, imagine an extreme case in which on a large fraction of instances in I (say: 90%) S_1 runs marginally slower than S_2 (say: 0.1% slower), while on the remaining instances, S_1 is dramatically faster than S_2 (say: 1000 times). In this case, S_1 would be much better than S_2 according to PAR10 score and preferred in most practical situations, while the Wilcoxon signed-rank test on I (rather than bootstrap samples of I) would indicate that S_1 performs significantly *worse* than S_2 , because used in this way, the test essentially ignores the magnitude of performance differences on the instances $i \in I$.

For the analysis of our experimental results described in Section 6, we used R to generate bootstrap samples and to carry out the Wilcoxon signed-rank test with a standard correction for ties (R Core Team, 2016).

4 Generating the Benchmark

First, we created a benchmark set of TSP instances based on the various instance sets introduced in Section 3.3. Using each of the instance generators, we produced an equal number of instances. In particular, for each of the instance sizes we considered, $n \in \{500, 1\,000, 1\,500, 2\,000\}$, we generated 150 RUE instances, 50 netgen instances each for $n_c \in \{2, 5, 10\}$ clusters, and 150 morphed instances obtained by combining those RUE and netgen instances with a morphing factor of $\alpha = 0.5$. Thus, we obtained a grand total of 600 instances (4×150) each in our RUE, netgen and morphed sets. While the (mostly) real-world instances in the TSPLIB, National and VLSI sets have between 48 and 11 849 cities, to achieve better balance of instance sizes across our entire benchmark, we included only instances with 500 to 2 000 cities, resulting in 22 TSPLIB, 18 VLSI and 5 National instances, as shown in Table 1.

We note that our overall benchmark set is heavily biased towards the artificially generated RUE, netgen, and morphed instances, because we considered it important for the training of the machine learning models used in our algorithm selectors to have at our disposal a large set of TSP instances. We also note that the clustered and morphed instances, which jointly account for more than half of the overall set, show structure that is quite typical for many real-world TSP instances and known to be challenging for TSP solvers (see, e.g., Christofides (1976); Solomon (1987)).

Since all five solvers we considered (EAX, EAX+restart, LKH, LKH+restart and

Table 1: Overview of the sizes and numbers of instances which were solved by at least one of the five algorithms and consist of 500 to 2 000 cities – per TSP set and in total. The problems are summarized by the minimum, first quartile (= 25%-quantile), median, arithmetic mean, third quartile (= 75%-quantile) and maximum.

TSP set	Number of Cities						Number of Instances
	Min.	1st Qu.	Median	Mean	3rd Qu.	Max.	
RUE	500.00	875.00	1 250.00	1 250.00	1 625.00	2 000.00	600 (32.5%)
VLSI	662.00	968.25	1 412.50	1 310.61	1 595.00	1 973.00	18 (1.0%)
TSPLIB	532.00	738.75	1 232.00	1 165.14	1 424.00	1 889.00	22 (1.2%)
National	734.00	929.00	980.00	1 248.60	1 621.00	1 979.00	5 (0.3%)
Netgen	500.00	875.00	1 250.00	1 250.00	1 625.00	2 000.00	600 (32.5%)
Morphed	500.00	875.00	1 250.00	1 250.00	1 625.00	2 000.00	600 (32.5%)
Total	500.00	813.00	1 304.00	1 249.58	1 583.00	2 000.00	1 845 (100.0%)

MAOS) are randomised, we assessed their performance based on ten independent runs (each with a different pseudo-random number seed) on each of the 1 845 instances from our benchmark set. As explained previously, each run was terminated as soon as an optimal solution was found or after a cutoff time of 3 600 seconds; in the first case, the run was considered successful and in the second case, unsuccessful. We then recorded the median running time over these ten runs; if at least six of the ten runs were successful, the performance on that instance thus corresponds to the arithmetic mean of the fifth- and sixth-lowest running time. When fewer than six runs were successful, the performance on that instance was recorded as ten times the cutoff time, i.e., 10 hours; this is the widely used PAR10 score (penalized average running time with penalisation factor 10).

Since the Pihera feature computation code is unable to handle instances with ATT edge weight type, we had to exclude the one instance in our set, `att532` from TSPLIB, in all experiments that included those edge weights.⁵ However, as the performance gap between the best (LKH) and worst (MAOS) solver on that instance is only 3.995 seconds, and we have 1 845 instances in total, the impact of `att532` on the PAR10 score of any solver or selector cannot exceed $3.995/1\,845 \approx 0.002$ seconds.

Analysis of the feature values revealed that six of the Pihera features (`nn3.Mdeg`, `nn3.mindeg`, `nn3.sc.min`, `nn3.avg`, `nn5.avg` and `nn7.avg`), as well as the features `mst_depth_min` (TSPmeta) and `bc_nols_q50` (UBC all) were constant across all 1 845 instances. As constant features do not provide any information to the machine learning algorithms used for algorithm selection, these eight features were discarded. We also detected that two further Pihera features, `nn5.Mdeg` and `nn3.q3deg`, were constant across 1 844 of our 1 845 instances. As explained in detail in Section 5, we evaluated our algorithm selectors using cross-validation (CV); we therefore discarded these two features as well, in order to avoid constant features within the training data of a single CV-fold. The detailed procedure for preprocessing the data is shown in Algorithm 1.

Note that we performed feature computation and solver runs on the same machine, since we later take both into account to assess the performance of our algorithm selectors.

⁵We decided not to exclude the instance from our overall benchmark set, as it is one of the most prominent TSPLIB instances.

Algorithm 1 Procedure for generating and preprocessing the training data.

```

1: procedure PREPROCESSING
2:   define available.instances as union of all instances that:
3:     - edge weight type is either EUC 2D, CEIL 2D or ATT
4:     - has solver run information and features
5:     - none of the solver runs or feature computations crashed
6:     - number of cities  $n$  per instance:
7:       - real-world problems (National, VLSI, TSPLIB):  $n \in [500, 2000]$ 
8:       - artificial problems (RUE, netgen, morphed):  $n \in \{500, 1000, 1500, 2000\}$ 
9:       - RUE instances: consider 150 instances per instance size  $n$ 
10:      - netgen problems: 50 instances per size  $n$  and number of clusters  $n_c \in \{2, 5, 10\}$ 
11:      - morphed instances: mix of RUE and netgen instances with  $\alpha = 0.5$ 
12:   create the following data sets based on available.instances:
13:     - runtimes: performance per instance, solver, and run
14:     - runstatus: status (successful / unsuccessful) per instance, solver, and run
15:     - feats: all features per instance
16:     - costs: cost of computing each feature set per instance
17:   aggregation of runtimes across all iterations per instance:
18:     if more than 5 out of the 10 runs are successful then
19:       - use median across the 10 runs (i.e., mean of the 5th and 6th fastest run)
20:       - define run as successful
21:     else
22:       - use penalized running time ( $= 10 \times \text{cutoff} = 36000$  s)
23:       - define run as unsuccessful
24:     end if
25:   clean up data sets by removing features that are constant across all (but one) instances
26:   partition instances and associated feature and performance data by uniformly random
    sampling into 10 cross-validation folds
27: end procedure

```

5 Experimental Setup

We trained the machine learning methods outlined in Section 3.4 to obtain algorithm selectors. With the exception of the scenarios in which we considered the Pihera features – and therefore had to discard TSPLIB instance `att532` (as explained in the previous section) – all selectors were trained on the same set of 1845 TSP instances, using the aggregated running times of the five TSP solvers that were introduced in Section 3.1, namely EAX, EAX+restart, LKH, LKH+restart and MAOS. Following common practice in machine learning, each selector was evaluated using 10-fold cross-validation.

Our selectors were either trained on one of the four feature sets from Section 3.2 or on the union of all four feature sets. Considering the high computational cost of these experiments and limitations of our computing resources, we did not explicitly consider combinations of two or three feature sets on top. However, as we performed feature selection based on the union of all feature sets (see Section 5.2), we implicitly also cover feature combinations that could stem from any combination of two or three feature sets.

In the context of each of the selector construction approaches outlined in Section 3.4, we considered several machine learning procedures: classification and regression trees, support vector machines, and two variants of random forests⁶. In addition,

⁶While the trees, support vector machines, and MARS models are deterministic, the random forests were trained twice per model configuration to account for stochasticity.

we also considered MARS models (see Section 3.4) in the two regression-based approaches, resulting in a total of 70 possible selectors (= 5 feature sets \times (4 classification + 5 regression + 5 paired regression machine learning procedures)).

5.1 Supervised learning strategies

Each of the three supervised learning strategies described below ultimately selects a TSP solver to use for each instance. Each of the selectors thus obtained is based on a specific feature set and consequently incurs a specific cost for determining the feature values for any given instance. These feature costs, as well as the running time of the selected TSP solver will then be averaged across all instances of the corresponding cross-validation fold, resulting in a PAR10-score per fold. These ten PAR10-scores are then again averaged across all ten folds, resulting in the algorithm selector’s PAR10-score.

Classification The first strategy trains a machine learning model that predicts the best TSP solver for a given instance. The decision of whether a solver performed best on an instance is based on the aggregated running times. However, for 30 out of 1845 instances (i.e., about 1.6% of our benchmark set), there was no clear best solver, i.e., two or more solvers shared the lowest runtime. For those instances, we used the arithmetic mean of the unaggregated running times (for that instance) as a tie-breaking criterion.

Regression While the classification approach does not consider the magnitude of the differences in running time, but solely relies on the best-performing solver on an instance, the regression approach directly models the running times of the solvers. This approach trains five models (one per solver) and afterwards selects the solver with the lowest predicted running time on the given TSP instance.

Paired Regression The third approach models the differences in running time between pairs of solvers. That is, for each pair of solvers, we model the differences in running time, resulting in a total of ten models. We sum the running time differences per solver and choose the solver with the best total difference in running time. We note that this corresponds to a weighted voting scheme over pairs of solvers.

5.2 Feature Selection

We performed feature selection on each of the 70 algorithm selectors (i.e., on the underlying machine learning models) mentioned previously, using three different feature selection approaches, resulting in a total of 210 additional algorithm selectors. We used 10-fold cross-validation to assess the performance of our models, based on PAR10-scores (including the feature computation times). The computation of many features within each of the sets from Section 3.2 is deeply intertwined; therefore, if one or more features from a given set are selected, the associated feature computation time is always the same as for the entire set. During feature selection, ties between selectors with equal performance were broken uniformly at random.

Greedy forward-backward selection Our first feature selection approach, `sffs` (sequential floating forward search), is a greedy forward-backward selection that starts with an empty set of features (Pudil et al., 1994); in our experiments, we used the variant implemented in the R-package `mlr` (Bischl et al., 2016b). In each iteration, this procedure extends the given set F by the single feature not yet in F that gives the largest improvement in selector performance. After a feature has been added, the feature whose exclusion leads to the largest improvement in performance is dropped and

the next iteration begins, i.e., a further feature is added. The process terminates when neither adding nor dropping any single feature leads to any improvement in selector performance.

Greedy backward-forward selection The second feature selection approach, *sffb*s (sequential floating backward search, Pudil et al. (1994)), works analogously to *sff*s, but in reverse. We used the variant implemented in *mlr* (Bischl et al., 2016b), which works as follows: Starting with the full feature set, in each iteration, first the feature whose removal causes the largest improvement in performance is removed from the given set F , and then the feature, which is not yet in F , but leads to the largest improvement in performance over F , is added and the next iteration begins.

Genetic Algorithm In addition to the first two feature selection approaches, we also considered a $(10 + 5)$ genetic algorithm (Eiben and Smith, 2015). Specifically, we generate ten *individuals* based on a set of randomly chosen features from the provided feature set(s), where an individual is a bit-string of selected / non-selected features, which are used for training the algorithm selector. These ten individuals form our initial *population*. Then, for each iteration (also called *generation*) of this feature selection strategy, these so-called *parents* from the current population are used to create five new feature sets, the so-called *offspring*, using a mutation rate of 5% and crossover probability of 50%. Out of the resulting 15 feature sets – ten parents and five offspring – the ten best are chosen as the starting population for the next generation of the evolutionary process. We performed 100 generations of this genetic algorithm and chose the best individual from the final population as the feature set for our final selector.

5.3 Baseline Algorithms

As done in previous work on algorithm selection, we assess our selectors against two baselines: the *virtual best solver* and the *single best solver*, to assess the performance of our algorithm selectors (Bischl et al., 2016a).

Virtual best solver (VBS) This baseline, often also called *oracle* or *perfect selector*, provides the best possible performance on the given data by always choosing the best solver at no cost in addition to that incurred by running the selected solver. This idealised procedure provides an upper bound for the performance of any algorithm selector; because of imperfect selection and feature computation cost, the performance of actual algorithm selectors usually falls short of that of the VBS.

Single-best solver (SBS) Our second baseline is given by the single solver available to any selector that shows the best aggregate performance on the given set of benchmark instances (here: the lowest PAR10 value). An algorithm selector is only useful if – taking feature costs into account – it performs better than the SBS. In principle, we could determine the SBS on our entire instance set or, for cross-validation, compute the SBS for each fold. On our data, this results in the same SBS.

6 Results

We now present a detailed performance analysis of the algorithm selectors we automatically constructed in our experiments.

6.1 Exploratory Data Analysis

We begin our analysis with some general observations on the feature values and performance data observed on our benchmark set. Table 2 shows how often a solver per-

Table 2: Solver performance in terms of the number of instances that were solved best (relative to all other solvers) or remained unsolved, as well as running times (PAR10 scores, in seconds). The performances of our best two selectors are also shown, and the performance of the SBS (= EAX+restart) is printed in **bold**. The PAR10 scores of the selectors are shown including and excluding the cost of feature computation (top and bottom values, respectively). The PAR10 scores in the last column (marked with *) are those for the VBS. Note that all PAR10 scores within this table are computed without cross-validation, i.e., they are simply averaged across all the instances of the corresponding TSP set(s). Consequently, the performance values slightly differ from the values obtained from 10-fold CV shown in Table 3.

Instance set	Measure	EAX	EAX +rest.	LKH	LKH +rest.	MAOS	Best Selectors #1 #2		Total
RUE	Unique	52	66	223	226	22	92	84	589
	Shared	1	1	10	10	0	3	4	11
	Failed	157	0	18	2	106	0	0	204
	PAR10	9430.92	21.21	1135.67	159.79	6377.80	20.65 20.42	21.10 20.87	10.70*
VLSI	Unique	3	5	4	6	0	4	6	18
	Shared	0	0	0	0	0	0	0	0
	Failed	2	0	2	0	2	0	0	4
	PAR10	4004.18	6.35	4008.29	47.23	4011.35	5.36 5.14	6.51 6.29	4.44*
TSPLIB	Unique	8	2	5	6	1	8	5	22
	Shared	0	0	0	0	0	0	0	0
	Failed	5	1	2	1	2	0	0	6
	PAR10	8185.08	1649.51	3332.99	1679.79	3282.30	13.68 13.46	15.34 15.12	10.76*
National	Unique	3	0	1	1	0	1	0	5
	Shared	0	0	0	0	0	0	0	0
	Failed	0	0	0	0	1	0	0	1
	PAR10	4.92	5.14	29.03	16.53	7209.53	5.25 5.05	5.34 5.14	4.12*
Netgen	Unique	156	216	92	108	20	216	216	592
	Shared	7	7	1	1	0	7	7	8
	Failed	77	0	18	13	92	0	0	149
	PAR10	4627.24	12.99	1190.55	862.97	5532.35	13.22 12.99	13.22 12.99	10.28*
Morphed	Unique	152	229	94	87	27	228	231	589
	Shared	8	7	3	4	0	8	7	11
	Failed	114	0	22	9	93	0	0	178
	PAR10	6846.10	16.72	1444.99	654.08	5593.25	16.93 16.70	16.94 16.71	11.43*
Total	Unique	374	518	419	434	70	549	542	1815
	Shared	16	15	14	15	0	18	18	30
	Failed	355	1	62	25	296	0	0	542
	PAR10	6934.81	36.30	1305.34	565.85	5789.98	16.75 16.52	16.93 16.70	10.73*

formed best on an instance from a specific instance set and also indicates whether multiple algorithms achieved the same best performance. For 75% of the RUE instances, one of the two LKH-versions was best (223 + 226 out of 600), and on more than 60% (753 out of 1 200) of the netgen and morphed instances either EAX or EAX+restart showed the best performance. Also, while EAX and EAX+restart, as well as LKH and LKH+restart, sometimes share the best performance, MAOS was either the unique best solver or fell short of the best performance, i.e., it never jointly ranked first with another solver. Noticeably, MAOS is usually outperformed by EAX+restart, LKH and LKH+restart; nevertheless, there exists a small fraction of instances (70 out of 1 845) in which it performed better than any of the other TSP solvers.

A closer look at the running times required by each solver across our instance sets for finding optimal solutions reveals that, while LKH and LKH+restart clearly performed best on the largest number of RUE instances, their mean PAR10 scores (1 135.67 s and 159.79 s) are substantially worse than those of EAX+restart (21.21 s). The reason for this lies in the fact that LKH and LKH+restart often find an optimal tour quickly, but when they fail to do so, they take quite long, while EAX+restart shows much less extreme performance variation across different instances from the same benchmark set.

Table 2 also clearly indicates that the restart mechanisms integrated into EAX+restart and LKH+restart achieve very substantial performance improvements compared to plain EAX and LKH. Interestingly, EAX benefits much more from restarting than LKH, as plain EAX shows the highest number of unsolved instances (355) and the worst PAR10 score of any of the solvers, whereas EAX+restart is the single best solver and fails to solve only a single instance across all of our benchmark sets. (We

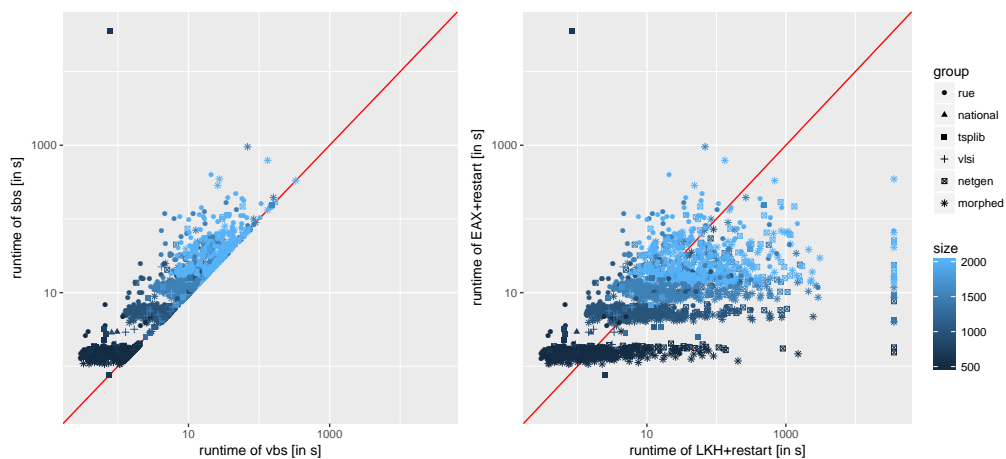


Figure 4: Scatterplots of the aggregated running times, shown on a log-scale, across our 1 845 benchmark instances. Left: Comparison of the virtual best solver (VBS) and the single best solver (SBS), i.e., EAX+restart. Right: Comparison of the two best performing algorithms, LKH+restart and EAX+restart. While there is only one instance (d657 from TSPLIB, shown as a small dark square in the top left) that could not be solved by the SBS, LKH+restart failed to find the optimal tour for 25 instances (squares on the right margin of the plot).

note that neither the benchmark sets nor the cutoff time were specifically chosen to achieve this level of performance.)

For LKH+restart, especially the clustered netgen and morphed instances turned out to be difficult to solve. On these instance sets, either variant of EAX ranked first more often than LKH+restart, and while EAX+restart always found an optimal tour within the given time budget, LKH+restart failed to solve 22 out of the 1 200 instances.

Figure 4 (left) provides a more detailed look at the performance of our two baselines for algorithm selection – the virtual best solver (VBS) and the single best solver, i.e., EAX+restart. As can be seen from this scatterplot, there is a substantial performance gap between the VBS and the SBS across all instance types and sizes, clearly indicating the performance gains that might be achieved using algorithm selection. (Recall that the performance of the VBS provides an upper bound on the performance of any algorithm selector.)

Further evidence for this performance potential comes from the fact that the SBS needs more than one second for all but one instance (p654 from TSPLIB, which is shown on the right of Figure 5), whereas the VBS finds an optimal tour in less than a second for 245 instances.

As can be seen in Figure 4 (right), LKH+restart and EAX+restart show qualitatively different performance characteristics: the running time of EAX+restart varies much less over instances of the same size. This is also reflected in the fact that the Spearman correlation coefficient between instance size and running time for LKH+restart is much lower (0.70) than for EAX+restart (0.91).

Instance d657 from TSPLIB could not be solved by the SBS. Analyzing the performances of all five solvers on that instance, we found that neither EAX, nor EAX+restart

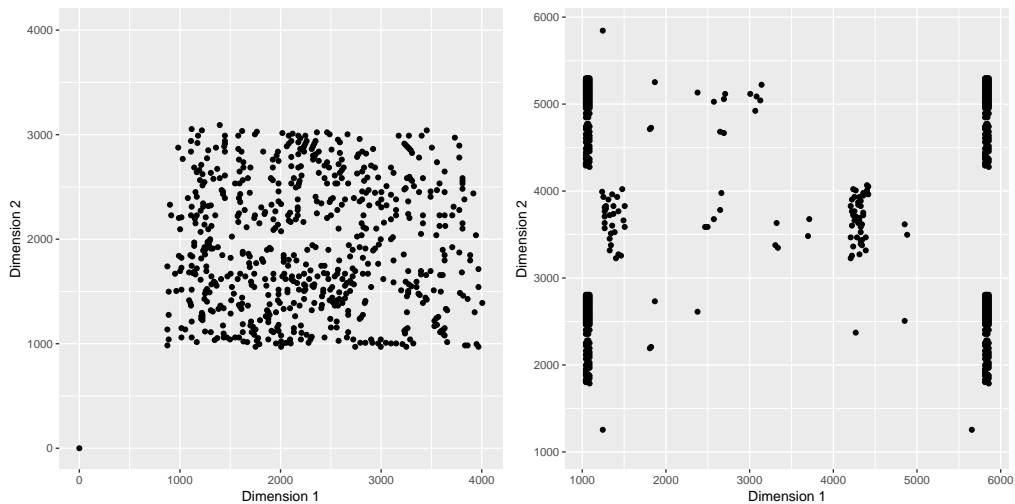


Figure 5: Visualization of two noteworthy TSPLIB instances. Left: d657, which was neither solved by EAX, EAX+restart nor MAOS, whereas LKH and LKH+restart managed to find the optimal tour in less than a second. Right: p654, which was the only instance that EAX+restart solved in less than a second (0.76 s), while LKH (1.76 s), LKH+restart (2.47 s) and MAOS (4.41 s) needed much longer.

or MAOS were able to solve this instance, whereas LKH and LKH+restart succeeded in less than one second. The instance is shown in Figure 5 (left); it resembles an RUE instance within the $[1\,000, 4\,000] \times [1\,000, 3\,000]$ box, with one additional city in the origin, $(0, 0)$. Interestingly, multiple runs of both EAX versions and of MAOS on the original instance failed, while all five solvers quickly succeeded on a customised replicate of that instance, which was created by sampling 656 cities random uniformly within the previously mentioned box-constraints extended by a city located in $(0, 0)$.

On the right side of Figure 5, we show the easiest instance for EAX+restart, p654 from TSPLIB. While it took EAX and EAX+restart between 0.75 and 0.76 s to find an optimal tour for this instance, the other three solvers needed 2.3 to 5.8 times longer. Comparing this instances with d657, as well as the performances of the solvers across our instance sets, there is a clear tendency for EAX+restart to be more effective on structured or clustered data than LKH (with and without restart strategy) or MAOS.

6.2 Comparison of Algorithm Selectors

Table 3 summarises the performance that was achieved by the best five algorithm selectors (according to their PAR10 scores) that we obtained from the experiment outlined in Section 5. All of them perform significantly better than the single best solver (i.e., EAX+restart, with a PAR10 score of 36.3 s), closing most of the gap to the virtual best solver (with a PAR10 score of 10.7 s). We also note that only a small fraction of the available features are used by the best selectors, and that the feature selection step is crucial for obtaining high-performance selectors – none of the 70 selectors that used all features was able to beat the SBS. The best-performing selectors from Table 3 were more than 10 times better (in terms of PAR10) than the same selectors constructed without feature selection.

Our two best selectors are SVMs, which handle the algorithm selection problem as a classification problem. These two selectors are based on 16 and 11 Pihera features, respectively, but only three of the features are identical across both selectors. Other approaches, e.g., a regression-based MARS model, a paired-regression SVM, or a classification tree, also exceed the performance of the single best solver.

Noticeably, features from the TSPmeta and the full UBC feature sets – which on average needed 9 to 12 s longer to compute (per instance) than features from the other two sets – were never selected by the best-performing selectors. Instead, these are either based on the cheap UBC or the Pihera feature set, but none of them uses features from both sets simultaneously, although the feature selection step in the construction of the the second-best selector began with the union of all our feature sets. Overall, our results suggest that all feature sets are similarly informative, and if features from one of those sets are present, adding features from any of the other sets does not improve performance sufficiently to justify the increased cost.

Comparing the PAR10 scores of our selectors and the individual TSP solvers we considered, our two best selectors always perform at least as well as any of the individual solvers (including the SBS) when ignoring the costs for the feature computation. The single best solver was only able to reach the performance of the best selector on the netgen instances, and the performance of the second-best selector on the netgen and National sets – for all other instance sets, our best two best selectors perform better than the SBS. When taking feature costs into account, our best selector beats the SBS on three of our six instance sets, and the second-best selector still beats the SBS on two instance sets.

We note that the SBS, EAX+restart, times out on TSPLIB instance d657. By choos-

Table 3: Overview of the configuration, performance and number of selected features (per feature set) of the five best performing algorithm selectors. All five selectors performed significantly better (p-value $< 10^{-16}$ in a Wilcoxon signed-rank test) than the single best solver (SBS), i.e., EAX+restart, whose PAR10 score was between 36.249 s (without att532) and 36.362 s (with att532). The table also lists how frequently each of the selectors performed better than the SBS, based on a comparison of 10 000 bootstrap samples.

Method	Selector Characteristics			Performance		# Selected Features			
	Learner	Initial Feat. Set	FS-Strat.	PAR10 (in s)	Boot. (in %)	UBC cheap	all	TSP-meta	Pihera
Classif.	SVM	Pihera	sffs	16.747	87.60	0	0	0	16
Classif.	SVM	All	sffs	16.930	65.34	0	0	0	11
Repairs	SVM	UBC (cheap)	sffs	17.712	76.57	13	0	0	0
Regr.	MARS	UBC (cheap)	sffs	19.670	64.34	9	0	0	0
Classif.	RPART	Pihera	sffs	20.799	63.88	0	0	0	5

ing a different solver on that instance, all of our selectors avoid a substantial performance penalty amounting to almost 20 on the PAR10 score on the entire benchmark set. LKH and LKH+restart solve this instance in less than a second. As a result, while the PAR10 score for EAX+restart across the instances from TSPLIB was 1 649.51 s, our best selector achieved a score of 13.68 s.

Nevertheless, even when excluding this instance from the training set, our two best-performing selectors still performed statistically significantly better than the SBS: The PAR10 score of the SBS is 16.8 s, while the retrained versions of the top two selectors from Table 3 achieved PAR10 scores of 16.7 s and 16.4 s, respectively (including feature computation costs); these differences are small, but statistically significant according to a Wilcoxon signed-rank test at $\alpha = 0.05$.

Figure 6 illustrates the behaviour of our best three selectors in more detail. The instance at the top left is d657 and all of our models correctly avoid selecting the SBS, EAX+restart, on this instance. For most instances however, the single best solver is the best choice – most points are very close to the diagonal, indicating that the selectors chose EAX+restart and that the costs for feature computation were negligible. The clouds of points on the upper left side of the diagonal show that especially the two best models improved over the SBS on many other instances as well.

7 Conclusion

In this article, we provided an in-depth study of automatic algorithm selection for the TSP. Specifically, we demonstrated that state-of-the-art inexact TSP solvers show substantial performance complementarity – different solvers perform best on different TSP instances – and that this performance complementarity can be exploited by constructing per-instance algorithm selectors.

Our study presented here expands our own preliminary work (Kotthoff et al., 2015) substantially in scope and depth. Specifically, we considered an additional solver (MAOS), two additional types of TSP instances (netgen and morphed) and an additional feature set (by Pihera and Musliu (2014)). More importantly, for the first time, we made extensive use of feature selection, which lead to substantial performance improvements in the selectors we were able to construct, as demonstrated using rigorous

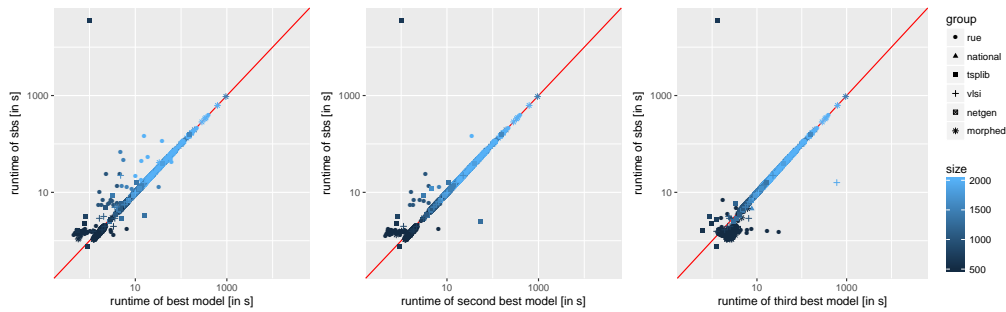


Figure 6: Scatterplots of the running times, shown on a log-scale, across 1 844 benchmark problems (`att532` was excluded). Comparison of the the single best solver (SBS), i.e., EAX+restart, and the best (left), second-best (middle), and third-best selector (right), respectively.

statistical tests. In addition to producing better TSP solver selectors, and hence an improvement in the state of the art in inexact TSP solving, we have also provided, for the first time, a detailed performance analysis of the best-performing inexact TSP solvers known previously.

There are several directions for future work which we believe may lead to even better algorithm selectors. In our earlier work, we explored the use of so-called probing features, which are derived from partial runs of a TSP solver (Kotthoff et al., 2015). While we did not require such features here to produce very effective algorithm selectors, their use might lead to further performance improvements. Likewise, hyperparameter optimisation of the machine learning procedures at the heart of our selectors, though costly, might prove beneficial. Finally, based on our results presented here, we believe that a hybrid TSP solver that initially tries to find an optimal tour with a very fast solver (such as LKH) and then switches to EAX+restart might hold significant promise.

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Appendix

While recursive partitioning and regression trees, random forests and kernel-based support vector machines are all popular machine learning approaches, *multivariate adaptive regression splines* (MARS, Friedman (1991)) are less prominent. We still considered MARS, because we observed promising performance in preliminary experiments; furthermore, the underlying idea is simple and appealing.

A MARS model is a combination of piecewise polynomial functions, so-called *splines*. More formally, the model can be written as a function

$$\hat{f}(\mathbf{z}) = c_0 + \sum_{k=1}^K c_k \cdot b_k(\mathbf{z})$$

with model coefficients c_0, \dots, c_K , basis functions $b_1(\mathbf{z}), \dots, b_K(\mathbf{z})$ and a p -dimensional input vector $\mathbf{z} = (z_1, \dots, z_p)^T$. The basis functions b_k , for $k = 1, \dots, K$, stem from a set

$$\mathcal{H} = \{(z_j - x_{1j})_+^q, (x_{1j} - z_j)_+^q, \dots, (z_j - x_{Nj})_+^q, (x_{Nj} - z_j)_+^q\}_{j=1, \dots, p}$$

of so-called *hinge functions*. Each of them is a polynomial (of order q) of the difference between the j -th input variable z_j and a realization of this variable from the training data, x_{1j}, \dots, x_{Nj} , the so-called *knots*. Furthermore, the hinge function limits the polynomial to non-negative values, i.e., $(z_j - x_{ij})_+^q := \max\{0, (z_j - x_{ij})^q\}$.