

Beginner's Guide to the MOEA Framework

David Hadka

For version 2.12 and later.

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Thank you for purchasing this book! All revenue received from book sales helps fund the continued development and maintenance of the MOEA Framework. We sincerely appreciate your support and hope you find this software useful in your endeavors.

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Chapter 1

Background

Optimization is the process of identifying the best solution among a set of alternatives (Miettinen, 1999). Whereas single objective optimization employs a single criterion for identifying the best solution among a set of alternatives, multiobjective optimization employs two or more criteria. The criteria used to compare solutions are known as *objectives*. As multiple objectives can conflict with one another — i.e., improving one objective leads to the deterioration of another — there is, generally speaking, no single optimal solution to multiobjective problems.

As an example, Figure 1.1 shows the tradeoff between two objectives: (1) cost and (2) error. The shaded region depicts the set of candidate solutions to this hypothetical problem. The top-left region contains low cost, high error candidate solutions. The bottom-right region contains high cost, low error candidate solutions. Between these two extremes lie the various degrees of tradeoff between the two objectives, where increases in cost lead to reduced error.

Figure 1.1 demonstrates a fundamental issue in multiobjective optimization. Given that there is no single optimal solution, rather a multitude of potential solutions with varying degrees of tradeoff between the objectives, decision-makers are subsequently responsible for exploring this set of potential solutions and identifying the solution(s) to be implemented. While ultimately the selection of the final solution is the responsibility of the decision-maker, optimization tools should assist this decision process to the best of their ability. For instance, it may prove useful to identify points of diminishing returns. For example, Figure 1.2 identifies the region where a large increase in cost is necessary to impart a marginal decrease in error. To perform this type of analysis, it is necessary to provide the decision-maker with an enumeration or approximation of these tradeoffs. This strategy of enumerating or approximating the tradeoffs is known as a posteriori optimization (Coello Coello et al., 2007), and is the focus of this book.

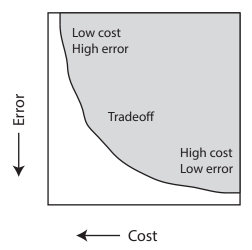


Figure 1.1: Example of the tradeoff between two objectives: (1) cost and (2) error. A tradeoff is formed between these two conflicting objectives where increases in cost lead to reduced error. All figures in this dissertation showing objectives include arrows pointing towards the ideal optimum.

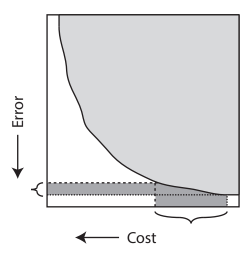


Figure 1.2: Example showing the effect of diminishing returns, where a large increase in cost is necessary to impart a marginal reduction in error.

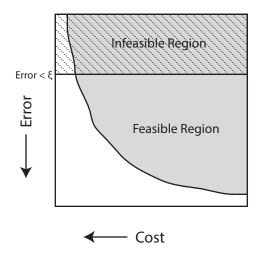


Figure 1.3: Example showing how constraints define an infeasible region (the hashed region). Valid solutions to the optimization problem are only found in the feasible region.

1.1 Multiobjective Problem

We can express the idea of a multiobjective problem (MOP) with M objectives formally as:

minimize
$$F(\mathbf{x}) = (f_1(\mathbf{x}), f_2(\mathbf{x}), \dots, f_M(\mathbf{x}))$$

subject to $c_i(\mathbf{x}) = 0, \ \forall i \in \mathcal{E},$
 $c_i(\mathbf{x}) \le 0, \ \forall j \in \mathcal{I}.$ (1.1)

We call \mathbf{x} the *decision variables*, which is the vector of variables that are manipulated during the optimization process:

$$\mathbf{x} = \begin{bmatrix} x_1 \\ x_2 \\ \vdots \\ x_L \end{bmatrix} \tag{1.2}$$

Decision variables can be defined in a variety of ways, but it is common to see the following types (Bäck et al., 1997):

• Real-Valued: 2.71

• Binary: 001100010010100001011110101101110011

• **Permutation:** 4,2,0,1,3

In some applications, it is possible for the number of decision variables, L, to not be a fixed value. In this book, however, we assume that L is constant for a given problem.

The decision space, Ω , is the set of all decision variables. The MOP may impose restrictions on the decision space, called *constraints*. As an example, in Figure 1.3, a hypothetical

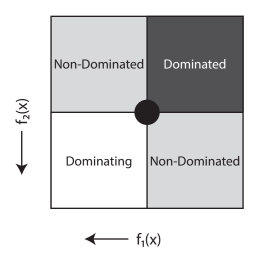


Figure 1.4: Depiction of the various Pareto dominance regions. These regions are relative to each solution, which is centered in the figure. The *dominated* region is inferior in all objectives, the *dominating* region is superior in all objectives and the *non-dominated* region is superior in one objective but inferior in the other.

constraint would prevent any solutions from exceeding an error threshold. In this manner, constraints inform the optimization process as to which solutions are infeasible or impractical. Equation (1.1) shows that zero or more constraints $c_i(\mathbf{x})$ can be defined to express both equality and inequality constraints. The sets \mathcal{E} and \mathcal{I} define whether the constraint is an equality or inequality constraint. The set of all decision variables in Ω which are feasible (i.e., satisfy all constraints) define the feasible region, Λ .

1.2 Pareto Optimality

The notion of optimality used today is adopted from the work of Francis Ysidro Edgeworth and Vilfredo Pareto (Coello Coello et al., 2007), and is commonly referred to as *Pareto optimality*. Pareto optimality considers solutions to be superior or inferior to another solution only when it is superior in all objectives or inferior in all objectives, respectively. The tradeoffs in an MOP are captured by solutions which are superior in some objectives but inferior in others. Such pairs of solutions which are both superior and inferior with respect to certain objectives are called *non-dominated*, as shown in Figure 1.4. More formally, the notion of Pareto optimality is defined by the Pareto dominance relation:

Definition 1 A vector $\mathbf{u} = (u_1, u_2, \dots, u_M)$ **Pareto dominates** another vector $\mathbf{v} = (v_1, v_2, \dots, v_M)$ if and only if $\forall i \in \{1, 2, \dots, M\}, u_i \leq v_i$ and $\exists j \in \{1, 2, \dots, M\}, u_j < v_j$. This is denoted by $\mathbf{u} \prec \mathbf{v}$.

Two solutions are non-dominated if neither Pareto dominates the other (i.e., $\mathbf{u} \not\prec \mathbf{v}$ and $\mathbf{v} \not\prec \mathbf{u}$). The set of all non-dominated solutions is captured by the Pareto optimal set and

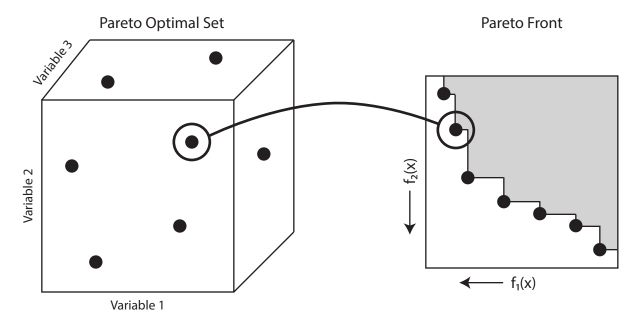


Figure 1.5: Shows a hypothetical mapping between a 3-dimensional Pareto optimal set and its associated 2-dimensional Pareto front. The shaded region in the Pareto front shows the space dominated by the Pareto front.

the Pareto front. The former contains the decision variables while the latter contains the objectives.

Definition 2 For a given multiobjective problem, the **Pareto optimal set** is defined by

$$\mathcal{P}^* = \{ \mathbf{x} \in \Omega \mid \neg \exists \mathbf{x}' \in \Lambda, F(\mathbf{x}') \prec F(\mathbf{x}) \}$$

Definition 3 For a given multiobjective problem with Pareto optimal set \mathcal{P}^* , the **Pareto** front is defined by

$$\mathcal{PF}^* = \{ F(\mathbf{x}) \mid \mathbf{x} \in \mathcal{P}^* \}$$

In this dissertation, the Pareto dominance relation is applied to the objectives. For convenience, we use $\mathbf{x} \prec \mathbf{y}$ interchangeably with $F(\mathbf{x}) \prec F(\mathbf{y})$.

Figure 1.5 shows an example Pareto optimal set and Pareto front, and the resulting mapping between the two. The Pareto optimal set defines the decision variables, whereas the Pareto front captures the objectives and their tradeoffs via Pareto optimality.

1.3 Multiobjective Evolutionary Algorithms

Evolutionary algorithms (EAs) are a class of search and optimization algorithms inspired by processes of natural evolution (Holland, 1975). A broad overview of the design and development of EAs is provided in Bäck et al. (1997). The outline of a simple EA is shown

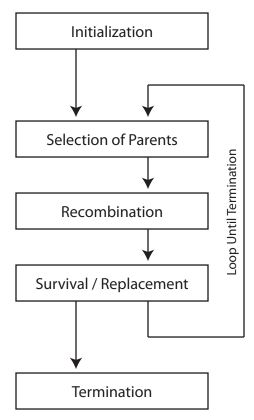


Figure 1.6: The outline of a simple EA. EAs begin with an initialization process, where the initial search population is generated. They next enter a loop of selecting parent individuals from the search population, applying a recombination operator (such as crossover and mutation in genetic algorithms) to generate offspring, and finally updating the search population with these offspring using a replacement strategy. This loop is repeated until some termination condition is met, usually after a fixed number of objective function evaluations (NFE). Upon termination, the EA reports the set of optimal solutions discovered during search.

in Figure 1.6. EAs begin with an initialization process, where the initial search population is generated. They next enter a loop of selecting parent individuals from the search population, applying a recombination operator to generate offspring, and finally updating the search population with these offspring using a replacement strategy. This loop is repeated until some termination condition is met, usually after a fixed number of objective function evaluations (NFE). Upon termination, the EA reports the set of optimal solutions discovered during search.

The behavior of the selection, recombination and survival/replacement processes typically depend on the "class" of EA. For instance, genetic algorithms (GAs) apply crossover and mutation operators that mimic genetic reproduction via DNA (Holland, 1975). Particle swarm optimization (PSO) algorithms simulate flocking behavior, where the direction of travel of each individual is steered towards the direction of travel of surrounding individuals (Kennedy and Eberhart, 1995). While the behavior of each class may be vastly different, they all share a common attribute of utilizing a search population.

Their ability to maintain a population of diverse solutions makes EAs a natural choice for solving MOPs. Early attempts at solving MOPs involved using aggregation-based approaches (Bäck et al., 1997). In aggregation-based approaches, the decision-maker defines an aggregate fitness function that transforms the MOP into a single objective problem, which can subsequently be solved with an EA. Two commonly-used aggregate fitness functions are linear weighting:

$$F_L(\mathbf{x}) = \sum_{i=1}^{M} \lambda_i f_i(\mathbf{x}), \tag{1.3}$$

and the weighted Chebyshev method:

$$F_T(\mathbf{x}) = \max_{i=1,2,\dots,M} \left(\lambda_i \left| z_i^* - f_i(\mathbf{x}) \right| \right), \tag{1.4}$$

where $\lambda = (\lambda_1, \lambda_2, \dots, \lambda_M)$ are the weights and $\mathbf{z}^* = (z_1^*, z_2^*, \dots, z_M^*)$ is a reference point identifying the decision-maker's goal (note: this reference point need not be a feasible solution).

Coello Coello et al. (2007) discusses the advantages and disadvantages of aggregate fitness approaches. The primary advantage is the simplicity of the approach and the ability to exploit existing EAs to solve MOPs. In addition, appropriately defined aggregate fitness functions can provide very good approximations of the Pareto front. However, poorly-weighted aggregate fitness functions may be unable to find non-convex regions of the Pareto front. This is problematic since selecting appropriate weights is non-trivial, especially if the relative worth of each objective is unknown or difficult to quantify. Lastly, in order to generate multiple Pareto optimal solutions, aggregate fitness approaches need to be run with differing weights to generate solutions across the entire Pareto front.

These limitations lead to the development of multiobjective evolutionary algorithms (MOEAs) that search for multiple Pareto optimal solutions in a single run. The first MOEA to search for multiple Pareto optimal solutions, the Vector Evaluated Genetic Algorithm (VEGA), was introduced by Schaffer (1984). VEGA was found to have problems similar

to aggregation-based approaches, such as an inability to generate concave regions of the Pareto front. Goldberg (1989) was first to suggest the use of Pareto-based selection, but this concept was not applied until 1993 in the Multiobjective Genetic Algorithm (MOGA) (Fonseca and Fleming, 1993). Between 1993 and 2003, several first-generation MOEAs were introduced demonstrating important design concepts such as elitism, diversity maintenance and external archiving. Notable first-generation algorithms include the Niched-Pareto Genetic Algorithm (NPGA) (Horn and Nafpliotis, 1993), the Non-dominated Sorting Genetic Algorithm (NSGA) (Srinivas and Deb, 1994), the Strength Pareto Evolutionary Algorithm (SPEA) (Zitzler and Thiele, 1999), the Pareto-Envelope based Selection Algorithm (PESA) (Corne and Knowles, 2000) and the Pareto Archived Evolution Strategy (PAES) (Knowles and Corne, 1999). Many of these MOEAs have since been revised to incorporate more efficient algorithms and improved design concepts. To date, Pareto-based approaches outnumber aggregate fitness approaches (Coello Coello et al., 2007). For a more comprehensive overview of the historical development of MOEAs, please refer to the text by Coello Coello et al. (2007).

1.4 Measuring Quality

When running MOEAs on a MOP, the MOEA outputs an approximation of the Pareto optimal set and Pareto front. The approximation of the Pareto front, called the *approximation set*, can be used to measure the quality of an MOEA on a particular problem. In some situations, such as with contrived test problems, a *reference set* of the globally optimal solutions may be known. If known, the reference set can be used to measure the absolute performance of an MOEA. If not known, the approximation sets from multiple MOEAs can be compared to determine their relative quality.

There is no consensus in the literature of the appropriate procedure with which to compare approximation sets. These procedures, called *performance metrics*, come in two forms: (1) unary and (2) binary performance metrics (Zitzler et al., 2002c). Unary performance metrics produce a single numeric value with which to compare approximation sets. Unary performance metrics have the advantage of permitting the comparison of approximation sets without requiring the actual approximation set, as one need only compare the numeric values. Binary performance metrics, on the other hand, compare pairs of approximation sets, identifying which of the two approximation sets is superior. In order to allow comparisons across studies, this book uses only unary performance metrics.

Zitzler et al. (2002b) contend that the number of unary performance metrics required to determine if one approximation set is preferred over another must be at least the number of objectives in the problem. Because different MOEAs tend to perform better in different metrics (Bosman and Thierens, 2003), Deb and Jain (2002) suggest only using metrics for the two main functional objectives of MOEAs: proximity and diversity. The following outlines several of the commonly-used unary performance metrics. For details of these performance metrics see Coello Coello et al. (2007).

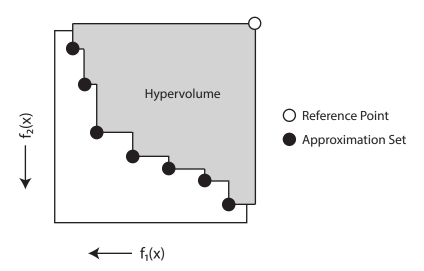


Figure 1.7: Hypervolume measures the volume of the space dominated by the approximation set, bounded by a reference point. This reference point is typically the nadir point (i.e., the worst-case value for each objective) of the reference set plus some fixed delta. This delta ensures extremal points contribute non-zero hypervolume.

Hypervolume As shown in Figure 1.7, the hypervolume metric computes the volume of the space dominated by the approximation set. This volume is bounded by a reference point, which is usually set by finding the nadir point (i.e., the worst-case objective value for each objective) of the reference set plus some fixed increment. This fixed increment is necessary to allow the extremal points in the approximation set to contribute to the hypervolume. Knowles and Corne (2002) suggest the hypervolume metric because it is compatible with the outperformance relations, scale independent, intuitive, and can reflect the degree of outperformance between two approximation sets.

The major disadvantage of the hypervolume metric is its runtime complexity of $O(n^{M-1})$, where n is the size of the non-dominated set. However, Beume and Rudolph (2006) provide an implementation with runtime $O(n \log n + n^{M/2})$ based on the Klee's measure algorithm by Overmars and Yap. This implementation permits computing the hypervolume metric on moderately sized non-dominated sets up to M = 8 objectives in a reasonable amount of time. Further improvements by While et al. (2012) improve the expected runtime further, allowing the efficient calculation of hypervolume with ten or more objectives.

Generational Distance Generational distance (GD) is the average distance from every solution in the approximation set to the nearest solution in the reference set, as shown in Figure 1.8. As such, it measures proximity to the reference set. GD by itself can be misleading, as an approximation set containing a single solution in close proximity to the reference set produces low GD measurements, and is often combined with diversity measures in practice (Hadka and Reed, 2012).

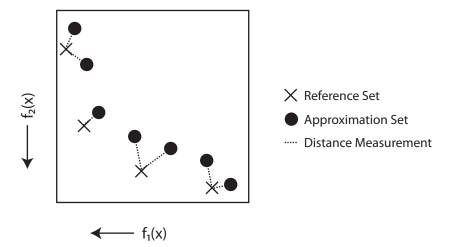


Figure 1.8: Generational distance is the average distance from every solution in the approximation set to the nearest solution in the reference set.

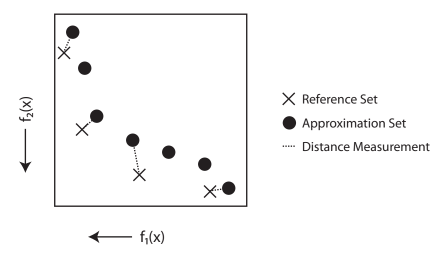


Figure 1.9: Inverted generational distance is the average distance from every solution in the reference set to the nearest solution in the approximation set.

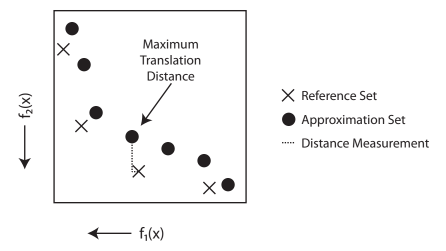


Figure 1.10: ϵ_+ -indicator (also known as the additive ϵ -indicator) is the smallest distance ϵ that the approximation set must be translated by in order to completely dominate the reference set (Coello Coello et al., 2007).

Inverted Generational Distance As its name indicates, the inverted generational distance (IGD) is the inverse of GD — it is the average distance from every solution in the reference set to the nearest solution in the approximation set. IGD measures diversity, as shown in Figure 1.9, since an approximation set is required to have solutions near each reference set point in order to achieve low IGD measurements (Coello Coello et al., 2007).

 ϵ_+ -Indicator The additive ϵ -indicator (ϵ_+ -indicator) measures the smallest distance ϵ that the approximation set must be translated by in order to completely dominate the reference set, as shown in Figure 1.10. One observes that good proximity and good diversity both result in low ϵ values, as the distance that the approximation needs to be translated is reduced. However, if there is a region of the reference set that is poorly approximated by the solutions in the approximation set, a large ϵ is required. Therefore, we claim the ϵ_+ -indicator measures the *consistency* of an approximation set (Hadka and Reed, 2013). An approximation set must be free from large gaps or regions of poor approximation in order to be consistent.

Spacing Spacing, shown in Figure 1.11, measures the uniformity of the spacing between solutions in an approximation set (Coello Coello et al., 2007). An approximation set that is well-spaced will not contain dense clusters of solutions separated by large empty expanses. Note that, since spacing does not involve a reference set in its calculation, an approximation can register good spacing while having poor proximity to the reference set. It is therefore recommended to use spacing in conjunction with a performance metric for proximity.

In academic works, it is common to see results published using GD, hypervolume and ϵ_{+} -indicator. These three metrics record proximity, diversity and consistency, respectively,

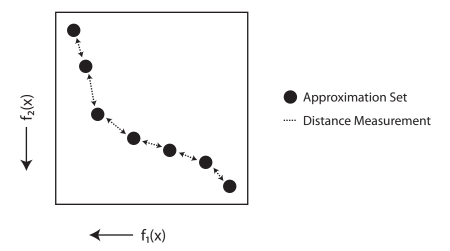


Figure 1.11: Spacing measures the uniformity of the spacing between solutions in an approximation set.

which we claim are the three main functional objectives of MOEAs (Fonseca and Fleming, 1996). Figure 1.12 provides a graphical representation of the importance of the ϵ_+ -indicator and consistency. MOEAs are expected to produce high-quality solutions covering the entire extent of the tradeoff surface, with few gaps or regions of poor approximation.

In order to report these performance metrics consistently, all performance metrics are normalized. This normalization converts all performance metrics to reside in the range [0,1], with 1 representing the optimal value. First, the reference set is normalized by its minimum and maximum bounds so that all points in the reference set lie in $[0,1]^N$, the N-dimensional unit hypercube. Second, each approximation set is normalized using the same bounds. Third, the performance metrics are calculated using these normalized sets. Finally, the performance metrics are transformed by the following equations to ensure a value of 1 represents the optimal value achievable by the metric. Hypervolume is transformed with:

$$\mathcal{M}(A_p^s) = \widehat{\mathcal{M}}(A_p^s) / \mathcal{M}^*, \tag{1.5}$$

where $\widehat{\mathcal{M}}$ represents the raw metric value. GD and the ϵ_+ -indicator are transformed with:

$$\mathcal{M}(A_p^s) = \max(1 - \widehat{\mathcal{M}}(A_p^s), 0). \tag{1.6}$$

When solving test problems, the reference set is known analytically. For most real-world problems, however, the reference set is not available. In these situations, it is often necessary to construct a reference set from the union of all approximation sets generated during experimentation. Then, performance metrics can be evaluated relative to this combined reference set.

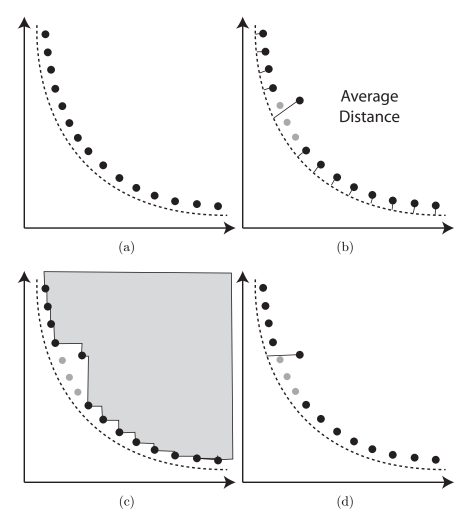


Figure 1.12: Demonstrates the importance of ϵ -indicator as a measure of consistency. (a) A good approximation set to the reference set, indicated by the dashed line. (b) Generational distance averages the distance between the approximation set and reference set, reducing the impact of large gaps. The missing points are shaded light gray. (c) The change in hypervolume due to a gap is small relative to the entire hypervolume. (d) ϵ -Indicator easily identifies the gap, reporting a metric 2-3 times worse in this example.

1.5 The MOEA Framework

The MOEA Framework is a free and open source Java library for developing and experimenting with multiobjective evolutionary algorithms (MOEAs) and other general-purpose optimization algorithms. We will be using the MOEA Framework throughout this book to explore multiobjective optimization. Its key features includes:

Fast, reliable implementations of many state-of-the-art multiobjective evolutionary algorithms. The MOEA Framework contains internally NSGA-II, NSGA-III, ϵ -MOEA, ϵ -NSGA-II, PAES, PESA2, SPEA2, IBEA, SMS-EMOA, GDE3, SMPSO, OMOPSO, CMA-ES, and MOEA/D. These algorithms are optimized for performance, making them readily available for high performance applications. By also supporting the JMetal and PISA libraries, the MOEA Framework provides access to 30 multiobjective optimization algorithms.

Extensible with custom algorithms, problems and operators. The MOEA Framework provides a base set of algorithms, test problems and search operators, but can also be easily extended to include additional components. Using a Service Provider Interface (SPI), new algorithms and problems are seamlessly integrated within the MOEA Framework.

Modular design for constructing new optimization algorithms from existing components. The well-structured, object-oriented design of the MOEA Framework library allows combining existing components to construct new optimization algorithms. And if needed functionality is not available in the MOEA Framework, you can always extend an existing class or add new classes to support any desired feature.

Permissive open source license. The MOEA Framework is licensed under the free and open GNU Lesser General Public License, version 3 or (at your option) any later version. This allows end users to study, modify, and distribute the MOEA Framework freely.

Fully documented source code. The source code is fully documented and is frequently updated to remain consistent with any changes. Furthermore, an extensive user manual is provided detailing the use of the MOEA Framework in detail.

Extensive support available online. As an actively maintained project, bug fixes and new features are constantly added. We are constantly striving to improve this product. To aid this process, our website provides the tools to report bugs, request new features, or get answers to your questions.

Over 1200 test cases to ensure validity. Every release of the MOEA Framework undergoes extensive testing and quality control checks. And, if any bugs are discovered that survive this testing, we will promptly fix the issues and release patches.

1.6 Getting Help

This beginner's guide is the most comprehensive resource for learning about the MOEA Framework. Additional resources are available on our website at http://www.moeaframework.org. This website also has links to file bugs or request new features. If you still can not find an answer to your question, feel free to contact us at support@moeaframework.org.

Chapter 2

Setup and First Example

In this chapter, we will setup the MOEA Framework on your computer and demonstrate a simple example. These instructions are tailored for Windows users, but similar steps can be taken to install the MOEA Framework on Linux or Mac OS.

2.1 Installing Java

The MOEA Framework runs on Java version 6 or any later version. Since we will need to compile examples, you will need to install the Java Development Kit (JDK) for version 6 or later. For Windows users, we recommend using Oracle's JDK available at http://www.oracle.com/technetwork/java/javase/. Make sure you download the JDK and not the JRE (Java Runtime Environment).

2.2 Installing Eclipse

If this is your first time using the MOEA Framework, we suggest using Eclipse to build projects. Eclipse is a free development environment for writing, debugging, testing, and running Java programs. First, download Eclipse from http://www.eclipse.org/. To install Eclipse, simply extract the ZIP archive to a location on your computer (e.g., your desktop). Within the extracted folder, run eclipse.exe. First-time users of Eclipse may be prompted to select a workspace location. The default location is typically fine. Click the checkbox to no longer show this dialog and click Ok.

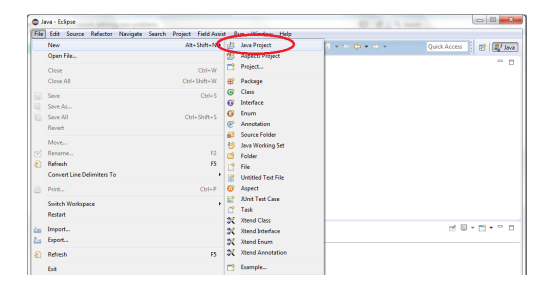
2.3 Setting up the MOEA Framework

We recommend starting with this book's supplemental materials, which includes a full installation of the MOEA Framework and all of the code samples found in this book. The supplemental materials can be downloaded by following this link: http://bit.ly/1N5sHjO.

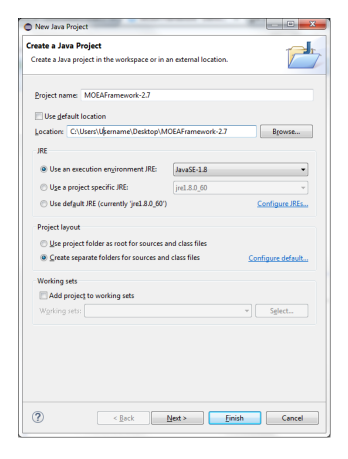
Note: that ends with the letter O and not the number 0. As you read this book, find the appropriate Java file within the book folder to follow along.

Alternatively, you can download the MOEA Framework's compiled binaries from http://www.moeaframework.org/ and manually type in the examples. The compiled binaries are distributed as a compressed TAR file (.tar.gz) and need to be extracted. We recommend using 7-Zip, a free and open source program, which can be downloaded from http://www.7-zip.org/. Extract to your Desktop or any other convenient location.

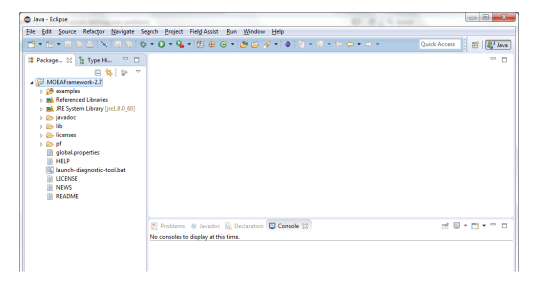
Next, we need to create a project within Eclipse. Select File \to New Java Project from the menu.



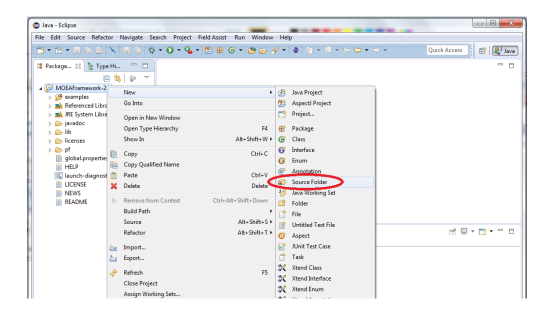
In the window that appears, uncheck "Use default location". Click the "Browse..." button and select the extracted MOEA Framework folder. Click Finish.



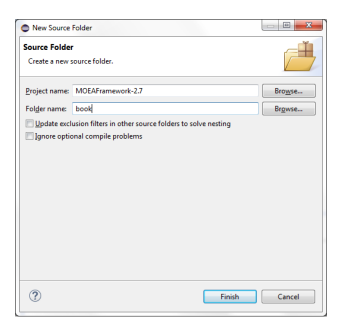
The MOEA Framework project will now appear within the "Package Explorer" in Eclipse, as shown below.



If you are using the supplemental materials, you can skip down to the next section titled "Your First Example." Otherwise, we will now create a source folder were our examples will reside. Right-click on the project and select New \rightarrow Source Folder.

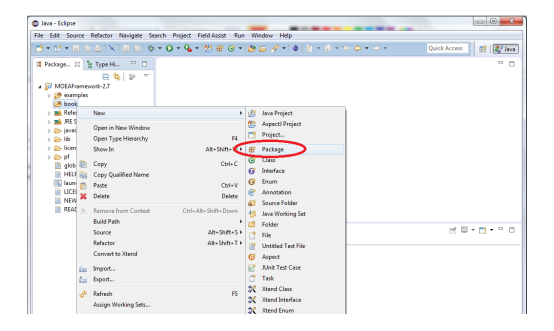


Give the new source folder the name "book" and click Finish.

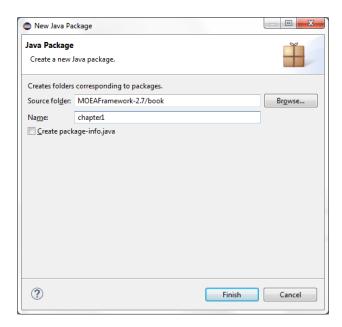


2.4 Your First Example

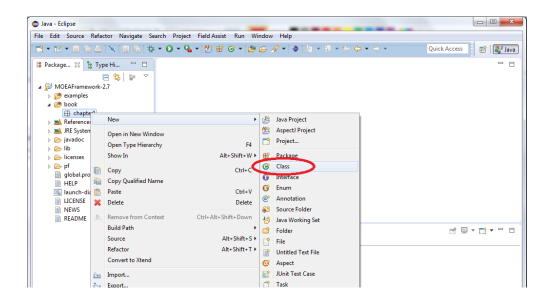
In Java, packages are used to organize source code into a hierarchical structure. We will organize the examples from each chapter into its own package. Thus, for the first chapter, we will create a package named chapter 2. Right-click on the source folder we just created and select New \rightarrow Package.



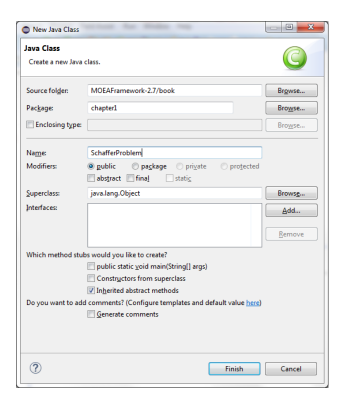
Enter the name chapter2 and click Finish.



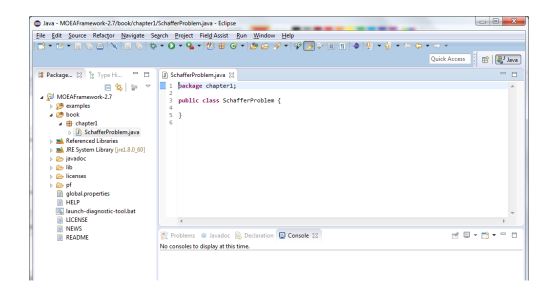
Finally, we create the Java file for the actual code. In Java, these are called classes. Right-click the chapter 2 folder and select New \rightarrow Class.



Type the name SchafferProblem and click Finish.



At this point, your Eclipse workspace will contain one Java file named SchafferProblem.java and that file will be opened in the text editor within Eclipse, as shown below.



Now we can begin defining the problem.

```
package chapter2;
  import org.moeaframework.core.Solution;
  import org.moeaframework.core.variable.EncodingUtils;
  import org.moeaframework.problem.AbstractProblem;
  public class SchafferProblem extends AbstractProblem {
7
8
    public SchafferProblem() {
9
10
       super(1, 2);
    @Override
13
    public void evaluate(Solution solution) {
14
      double x = EncodingUtils.getReal(solution.getVariable(0));
16
      solution.setObjective(0, Math.pow(x, 2.0));
17
      solution.setObjective(1, Math.pow(x - 2.0, 2.0));
18
19
    }
20
    @Override
    public Solution newSolution() {
22
      Solution solution = new Solution(1, 2);
23
      solution.setVariable(0, EncodingUtils.newReal(-10.0, 10.0));
24
25
      return solution;
26
27
```

MOEAFramework/book/chapter2/SchafferProblem.java

The anatomy of a problem is as follows. First, it must implement the Problem interface. Rather that implement the Problem interface directly, it is often more convenient to extend the AbstractProblem class, as seen on Line 7. Three methods are required when extending AbstractProblem: the constructor, newSolution, and evaluate. The constructor, shown on lines 9-11, is responsible for initializing the problem. For this problem, we call **super** (1, 2) to indicate this problem will consist of one decision variable and two objectives. The newSolution method, shown on lines 14-18, generates a prototype solution for the problem. A prototype solution describes each decision variable, and were applicable, any bounds on the variables. For this problem, we create a single real-valued decision variable bounded by [-10, 10]. Thus, on line 15 we create the prototype solution with one variable and two objectives (e.g., **new** Solution (1, 2)), assign the variable on line 16 (e.g., solution.setVariable(0, EncodingUtils.newReal(-10.0, 10.0); and return the solution on line 17. Finally, we define the evaluate method on lines 21-26, which is responsible for computing the objectives for a given candidate solution. On line 22, we read the value of the decision variable (e.g., EncodingUtils.getReal(solution.getVariable(0)), and on lines 24 and 25 evaluate the two objectives. For the Schaffer problem, the two objectives are $f_1(x) = x^2$ and $f_2(x) = (x-2)^2$. Type this code into the SchafferProblem. java file and save.

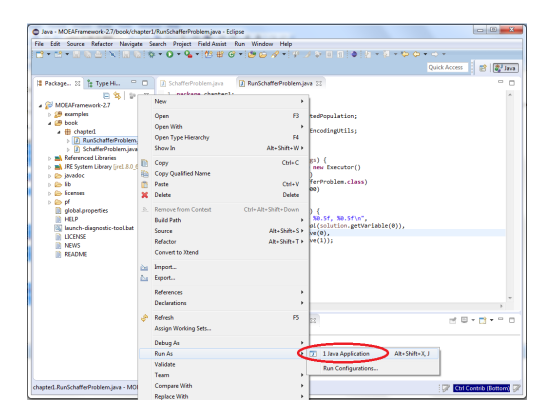
At this point, the problem is defined, but we also need to create the code to solve the problem. To begin, create a new class called RunSchafferProblem with the following code:

```
package chapter2;
2
  import org.moeaframework.Executor;
  import org.moeaframework.core.NondominatedPopulation;
  import org.moeaframework.core.Solution;
  import org.moeaframework.core.variable.EncodingUtils;
6
7
  public class RunSchafferProblem {
8
9
    public static void main(String[] args) {
10
11
      NondominatedPopulation result = new Executor()
           .withAlgorithm("NSGAII")
12
           .withProblemClass(SchafferProblem.class)
           .withMaxEvaluations(10000)
14
           .run();
16
       for (Solution solution : result) {
17
         System.out.printf("%.5f \Rightarrow %.5f, %.5f\n",
18
             EncodingUtils.getReal(solution.getVariable(0)),
19
             solution.getObjective(0),
20
             solution.getObjective(1));
21
22
    }
23
24
25 }
```

MOEAFramework/book/chapter2/RunSchafferProblem.java

On line 10, we create a the main method. In Java, main methods are the starting points for applications. This is the first method that is invoked when we run the application. To solve our problem, we will use the Executor class. The executor is responsible for creating instances of optimization algorithms and using them to solve problems. It is a sophisticated class, but at the bare minimum it requires three pieces of information: 1) the name of the optimization algorithm, 2) the problem, and 3) the maximum number of function evaluations (NFE) permitted to solve the problem. We set the values on lines 12-14 and call run() on line 15 to solve the problem. The result, a Pareto approximation set, is saved on line 10 to a NondominatedPopulation. Lines 17-22 format and print the Pareto approximate set to the screen. Each line on the output is a Pareto approximate solution where the left-most value is the decision variable and the two values to the right of => are the objective values.

Run this application by right-clicking the file \blacksquare RunSchafferProblem.java and selecting Run \rightarrow Java Application.



The output will appear in the Console within Eclipse and should appear similar to below.

Congratulations, you have just successfully optimized a problem using the MOEA Framework!

2.5 Running from Command Line

If you are not using Eclipse to run these examples, they can also be run manually from the command line. On Windows, open a new Command Prompt window and change the directory to the MOEA Framework folder. Then type the following commands:

```
javac -cp "lib/*;book" book/chapter2/SchafferProblem.java
javac -cp "lib/*;book" book/chapter2/RunSchafferProblem.java
java -cp "lib/*;book" chapter2.RunSchafferProblem
```

The first two lines compile the two class files we created. Note the use of the -cp "lib/*;book" argument that specifies the Java classpath. This tells Java where to locate any referenced files. We will be referencing files in the lib folder, which contains all of the MOEA Framework libraries, and the book folder, which contains the files we are compiling. The last line runs the example. We again must specify the classpath, but note that we are running the class chapter2.RunShafferProblem. This is the full class path for our problem. It consists of the package (e.g., chapter2), followed by a period, followed by the class name (e.g., RunSchafferProblem).

2.6 Plotting Results

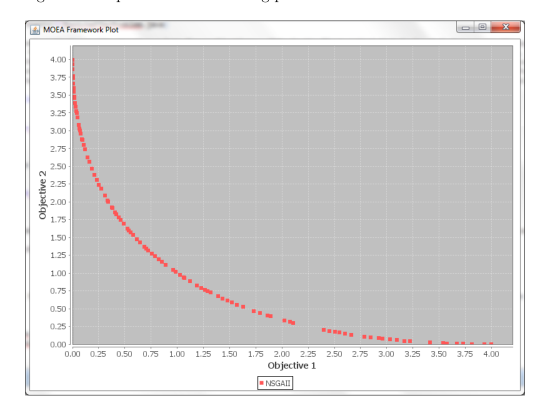
In the previous example, we output the two objectives to the console. Outputting the raw data like this is useful as you can save the data to a text file, load the data into Excel, etc. For problems like this with only two objectives, we can plot the solutions as points in a scatter plot, as shown below:

```
package chapter2;

import org.moeaframework.Executor;
import org.moeaframework.analysis.plot.Plot;
import org.moeaframework.core.NondominatedPopulation;
```

```
public class PlotSchafferProblem {
7
8
    public static void main(String[] args) {
9
       NondominatedPopulation result = new Executor()
10
           .withAlgorithm("NSGAII")
           .withProblemClass(SchafferProblem.class)
12
13
           .withMaxEvaluations (10000)
           .run();
14
15
       new Plot()
16
           .add("NSGAII", result)
17
           .show();
19
20
21
```

 $MOEAF ramework/book/chapter 2/PlotSchaffer Problem. java \\ Running this code produces the following plot:$



Now we can see the shape of the Pareto approximation set produced by the optimization algorithm.

Chapter 3

Constrained Optimization

In the previous chapter, we solved the two objective Schaffer problem. This was an unconstrained problem, meaning that any solution we generate is a feasible design. Many real-world problems have constraints, either caused by physical limitations (e.g., maximum operating temperature), monetary (e.g., available capital), are risk-based (e.g., maximum failure rate for a product), etc. In general, there are two types of constraints: equality and inequality. Equality constraints are of the form g(x) = c for some constant c. Inequality constraints are of the form $h(x) \geq d$ for some constant c. For example, takes the Srinivas problem as defined below.

$$F = (f_1(x, y), f_2(x, y)), \text{ where}$$

$$f_1(x, y) = (x - 2)^2 + (y - 1)^2 + 2,$$

$$f_2(x, y) = 9x - (y - 1)^2$$

$$0 \ge x^2 + y^2 - 225,$$

$$0 \ge x - 3y + 10$$

On the left, we see the two objective functions that we are minimizing. On the right, we have the constraints. Note they are all inequality constraints.

The MOEA Framework represents constraints a bit differently. Instead of needing to know the constraint formula, the MOEA Framework simply says "set the constraint value to 0 if the solution is feasible, set it to any non-zero value if the constraint is violated." For example, take the constraint $0 \ge x - 3y + 10$. You would typically express this constraint within the MOEA Framework using the ternary if-else expression $x - 3y + 10 \le 0$? 0: x - 3y + 10. This expression begins with the comparator $x - 3y + 10 \le 0$ that tests if the constraint is satisfied. If satisfied, the resulting is 0. Otherwise, the result is x - 3y + 10. Why do we set the value to x - 3y + 10 when the constraint is violated? It is useful in optimization to know how far a solution is from the feasibility boundary. By setting the constraint value to smaller values the closer a solution likes in proximity to the feasibility boundary, the optimization algorithm can guide search towards the feasible region. For the Srinivas problem, we would evaluate the problem as follows:

```
public void evaluate(Solution solution) {
    double x = EncodingUtils.getReal(solution.getVariable(0));
```

```
double y = EncodingUtils.getReal(solution.getVariable(1));
double f1 = Math.pow(x - 2.0, 2.0) + Math.pow(y - 1.0, 2.0) + 2.0;
double f2 = 9.0*x - Math.pow(y - 1.0, 2.0);
double c1 = Math.pow(x, 2.0) + Math.pow(y, 2.0) - 225.0;
double c2 = x - 3.0*y + 10.0;

solution.setObjective(0, f1);
solution.setObjective(1, f2);
solution.setConstraint(0, c1 <= 0.0 ? 0.0 : c1);
solution.setConstraint(1, c2 <= 0.0 ? 0.0 : c2);
}</pre>
```

Lets try another example. Suppose we have the constraint $x^2 + y \le 10$. The trick here is to remember that we want to assign non-zero values when the constraint is violated. It is useful to convert the constraint into the form $h(x) \le 0$, or $x^2 + y - 10 \le 0$. The resulting constraint calculation would be:

```
double c = Math.pow(x, 2.0) + y - 10;
solution.setConstraint(0, c <= 0.0 ? 0.0 : c);</pre>
```

Great! You'll probably notice that there is an additional constraint in our Srinivas problem: $-20 \le x, y \le 20$. This is different from equality and inequality constraints because these are constraints placed on the decision variables. In the MOEA Framework, we do not need to explicitly specify this as a constraint. As shown below, we set these bounds when defining the decision variables.

```
public Solution newSolution() {
    Solution solution = new Solution(2, 2, 2);

solution.setVariable(0, EncodingUtils.newReal(-20.0, 20.0));
solution.setVariable(1, EncodingUtils.newReal(-20.0, 20.0));

return solution;
}
```

3.1 Constrained Optimization Example

Ok, now we can fully define the constrained Srinivas problem. From the problem statement, we see that the Srinivas problem has two decision variables, two objectives, and two constraints. The two decision variables are real-valued and bounded by [-20, 20]. The equations for the objectives and constraints are given.

Within Eclipse, create a new package named chapter3 and create the class SrinivasProblem. Enter the following code:

```
package chapter3;
2
  import org.moeaframework.core.Solution;
3
  import org.moeaframework.core.variable.EncodingUtils;
  import org.moeaframework.problem.AbstractProblem;
  public class SrinivasProblem extends AbstractProblem {
7
8
    public SrinivasProblem() {
9
      super(2, 2, 2);
    @Override
13
    public void evaluate(Solution solution) {
14
      double x = EncodingUtils.getReal(solution.getVariable(0));
      double y = EncodingUtils.getReal(solution.getVariable(1));
16
17
      double f1 = Math.pow(x - 2.0, 2.0) + Math.pow(y - 1.0, 2.0) + 2.0;
      double f2 = 9.0 * x - Math.pow(y - 1.0, 2.0);
      double c1 = Math.pow(x, 2.0) + Math.pow(y, 2.0) - 225.0;
19
      double c2 = x - 3.0*y + 10.0;
20
      solution.setObjective(0, f1);
22
      solution.setObjective(1, f2);
23
      solution.setConstraint(0, c1 \le 0.0 ? 0.0 : c1);
24
      solution.setConstraint(1, c2 \le 0.0 ? 0.0 : c2);
25
26
27
    @Override
28
    public Solution newSolution() {
29
      Solution solution = new Solution(2, 2, 2);
30
31
      solution.setVariable(0, EncodingUtils.newReal(-20.0, 20.0));
32
      solution.setVariable(1, EncodingUtils.newReal(-20.0, 20.0));
33
34
      return solution;
35
36
37
```

MOEAFramework/book/chapter3/SrinivasProblem.java

We already explained the components of this code. The primary difference is the addition of the constraints. First, on lines 10 and 28, we pass in three arguments instead of two. The third argument indicates the number of constraints (e.g., **super** (2, 2, 2) and **new** Solution (2, 2, 2)). Secondly, on lines 23-24 we set the constraints. Again, the constraint is 0 when a solution is feasible.

As before, we also need a class to run this example. Create the class

RunSrinivasProblem with the code below:

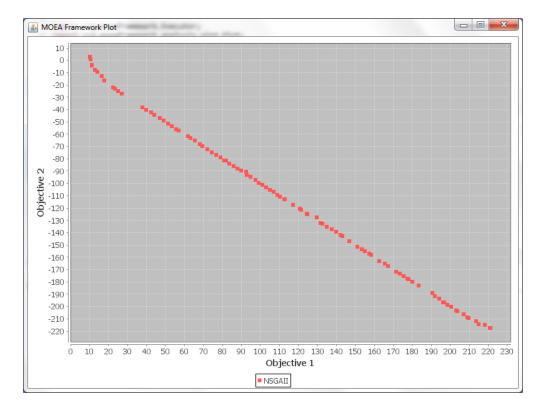
```
package chapter3;
  import org.moeaframework.Executor;
3
  import org.moeaframework.core.NondominatedPopulation;
  import org.moeaframework.core.Solution;
  public class RunSrinivasProblem {
7
8
    public static void main(String[] args) {
9
      NondominatedPopulation result = new Executor()
           .withAlgorithm("NSGAII")
           .withProblemClass (SrinivasProblem.class)
           .withMaxEvaluations(10000)
           .run();
14
15
      for (Solution solution : result) {
16
         if (!solution.violatesConstraints()) {
17
           System.out.format("%10.3f
                                            %10.3f%n",
18
               solution.getObjective(0),
19
               solution.getObjective(1));
20
21
22
23
24
```

MOEAFramework/book/chapter3/RunSrinivasProblem.java

There are a few changes to this code from the previous chapter. First, on line 12 we use the new SrinivasProblem class. Second, on line 17, we check if the solutions are feasible prior to printing the output. Most algorithms will only store feasible solutions, but it's always good practice to check if a solution violates any constraints. With this code input, you can then run the RunSrinivasProblem class and view the output. We could alternatively plot the results for easier viewing:

```
package chapter3;
2
  import org.moeaframework.Executor;
3
  import org.moeaframework.analysis.plot.Plot;
5
  import org.moeaframework.core.NondominatedPopulation;
  public class PlotSrinivasProblem {
8
    public static void main(String[] args) {
9
      NondominatedPopulation result = new Executor()
           .withAlgorithm("NSGAII")
           .withProblemClass(SrinivasProblem.class)
12
           .withMaxEvaluations(10000)
13
```

 ${\it MOEAF ramework/book/chapter 3/PlotSrinivas Problem.} java which produces the following plot:$



3.2 The Knapsack Problem

Ok, now for a more complex example. Have you ever heard of the famous Knapsack problem? If not, check out the Wikipedia page at http://en.wikipedia.org/wiki/Knapsack_problem for more details. This is a famous combinatorial problem that involves choosing which items to place in a knapsack to maximize the value of the items carried without exceeding the weight capacity of the knapsack. More formally, we are given N items. Each item has a profit, P(i), and weight, W(i), for $i=1,2,\ldots,N$. Let d(i) represent our decision to place the i-th item in the knapsack, where d(i)=1 if the item is put into the knapsack and d(i)=0 otherwise. If the knapsack has a weight capacity of C, then the knapsack problem is defined as:

Maximize
$$\sum_{i=1}^{N} d(i) * P(i)$$
 such that $\sum_{i=1}^{N} d(i) * W(i) \leq C$

The summation on the left (which we are maximizing) calculates the total profit we gain from the items placed in the knapsack. The summation on the right side is a constraint that ensures the items placed in the knapsack do not exceed the weight capacity of the knapsack.

Lets make it a little more interesting, after all this is a library for multiobjective optimization. Instead of having one knapsack, lets have two (in fact, this can be generalized to any number of knapsacks). Additionally, the profit and weights vary depending on which knapsack is holding each item. For example, an item will have a profit of \$25 and a weight of 5 pounds in the first knapsack, but will have a profit of \$15 and a weight of 8 pounds in the second knapsack. (It may seem unusual that the weight changes, but that is how the problem is defined in the literature.) Thus, profit is now defined by P(i, j) and weight by W(i, j), where the j = 1, 2 term is the knapsack index. Lastly, each knapsack defines its own capacity, C_1 and C_2 . Combining all of this, the multiobjective knapsack problem is formally defined as:

Maximize
$$\sum_{i=1}^{N} d(i) * P(i,1)$$
 such that $\sum_{i=1}^{N} d(i) * W(i,1) \leq C_1$ and Maximize $\sum_{i=1}^{N} d(i) * P(i,2)$ such that $\sum_{i=1}^{N} d(i) * W(i,2) \leq C_2$

This problem is a bit different from the others we have seen thus far. For the knapsack problem, we are picking items to fit into the knapsack. The bit string representation works well for situation where we are making many yes/no decisions (yes if it is included in the knapsack). For example, if we have 5 items, we can represent the decision to include each item using a bit string with 5 bits. Each bit in the string corresponds to an item, and is set to 1 if the item is included and 0 if the item is excluded. For instance, the bit string 00110 would place items 3 and 4 inside the knapsacks, excluding the rest. Our newSolution method is defined as follows:

```
public Solution newSolution() {
   Solution solution = new Solution(1, nsacks, nsacks);
   solution.setVariable(0, EncodingUtils.newBinary(nitems));
   return solution;
}
```

Observe on line 3 how we create the bit string representation (also called a binary string) with nitems bits. Also note that the 5 bits are contained within a single decision variable, so we define this problem with only a single decision variable.

Now for the evaluate method. Summing up the profits is straightforward. But there is also a constraint we must deal with. We must ensure the weight of the items does not exceed the capacity of the knapsack. Thus, we need to sum up the weights of the selected items and compare to the capacity. The resulting method is shown below:

```
public void evaluate(Solution solution) {
      boolean[] d = EncodingUtils.getBinary(solution.getVariable(0));
2
3
       double[] f = new double[nsacks];
       double[] g = new double[nsacks];
4
       // calculate the profits and weights for the knapsacks
6
       for (int i = 0; i < nitems; i++) {</pre>
7
         if (d[i]) {
           for (int j = 0; j < nsacks; j++) {</pre>
9
             f[j] += profit[j][i];
             g[j] += weight[j][i];
         }
       }
14
       // check if any weights exceed the capacities
16
       for (int j = 0; j < nsacks; j++) {</pre>
17
         if (g[j] <= capacity[j]) {
18
           g[j] = 0.0;
19
20
         } else {
21
           g[j] = g[j] - capacity[j];
         }
22
23
24
       // negate the objectives since Knapsack is maximization
25
       solution.setObjectives(Vector.negate(f));
26
       solution.setConstraints(g);
27
```

Note the comment on line 25. We are negating the objective values since our objectives are being maximized. The MOEA Framework is designed to minimize objectives. To handle maximized objectives, simply negate the value. Minimizing the negated value is equivalent to maximizing the original value. Take caution, however, as the output from the MOEA Framework will include the negative values. You should always negate the outputs to restore them to their correct sign.

Below is the full implementation of the Knapsack problem. We have configured this instance for a simple problem with 2 knapsacks and 5 items. Copy this code into the KnapsackProblem class.

```
package chapter3;

import org.moeaframework.core.Solution;
import org.moeaframework.core.variable.EncodingUtils;
import org.moeaframework.problem.AbstractProblem;
import org.moeaframework.util.Vector;

public class KnapsackProblem extends AbstractProblem {
```

```
9
     /**
      * The number of sacks.
11
    public static int nsacks = 2;
13
14
15
     * The number of items.
16
17
    public static int nitems = 5;
18
19
20
      * Entry {@code profit[i][j]} is the profit from including item {@code j}
21
      * in sack {@code i}.
22
      */
23
     public static int[][] profit = {
24
         {2, 5},
25
         {1, 4},
26
         {6, 2},
27
         {5, 1},
28
         {3, 3}
29
30
     };
31
32
      * Entry {@code weight[i][j]} is the weight incurred from including item
33
      * {@code j} in sack {@code i}.
34
35
     public static int[][] weight = {
36
         {3, 3},
37
         {4, 2},
38
         \{1, 5\},\
39
         {5, 3},
40
         {5, 2}
42
     };
43
44
      * Entry {@code capacity[i]} is the weight capacity of sack {@code i}.
45
46
    public static int[] capacity = { 10, 8 };
47
48
     public KnapsackProblem() {
49
       super(1, nsacks, nsacks);
50
51
     public void evaluate(Solution solution) {
54
       boolean[] d = EncodingUtils.getBinary(solution.getVariable(0));
       double[] f = new double[nsacks];
55
       double[] g = new double[nsacks];
56
57
       // calculate the profits and weights for the knapsacks
58
       for (int i = 0; i < nitems; i++) {</pre>
59
```

```
60
         if (d[i]) {
           for (int j = 0; j < nsacks; j++) {</pre>
61
             f[j] += profit[j][i];
62
             g[j] += weight[j][i];
63
64
         }
65
       }
66
67
       // check if any weights exceed the capacities
68
       for (int j = 0; j < nsacks; j++) {</pre>
         if (g[j] <= capacity[j]) {
70
           g[j] = 0.0;
71
         } else {
72
           g[j] = g[j] - capacity[j];
74
       }
75
76
       // negate the objectives since Knapsack is maximization
       solution.setObjectives(Vector.negate(f));
78
       solution.setConstraints(g);
79
    }
80
81
    public Solution newSolution() {
82
       Solution solution = new Solution(1, nsacks, nsacks);
83
       solution.setVariable(0, EncodingUtils.newBinary(nitems));
84
       return solution;
85
86
87
```

MOEAFramework/book/chapter3/KnapsackProblem.java Next, copy the following code into the RunKnapsackProblem class.

```
package chapter3;
2
  import java.io.IOException;
3
  import org.moeaframework.Executor;
  import org.moeaframework.core.NondominatedPopulation;
6
  import org.moeaframework.core.Solution;
  import org.moeaframework.examples.ga.knapsack.Knapsack;
  import org.moeaframework.util.Vector;
  public class RunKnapsackProblem {
    public static void main(String[] args) throws IOException {
13
      NondominatedPopulation result = new Executor()
14
           .withProblemClass (Knapsack.class)
           .withAlgorithm("NSGAII")
16
           .withMaxEvaluations (10000)
17
           .distributeOnAllCores()
18
```

```
19
           .run();
20
       for (int i = 0; i < result.size(); i++) {</pre>
21
         Solution solution = result.get(i);
22
         double[] objectives = solution.getObjectives();
23
24
25
         // negate objectives to return them to their maximized form
         objectives = Vector.negate(objectives);
26
27
         System.out.println("Solution " + (i+1) + ":");
28
         System.out.println("
                                   Sack 1 Profit: " + objectives[0]);
29
                                   Sack 2 Profit: " + objectives[1]);
         System.out.println("
30
         System.out.println("
                                   Binary String: " + solution.getVariable(0));
31
32
33
34
35
```

MOEAFramework/book/chapter3/RunKnapsackProblem.java

As before, we specify the problem class when creating the Executor. Observe that we do not need to thell the algorithm about any new features of our problem. It automatically detects that the problem uses a bit string representation and constructs the algorithm with the appropriate crossover and mutation operators. Also note on line 18 that we call distributeOnAllCores(), which enables multithreading. For problems with time-consuming evaluations, you can gain substantial performance improvements by spreading the work across multiple processors on your computer. The MOEA Framework automatically handles this for you.

Running this example will produce output similar to the following. Since this is a multiobjective problem, there is typically no single optimal solution. Instead, there are several options, all equally "good". Solution 1 has the maximum profit of 9.0 for Sack 1, but the worst profit of 5.0 for Sack 2.

```
Solution 1:
    Sack 1 Profit: 9.0
    Sack 2 Profit: 5.0
    Binary String: 01010
Solution 2:
    Sack 1 Profit: 5.0
    Sack 2 Profit: 8.0
    Binary String: 00110
Solution 3:
    Sack 1 Profit: 7.0
    Sack 2 Profit: 6.0
    Binary String: 00011
Solution 4:
    Sack 1 Profit: 6.0
    Sack 2 Profit: 7.0
    Binary String: 01100
```

In this Knapsack problem, we hard-coded the profits, weights, and capacities. A more generalized implementation is available with the MOEA Framework in the examples folder.

3.3 Feasibility

You may encounter problems that are severely constrained where the majority of the search space is infeasible. Under these circumstances, it becomes very challenging for an optimization algorithm to find feasible solutions. When this happens, the results from the MOEA Framework will contain only infeasible solutions. It is important to check if solutions are infeasible with the solution.violatesConstraints() method. This can be a challenging problem to address. In some cases, the problem can be reformulated or represented in a different way to relax the constraints. This is outside the scope of this beginner's guide, but more details can be found in academic literature.

Chapter 4

Choice of Optimization Algorithm

Up to this point, we have been using one of the most popular MOEAs: NSGA-II. NSGA-II is frequently used in practice as it shows strong performance on a range of problems and has been studied extensively in the academic literature. However, the MOEA Framework is not limited to these algorithms. This chapter will discuss how to run different algorithms, how to parameterize the algorithms, and how to compare the performance of algorithms.

4.1 Running Different Algorithms

The MOEA Framework contains an assortment of optimization algorithms for your use, each identified by name. For example, the following optimizes the Schaffer problem using the NSGA-II algorithm:

```
NondominatedPopulation result = new Executor()
    .withProblemClass(SchafferProblem.class)
    .withAlgorithm("NSGAII")
    .withMaxEvaluations(10000)
    .run();
```

Likewise, we can optimize the problem with GDE3 as demonstrated below. Observe that we only needed to change the name of the algorithm.

```
NondominatedPopulation result = new Executor()
    .withProblemClass(SchafferProblem.class)
    .withAlgorithm("GDE3")
    .withMaxEvaluations(10000)
    .run();
```

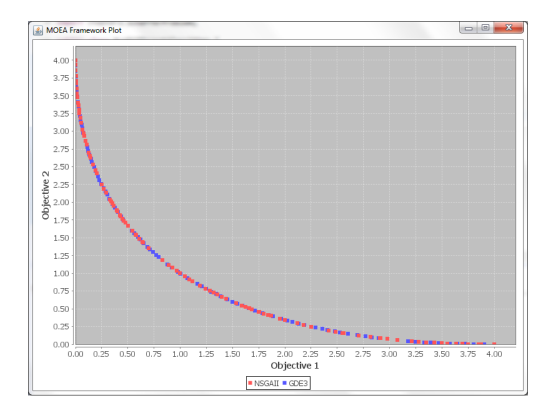
The MOEA Framework handles the rest. It ensures the algorithms are configured appropriately for the given problem or issues a warning or error if it detects any problems. The

full code to run this example and plot the results is provided below.

```
package chapter4;
  import java.io.IOException;
3
  import org.moeaframework.Executor;
5
  import org.moeaframework.analysis.plot.Plot;
6
  import org.moeaframework.core.NondominatedPopulation;
  import chapter2.SchafferProblem;
  public class RunDifferentAlgorithms {
10
    public static void main(String[] args) throws IOException {
      NondominatedPopulation result1 = new Executor()
13
           .withProblemClass(SchafferProblem.class)
14
           .withAlgorithm("NSGAII")
15
           .withMaxEvaluations(10000)
16
           .run();
17
18
      NondominatedPopulation result2 = new Executor()
19
           .withProblemClass(SchafferProblem.class)
20
           .withAlgorithm("GDE3")
           .withMaxEvaluations(10000)
22
           .run();
24
      new Plot()
25
           .add("NSGAII", result1)
26
           .add("GDE3", result2)
27
           .show();
28
29
30
31
```

MOEAFramework/book/chapter4/RunDifferentAlgorithms.java

The output, which shows the two algorithms colored red and blue, is shown below. Visually, we can see the two algorithms produce nearly identical results. This visual approach is useful to quickly observe the relative performance of two or more algorithms. Later in this chapter we'll demonstrate how to statistically compare the performance of algorithms.



Note that not all algorithms can solve all problems. Table 4.1 provides a list of the algorithms supported natively by the MOEA Framework. The first column provides the name (which you specify in the string to withAlgorithm), and the remaining columns indicate if the algorithm supports different encodings and constraints.

In addition to these algorithms, which are implemented natively within the MOEA Framework, algorithms from the population JMetal and PISA libraries can also be executed within the MOEA Framework with little additional work. Refer to See Appendix B for more details.

4.2 Parameterization

In addition to selecting the specific algorithm to use, each algorithm has its own parameters that can be customized. If left untouched, the algorithm uses default parameters based on best practices from the literature. All parameters can also be overridden. Appendix A lists the parameters for each algorithm, provides a short description, and indicates the default value. For example, lets compare the default parameters for NSGA-II to a custom set of parameters. Here, we are reducing the population size from 100 (the default) to 50 and increasing the distribution indices which affect how offspring are generated. We would expect these custom parameters to perform worse in practice, but lets see for ourselves:

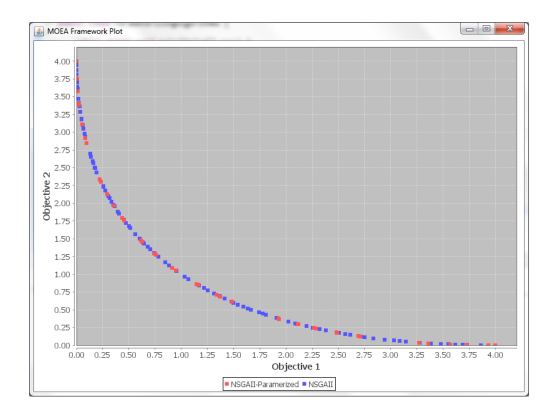
```
package chapter4;
```

Table 4.1: List of available optimization algorithms.

```
import org.moeaframework.Executor;
  import org.moeaframework.analysis.plot.Plot;
  import org.moeaframework.core.NondominatedPopulation;
5
  import chapter2.SchafferProblem;
7
9
  public class ParameterizingAlgorithms {
10
    public static void main(String[] args) {
      NondominatedPopulation result1 = new Executor()
           .withProblemClass(SchafferProblem.class)
           .withAlgorithm("NSGAII")
14
           .withMaxEvaluations(10000)
           .run();
16
17
      NondominatedPopulation result2 = new Executor()
18
           .withProblemClass(SchafferProblem.class)
19
           .withAlgorithm("NSGAII")
20
           .withMaxEvaluations(10000)
21
           .withProperty("populationSize", 50)
           .withProperty("sbx.rate", 1.0)
           .withProperty("sbx.distributionIndex", 250.0)
24
           .withProperty("pm.rate", 0.0)
25
           .withProperty("pm.distributionIndex", 300.0)
26
           .run();
27
28
      new Plot()
29
           .add("NSGAII-Paramerized", result2)
30
           .add("NSGAII", result1)
           .show();
32
33
34
```

MOEAF ramework/book/chapter 4/Parameterizing Algorithms. java

As shown below, the custom parameters (shown in red) does not approximate the Pareto surface as well as the default parameters (shown in blue). The smaller population size results in fewer points, and the larger distribution indices prevents the algorithm from effectively generating new offspring.



Many optimization algorithms support customizable crossover and mutation operators, and these operators have their own parameters founds in Appendix C. For example, NSGA-II has a single parameter, "populationSize" and uses the Simulated Binary Crossover and Polynomial Mutation operators for real-valued problems. These two operators have two parameters each. We can customize these parameters as demonstrated below.

```
NondominatedPopulation result = new Executor()
    .withProblemClass(SchafferProblem.class)
    .withAlgorithm("NSGAII")
    .withMaxEvaluations(10000)
    .withProperty("populationSize", 200)
    .withProperty("sbx.rate", 0.9)
    .withProperty("sbx.distributionIndex", 25.0)
    .withProperty("pm.rate", 0.1)
    .withProperty("pm.distributionIndex", 30.0)
    .run();
```

Some algorithms also allow overriding which operators are used. For example, we can choose to use Parent Centric Crossover with NSGA-II:

```
NondominatedPopulation result = new Executor()
.withProblemClass(SchafferProblem.class)
.withAlgorithm("NSGAII")
.withMaxEvaluations(10000)
```

```
.withProperty("populationSize", 200)
.withProperty("operator", "pcx")
.withProperty("pcx.parents", 5)
.withProperty("pcx.offspring", 1)
.withProperty("pcx.eta", 0.1)
.withProperty("pcx.zeta", 0.1)
.run();
```

Note the use of the "operator" parameter to specify the use of Parent Centric Crossover with the string "pcx". Refer to Appendix C for a list of available operators and their string abbreviations. Operators can be combined using the + symbol. For example, by default we use Simulated Binary Crossover (SBX) and Polynomial Mutation (PM), which can be expressed as "sbx+pm". This says to first apply Simulated Binary Crossover on two parents to produce two offspring, then apply Polynomial Mutation to each offspring. Alternatively, we could combine Differential Evolution (DE) and Polynomial Mutation (PM) using the string "de+pm". This is the operator used by GDE3, but as shown below, it can also be used with NSGA-II:

```
NondominatedPopulation result = new Executor()
    .withProblemClass(SchafferProblem.class)
    .withAlgorithm("NSGAII")
    .withProperty("operator", "de+pm")
    .withMaxEvaluations(10000)
    .run();
```

Several precautions should be observed. First, not all combinations of operators are valid. You will see an exception if you request an invalid combination, typically saying invalid number of parents. See the User Manual for more details. Second, be careful when passing in the parameters. Parameter names are case sensitive. "populationsize" is not the same as "populationsize". Additionally, you will not be notified if an invalid parameter is provided, it will simply be ignored.

4.3 Comparing Algorithms

In academic work, it is often useful to compare the performance of two or more MOEAs. For instance, you could have developed a new search operator or algorithm and wish to compare the performance against existing MOEAs to determine the level of improvement.

The MOEA Framework provides the Analyzer class, which is typically used in conjunction with the Executor class we have seen previously, to analyze the performance of algorithms. Additionally, it can statistically compare the performance of two or more algorithms to determine, for a given confidence level, if there is any statistical difference between the two algorithms. For example, the following code compares three algorithms — NSGA-II,

GDE3, and ϵ -MOEA — on the UF1 test problem.

```
package chapter4;
  import java.io.IOException;
3
  import org.moeaframework.Analyzer;
5
  import org.moeaframework.Executor;
6
  import org.moeaframework.analysis.plot.Plot;
  public class ComparingAlgorithms {
9
    public static void main(String[] args) throws IOException {
       String problem = "UF1";
       String[] algorithms = { "NSGAII", "GDE3", "eMOEA" };
13
14
15
       //setup the experiment
      Executor executor = new Executor()
16
           .withProblem(problem)
17
           .withMaxEvaluations(10000);
18
19
      Analyzer analyzer = new Analyzer()
20
           .withSameProblemAs(executor)
           .includeHypervolume()
22
           .showStatisticalSignificance();
24
       //run each algorithm for 50 seeds
25
       for (String algorithm : algorithms) {
26
         analyzer.addAll(algorithm,
27
             executor.withAlgorithm(algorithm).runSeeds(50));
28
29
30
       //print the results
       analyzer.printAnalysis();
32
       //plot the results
34
      new Plot()
35
           .add(analyzer)
36
           .show();
38
39
```

MOEAFramework/book/chapter4/ComparingAlgorithms.java

On lines 16-18, we setup the Executor for our test problem, UF1. Note that unlike the previous examples of using the Executor, we do not specify the algorithm. We will do this later. Next, on lines 20-23, the setup the Analyzer. We again must indicate on line 21 which problem we are solving so it can compute the performance metrics correctly. On line 22 we specify that we are interested in the hypervolume performance metric and on line 23 request that we compute the statistical significance of the results (with 95% confidence level

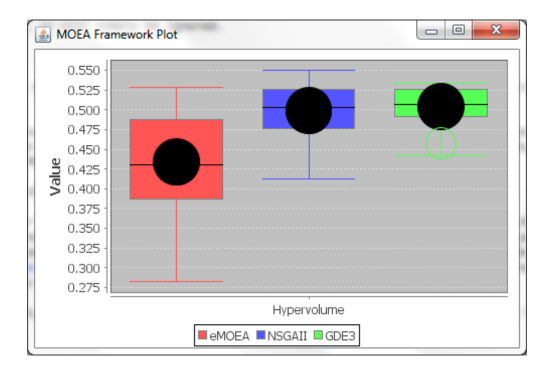
by default). The for loop in lines 26-29 actually performs the analysis. Looping over each algorithm in our study, we setup the Executor to run the given algorithm for 50 seeds on line 28. The results of the 50 seeds is added to the analyzer on line 27. Finally, we print the output on line 32. Alternatively, we can plot the differences visually on lines 35-37.

The text output of this code displays the minimum, median, and maximum hypervolume value achieved by each algorithm and the statistical significance.

```
GDE3:
    Hypervolume:
        Min: 0.46862049421426344
        Median: 0.5035596089917667
        Max: 0.534267455034584
        Count: 50
        Indifferent: []
eMOEA:
    Hypervolume:
        Min: 0.3784906829802386
        Median: 0.4503635874660789
        Max: 0.5322735014135916
        Count: 50
        Indifferent: [NSGAII]
NSGAII:
    Hypervolume:
        Min: 0.33173049116232906
        Median: 0.45913525261150123
        Max: 0.5370813537070005
        Count: 50
        Indifferent: [eMOEA]
```

When checking for the statistical significance as we have done, the output will contain the Indifferent field listing, if any, the algorithms with statistically indifferent performance. That is to say, with 95% confidence, the performance of NSGA-II and eMOEA are identical, or not statistically different.

The graphical output of this code produces the box-and-whisker plot shown below. The results of each algorithm is represented by a colored box. The filled, rectangular box shows the range encompassed by the 25% to 75% percentile (i.e., the middle 50% of the data) and the horizontal line shows the median. The colored whiskers (the thin lines at the top and bottom of the rectangular box) show the regular min and max of the data. The solid black circle shows the mean. Using this plot, we can see quickly that ϵ -MOEA performs worse than GDE3 and NSGA-II (larger values are preferred).



In addition to the hypervolume performance metric show above, the MOEA Framework also supports generational distance, inverted generational distance, additive ϵ -indicator, maximum Pareto front error, spacing, contribution, and the R2 indicator (Knowles and Corne, 2002; Coello Coello et al., 2007; Hansen et al., 1998). The R1 and R3 indicators are also available, but are not often used in practice as the R2 indicator exhibits better properties.

Table 4.2 shows the supported indicators and key properties. Pareto compliant implies that better indicator values correspond to approximation sets that are preferred by weak Pareto dominance. This is a good property for performance metrics as better values (closer to the target) indicate better Pareto approximate sets.

Each indicator is enabled by calling the corresponding include method, such as includeHypervolume() for the hypervolume metric. In this example, we analyzed the UF1 test problem which has a known reference set. The reference set contains all Pareto optimal solutions for a given problem (or a good approximation of the set). The MOEA Framework automatically loads the reference sets for built-in problems. For custom problems, there are two options. First, if you do not explicitly load a reference set, then the Analyzer automatically combines all results to produce a reference set. In this case, the reference set contains all Pareto optimal solutions generated by the runs. Alternatively, you can call analyzer.withReferenceSet(new File("set.txt")) to load the reference set from a file. This file simply contains the objective values for each Pareto optimal solution. For example, if the Pareto set was a straight line from (0, 1) to (1, 0), then the reference set file would be:

^{0.0 1.0}

^{0.1 0.9}

^{0.2 0.8}

Table 4.2: List of available performance metrics.

Indicator	Pareto Compliant	Pareto Compliant Reference Set Required Normalized Target	Normalized	Target
Hypervolume	Yes	Yes	Yes	Maximize $\rightarrow 1$
Generational Distance	No	Yes	Yes	Minimize $\rightarrow 0$
Inverted Generational Distance	$N_{\rm O}$	Yes	Yes	Minimize $\rightarrow 0$
Additive ϵ -Indicator	Yes	m No	Yes	Minimize $\rightarrow 0$
Maximum Pareto Front Error	$N_{\rm O}$	Yes	Yes	Minimize $\rightarrow 0$
Spacing	$N_{\rm O}$	m No	$N_{\rm o}$	Minimize $\rightarrow 0$
Contribution	Yes	Yes	$N_{\rm o}$	Maximize $\rightarrow 1$
R2 Indicator	$N_{\rm O}$	Yes	Yes	Minimize \rightarrow -1

```
0.3 0.7

0.4 0.6

0.5 0.5

0.6 0.4

0.7 0.3

0.8 0.2

0.9 0.1

1.0 0.0
```

The ordering of solutions in the reference set file does not matter. When calculating hypervolume, it is sometimes a good practice to specify a custom ideal and reference point. This can be achieved by calling, for example, analyzer.withIdealPoint(0.0, 0.0) and analyzer.withReferencePoint(2.0, 2.0), to compute the hypervolume within the bounds (0,0) and (2,2).

4.4 Runtime Dynamics

In the previous section, we saw how to statistically compare the performance of two or more algorithms. In doing so, we are only comparing the end-of-run performance of the algorithm using only the final Pareto approximation. It can also be useful to investigate the intermediate performance of an algorithm. That is, we want to see how the performance of the algorithm evolves over time. Additionally, we can also inspect other properties of the algorithm to see how, internally, they adapt over time.

To collect runtime dynamics, we introduce the Instrumenter class. The Instrumenter gets its name from its ability to add instrumentation, which are pieces of code that record information, to an algorithm. A variety of data can be collected using the Instrumenter, including:

- 1. Elapsed time
- 2. Population size / archive size
- 3. The approximation set
- 4. Performance metrics
- 5. Restart frequency

The Instrumenter works hand-in-hand with the Executor to collect its data. The Executor is responsible for configuring and running the algorithm, but it allows the Instrumenter to record the necessary data while the algorithm is running. To start collecting run-time dynamics, we first create and configure an Instrumenter instance.

First, line 1 creates the new Instumenter instance. Next, line 2 specifies the problem. This allows the instrumenter to access the known reference set for this problem, which is necessary for evaluating performance metrics. Third, line 3 sets the frequency at which the data is recorded. In this example, data is recorded every 100 evaluations. Lastly, lines 4 and 5 indicate that only the elapsed time and generational distance should be recorded. Calling .attachAll() will collect all available data.

Next, we create and configure the Executor instance with the following code snippet:

```
new Executor()
    .withSameProblemAs(instrumenter)
    .withAlgorithm("NSGAII")
    .withMaxEvaluations(10000)
    .withInstrumenter(instrumenter)
    .run();
```

This code snippet is similar to the previous examples of the Executor, but includes the addition of line 5. Line 5 tells the executor that all algorithms it executes will be instrumented with our Instrumenter instance. Once the instrumenter is set and the algorithm is configured, we can run the algorithm on line 6.

When the run completes, we can access the data collected by the instrumenter. The data is stored in an Accumulator object. The Accumulator for the run we just executed can be retrieved with the following line:

```
Accumulator accumulator = instrumenter.getLastAccumulator();
```

An Accumulator is similar to a Map in that it stores key-value pairs. The key identifies the type of data recorded. However, instead of storing a single value, the Accumulator stores many values, one for each datapoint collected by the Instrumenter. Recall that in this example, we are recording a datapoint every 100 evaluations (i.e., the frequency). We could choose to either print the contents to the console:

```
System.out.println(accumulator.toCSV());
```

or save the contents to a CSV file:

```
accumulator.saveCSV(new File("output.csv"));
```

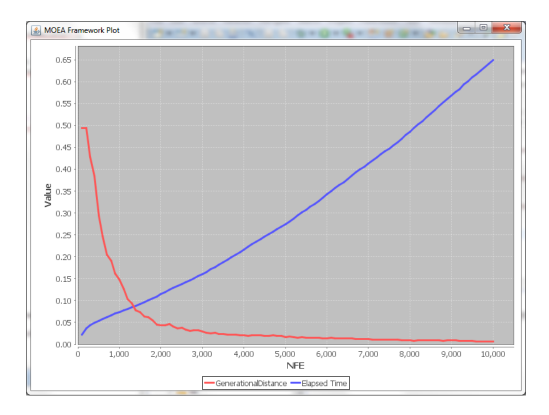
or plot the data in a line graph:

The full code example for collecting the runtime dynamics and plotting the results is shown below:

```
package chapter4;
  import java.io.IOException;
3
  import org.moeaframework.Executor;
  import org.moeaframework.Instrumenter;
  import org.moeaframework.analysis.collector.Accumulator;
  import org.moeaframework.analysis.plot.Plot;
  public class RuntimeDynamics {
    public static void main(String[] args) throws IOException {
12
      Instrumenter instrumenter = new Instrumenter()
           .withProblem("UF1")
14
           .withFrequency(100)
           .attachElapsedTimeCollector()
16
           .attachGenerationalDistanceCollector();
17
18
      new Executor()
19
           .withSameProblemAs(instrumenter)
20
           .withAlgorithm("NSGAII")
           .withMaxEvaluations(10000)
           .withInstrumenter(instrumenter)
           .run();
24
25
      Accumulator accumulator = instrumenter.getLastAccumulator();
26
      new Plot()
28
           .add(accumulator)
           .show();
30
31
32
33
```

MOEAFramework/book/chapter4/RuntimeDynamics.java

This code produces the following plot:



Rather than plotting all the data in an accumulator, you can also plot specific metrics. As shown below, we only plot the generational distance:

Chapter 5

Customizing Algorithms

In previous chapters, we used the Executor class to optimize problems. The Executor is great for quickly solving problems using predefined optimization algorithms. However, it is not very customizable. This chapter discusses step for customizing algorithms.

5.1 Manually Running Algorithms

Before we discuss how to customize an algorithm, lets start by looking at what happens behind the scenes in the Executor class. Take the following code, for example.

```
NondominatedPopulation result = new Executor()
.withAlgorithm("NSGAII")
.withProblemClass(SchafferProblem.class)
.withMaxEvaluations(10000)
.run();
```

This code creates a new instance of the NSGA-II algorithm and uses it to optimize the Schaffer problem. During optimization, it spends at most 10,000 evaluations. Finally, the Pareto approximate result is saved to the result variable. Underneath the hood, these same steps can be accomplished as shown below:

```
package chapter5;

import org.moeaframework.algorithm.NSGAII;
import org.moeaframework.core.Algorithm;
import org.moeaframework.core.Initialization;
import org.moeaframework.core.NondominatedPopulation;
import org.moeaframework.core.NondominatedSortingPopulation;
import org.moeaframework.core.Problem;
import org.moeaframework.core.Variation;
import org.moeaframework.core.Variation;
import org.moeaframework.core.comparator.ChainedComparator;
import org.moeaframework.core.comparator.CrowdingComparator;
```

```
import org.moeaframework.core.comparator.ParetoDominanceComparator;
  import org.moeaframework.core.operator.GAVariation;
  import org.moeaframework.core.operator.RandomInitialization;
  import org.moeaframework.core.operator.TournamentSelection;
  import org.moeaframework.core.operator.real.PM;
  import org.moeaframework.core.operator.real.SBX;
  import chapter2.SchafferProblem;
18
19
  public class ManualRun {
20
    public static void main(String[] args) {
22
      // define the problem
23
      Problem problem = new SchafferProblem();
24
      // create an initial random population of 100 individuals
26
      Initialization initialization = new RandomInitialization(
27
          problem,
2.8
          100);
30
      // define the selection operator
      TournamentSelection selection = new TournamentSelection(2,
32
33
          new ChainedComparator(
               new ParetoDominanceComparator(),
34
               new CrowdingComparator()));
35
36
      // define the crossover / mutation operator
      Variation variation = new GAVariation (
38
          new SBX(1.0, 25.0),
39
          new PM(1.0 / problem.getNumberOfVariables(), 30.0));
40
41
      // construct the algorithm
42
      Algorithm algorithm = new NSGAII(
43
          problem,
          new NondominatedSortingPopulation(),
45
          null, // no archive
          selection,
47
          variation,
48
          initialization);
49
      // run the algorithm for 10,000 evaluations
      while (algorithm.getNumberOfEvaluations() < 10000) {</pre>
        algorithm.step();
54
      // get the Pareto approximate results
56
      NondominatedPopulation result = algorithm.getResult();
57
58
59
60
```

MOEAFramework/book/chapter5/ManualRun.java

As can be seen, we must define each component of the algorithm manually, as listed below:

Line 24 Problem definition

Lines 27-29 Initialization operator - Produces a randomly-generated initial population with 100 individuals

Lines 32-35 Selection operator - Binary tournament selecting using Pareto dominance and crowding distance

Lines 38-40 Variation operator - Simulated binary crossover with polynomial mutation

Lines 43-49 Algorithm definition - NSGA-II with the given operators

Lines 52-54 Runs the algorithm for 10,000 evaluations

Line 57 Retrieve the Pareto approximate results

Customizing an algorithm requires changing the components above as desired. In the remainder of this chapter we'll consider several customizations.

5.2 Custom Initialization

Lets say instead of the randomly-generated initial population, we would rather use the Latin hypercube distribution. We would begin by implementing the Initialization interface and generating the initial population using a Latin hypercube distribution:

```
package chapter5;
2
  import org.moeaframework.core.Initialization;
  import org.moeaframework.core.Problem;
  import org.moeaframework.core.Solution;
  import org.moeaframework.core.variable.RealVariable;
  import org.moeaframework.util.sequence.LatinHypercube;
  public class LatinHypercubeInitialization implements Initialization {
9
10
    private Problem problem;
    private int size;
14
    public LatinHypercubeInitialization(Problem problem, int size) {
      super();
16
      this.problem = problem;
17
      this.size = size;
18
    }
19
20
    @Override
21
```

```
22
    public Solution[] initialize() {
       LatinHypercube lhs = new LatinHypercube();
       double[][] samples = lhs.generate(size, problem.getNumberOfObjectives());
24
       Solution[] result = new Solution[size];
25
26
       for (int i = 0; i < size; i++) {</pre>
27
         Solution solution = problem.newSolution();
28
29
         for (int j = 0; j < problem.getNumberOfVariables(); j++) {</pre>
30
           RealVariable variable = (RealVariable) solution.getVariable(j);
           variable.setValue(samples[i][j] * (variable.getUpperBound()-variable.
32
              getLowerBound())
               + variable.getLowerBound());
         result[i] = solution;
37
38
      return result;
39
40
41
```

MOEAFramework/book/chapter5/LatinHypercubeInitialization.java

When implementing the Initialization interface, we must include the Solution [] initialize() method to return the initial population. Fortunately, we already have code that produces the Latin hypercube samples (LHS), but they are scaled on the range [0,1]. Thus, we first generate the LHS samples on the range [0,1] on lines 23-24. Next, we loop over each of these samples on line 27. For each sample, we create a new Solution instance (line 28). On lines 30-34, we assign the value of each decision variable, being sure to scale from the range [0,1] to the lower and upper bounds of the decision variable. Each solution is added to the result array and returned on line 39.

We then replace the RandomInitialization class with our new LatinHypercubeInitialization on line 27-29 below:

```
package chapter5;

import org.moeaframework.algorithm.NSGAII;
import org.moeaframework.core.Algorithm;
import org.moeaframework.core.Initialization;
import org.moeaframework.core.NondominatedPopulation;
import org.moeaframework.core.NondominatedSortingPopulation;
import org.moeaframework.core.Problem;
import org.moeaframework.core.Variation;
import org.moeaframework.core.comparator.ChainedComparator;
import org.moeaframework.core.comparator.CrowdingComparator;
import org.moeaframework.core.comparator.ParetoDominanceComparator;
import org.moeaframework.core.operator.GAVariation;
import org.moeaframework.core.operator.TournamentSelection;
```

```
import org.moeaframework.core.operator.real.PM;
  import org.moeaframework.core.operator.real.SBX;
16
17
  import chapter2.SchafferProblem;
18
19
  public class CustomInitialization {
20
21
    public static void main(String[] args) {
22
       // define the problem
23
      Problem problem = new SchafferProblem();
24
       // create an LHS initial population with 100 individuals
26
       Initialization initialization = new LatinHypercubeInitialization(
27
           problem,
           100);
30
       // define the selection operator
31
      TournamentSelection selection = new TournamentSelection(2,
           new ChainedComparator(
33
               new ParetoDominanceComparator(),
34
               new CrowdingComparator()));
35
36
       // define the crossover / mutation operator
37
      Variation variation = new GAVariation (
38
           new SBX(1.0, 25.0),
39
           new PM(1.0 / problem.getNumberOfVariables(), 30.0));
40
41
       // construct the algorithm
42
      Algorithm algorithm = new NSGAII(
          problem,
44
           new NondominatedSortingPopulation(),
45
           null, // no archive
46
           selection,
           variation,
48
           initialization);
49
50
       // run the algorithm for 10,000 evaluations
      while (algorithm.getNumberOfEvaluations() < 10000) {</pre>
         algorithm.step();
53
54
       // get the Pareto approximate results
56
      NondominatedPopulation result = algorithm.getResult();
57
58
59
60
```

MOEAFramework/book/chapter5/CustomInitialization.java

5.3 Custom Algorithms

It is very convenient to develop new optimization algorithms within the MOEA Framework. Not only can you build the algorithm using the existing components already built into the MOEA Framework, but your algorithm will work with all of the analytical and plotting tools available.

All optimization algorithms implemented within the MOEA Framework must implement the Algorithm interface. This interfaces defines several methods that must be implemented. For convenience, several abstract classes are also provided, including:

- **Algorithm** Used when implementing an algorithm that can not use any of the abstract classes discussed below. Provides the greatest flexibility, but requires the most development.
- **AbstractAlgorithm** Used when creating iterative optimization algorithms. The developer must provide the initialize() and iterate() method.
- **AbstractEvolutionaryAlgorithm** Used when creating evolutionary algorithms that typically optimize a population of candidate solutions.
- **AbstractPSOAlgorithm** Used when implementing particle swarm optimization algorithms. Provides structures for storing and updating particle positions, velocities, etc.

To demonstrate the creation of a new algorithm, we will construct a very simple optimization algorithm called the RandomWalker. The overall algorithm is as follows:

- 1. Initialize the population with random solutions
- 2. While NFE < MaxNFE
 - (a) Randomly pick a solution from the population
 - (b) Mutate the individual with polynomial mutation
 - (c) Add the offspring to the population
 - (d) Use non-dominated sorting to remove the worst member of the population

From the above description, we see that the algorithm uses a population of solutions. Therefore, we will extend the AbstractEvolutionaryAlgorithm, as shown below.

```
package chapter5;

import org.moeaframework.algorithm.AbstractEvolutionaryAlgorithm;
import org.moeaframework.core.Initialization;
import org.moeaframework.core.NondominatedSortingPopulation;
import org.moeaframework.core.PRNG;
import org.moeaframework.core.Problem;
```

```
import org.moeaframework.core.Solution;
  import org.moeaframework.core.Variation;
  public class RandomWalker extends AbstractEvolutionaryAlgorithm {
    private final Variation variation;
14
    public RandomWalker (Problem problem, Initialization initialization,
15
        Variation variation) {
16
      super(problem, new NondominatedSortingPopulation(), null, initialization);
      this.variation = variation;
18
    }
19
20
    @Override
21
    protected void iterate() {
22
      // get the current population
23
      NondominatedSortingPopulation population = (NondominatedSortingPopulation)
24
          getPopulation();
25
      // randomly select a solution from the population
26
      int index = PRNG.nextInt(population.size());
28
      Solution parent = population.get(index);
29
      // mutate the selected solution
30
      Solution offspring = variation.evolve(new Solution[] { parent })[0];
      // evaluate the objectives/constraints
33
      evaluate (offspring);
34
35
      // add the offspring to the population
36
      population.add(offspring);
37
38
      // use non-dominated sorting to remove the worst solution
39
      population.truncate(population.size()-1);
40
41
42
```

MOEAFramework/book/chapter5/RandomWalker.java

Observe that our constructor on line 15 accepts three arguments: the problem definition, the initialization procedure, and the variation operator. We pass the problem and initialization procedure to the super class (AbstractEvolutionaryAlgorithm). Also note that we pass a new NondominatedSortingPopulation to the super class, which is used to store the population. AbstractEvolutionaryAlgorithm will handle all of the messy details of initializing and maintaining the population.

Since we extended the AbstractEvolutionaryAlgorithm class, we only need to implement the iterate() method. The iterate() method, starting on line 22, describes one iteration of the algorithm. Depending on the type of algorithm, the iterate() method could perform a little or a lot of work. For example, in our RandomWalker example, each

iteration will generate one offspring. In a generational algorithm like NSGA-II, all members of the population are evolved every iteration. It is up to the designer of the algorithm to determine how much work is performed in the iterate() method. At the very least, however, at least one offspring should be generated each iteration.

We can test our RandomWalker class as follows:

```
package chapter5;
2
  import org.moeaframework.core.Algorithm;
  import org.moeaframework.core.Initialization;
  import org.moeaframework.core.NondominatedPopulation;
  import org.moeaframework.core.Problem;
  import org.moeaframework.core.Variation;
  import org.moeaframework.core.operator.RandomInitialization;
  import org.moeaframework.core.operator.real.PM;
  import chapter2.SchafferProblem;
  public class RunRandomWalker {
13
    public static void main(String[] args) {
14
      Problem problem = new SchafferProblem();
16
17
      Initialization initialization = new RandomInitialization(
           problem,
18
           100);
19
20
      Variation variation = new PM(
21
           1.0 / problem.getNumberOfVariables(),
22
23
           30.0);
24
      Algorithm algorithm = new RandomWalker(
25
           problem,
           initialization,
27
           variation);
29
      while (algorithm.getNumberOfEvaluations() < 10000) {</pre>
30
        algorithm.step();
31
32
33
      NondominatedPopulation result = algorithm.getResult();
34
35
36
```

MOEAFramework/book/chapter5/RunRandomWalker.java

5.4 Creating Service Providers

The MOEA Framework is designed to be flexible. It uses what's called a Service Provider Interface, or SPI, to dynamically load algorithms. This allows new optimization algorithms to be introduced within the MOEA Framework without modifying the MOEA Framework's code. To do this, we will create a new Java class called RandomWalkerProvider and extend the AlgorithmProvider interface. The contents of this file are shown below.

```
package chapter5;
  import java.util.Properties;
3
5
  import org.moeaframework.core.Algorithm;
  import org.moeaframework.core.Initialization;
  import org.moeaframework.core.Problem;
  import org.moeaframework.core.Variation;
  import org.moeaframework.core.operator.RandomInitialization;
  import org.moeaframework.core.spi.AlgorithmProvider;
  import org.moeaframework.core.spi.OperatorFactory;
11
  import org.moeaframework.util.TypedProperties;
  public class RandomWalkerProvider extends AlgorithmProvider {
14
    @Override
16
    public Algorithm getAlgorithm(String name, Properties properties, Problem
17
        problem) {
      if (name.equalsIgnoreCase("RandomWalker")) {
18
        // if the user requested the RandomWalker algorithm
19
        TypedProperties typedProperties = new TypedProperties(properties);
20
        // allow the user to customize the population size (default to 100)
22
        int populationSize = typedProperties.getInt("populationSize", 100);
24
        // initialize the algorithm with randomly-generated solutions
25
        Initialization initialization = new RandomInitialization(problem,
26
            populationSize);
        // use the operator factory to create a polynomial mutation operator
        Variation variation = OperatorFactory.getInstance().getVariation("pm",
29
            properties, problem);
30
        // construct and return the RandomWalker algorithm
31
        return new RandomWalker(problem, initialization, variation);
33
         // return null if the user requested a different algorithm
34
        return null;
35
36
37
38
39 }
```

MOEAFramework/book/chapter5/RandomWalkerProvider.java

If the user requests an optimization algorithm with the name "RandomWalker", the above class will create and return an instance of our RandomWalker class. Note that a properties object is passed to the getAlgorithm method. This properties object contains any custom properties set by the user, which allows the user to provide special parameters for the algorithm such as the population size.

Before we can use our new algorithm, we must register it with the MOEA Framework. The code below shows how to register and use our new optimization algorithm:

```
package chapter5;
2
  import org.moeaframework.Executor;
  import org.moeaframework.core.NondominatedPopulation;
  import org.moeaframework.core.spi.AlgorithmFactory;
  import chapter2.SchafferProblem;
7
  public class RunWithProvider {
9
    public static void main(String[] args) {
      AlgorithmFactory.getInstance().addProvider(new RandomWalkerProvider());
      NondominatedPopulation result = new Executor()
14
           .withAlgorithm("RandomWalker")
           .withProblemClass(SchafferProblem.class)
16
           .withMaxEvaluations(10000)
17
           .run();
18
19
20
21
```

MOEAF ramework/book/chapter 5/RunWith Provider.java

Great! Now we can test our new algorithm with the Analyzer class or inspect its runtime dynamics with the Instrumenter class. For example, below plots the convergence of the algorithm using generational distance:

```
package chapter5;

import org.moeaframework.Executor;
import org.moeaframework.Instrumenter;
import org.moeaframework.analysis.collector.Accumulator;
import org.moeaframework.analysis.plot.Plot;
import org.moeaframework.core.spi.AlgorithmFactory;

public class PlotDynamicsWithProvider {
    public static void main(String[] args) {
```

```
AlgorithmFactory.getInstance().addProvider(new RandomWalkerProvider());
12
       Instrumenter instrumenter = new Instrumenter()
14
           .withProblem("UF1")
           .withFrequency(100)
16
           .addAllowedPackage("chapter5")
17
           .attachGenerationalDistanceCollector();
18
19
      new Executor()
20
           .withSameProblemAs(instrumenter)
21
           .withAlgorithm("RandomWalker")
           .withMaxEvaluations (10000)
           .withInstrumenter(instrumenter)
24
           .run();
25
26
      Accumulator accumulator = instrumenter.getLastAccumulator();
27
28
      new Plot().add(accumulator).show();
29
30
31
32
```

MOEAFramework/book/chapter5/PlotDynamicsWithProvider.java

Note we must call addAllowedPackage ("chapter5") on line 17 to allow the instrumenter to work with code in our custom package.

5.5 Hyperheuristics

A hyperheuristic combines two or more other heuristics¹. The general idea is that a given heuristic works well one some problems but performs poorly on others. By utilizing more than one heuristic, we avoid being stuck using a poorly performing heuristic. There are many examples of this in the literature, including work on AMALGAM (Vrugt and Robinson, 2007; Vrugt et al., 2009) and Borg (Hadka and Reed, 2013). These hyperheuristics employ some learning mechanism to determine which heuristics perform better on a given problem. In this book, for simplicity, we won't use any learning mechanism and instead simply run both heuristics an equal number of times. In this example, we will combine NSGA-II and GDE3. This exercise is intended to demonstrate how using modular designs is beneficial. We will combine these two algorithms with a minimal amount of code:

```
package chapter5;

import org.moeaframework.algorithm.AbstractEvolutionaryAlgorithm;
import org.moeaframework.algorithm.GDE3;
import org.moeaframework.algorithm.NSGAII;
import org.moeaframework.core.Initialization;
```

¹Heuristic is another name for an MOEA

```
import org.moeaframework.core.NondominatedSortingPopulation;
  import org.moeaframework.core.Problem;
  import org.moeaframework.core.Variation;
import org.moeaframework.core.comparator.ChainedComparator;
import org.moeaframework.core.comparator.CrowdingComparator;
12 | import org.moeaframework.core.comparator.ParetoDominanceComparator;
  import org.moeaframework.core.operator.GAVariation;
13
14 import org.moeaframework.core.operator.TournamentSelection;
  import org.moeaframework.core.operator.real.DifferentialEvolution;
15
  import org.moeaframework.core.operator.real.DifferentialEvolutionSelection;
  import org.moeaframework.core.operator.real.PM;
17
  import org.moeaframework.core.operator.real.SBX;
19
  public class Hyperheuristic extends AbstractEvolutionaryAlgorithm {
20
21
    private NSGAII nsgaii;
22
    private GDE3 gde3;
24
25
    private int iteration;
26
    public Hyperheuristic (Problem problem, NondominatedSortingPopulation
28
        population,
        Initialization initialization) {
29
      super(problem, population, null, initialization);
30
      // define our NSGAII instance
      TournamentSelection selection = new TournamentSelection(2,
33
          new ChainedComparator(
               new ParetoDominanceComparator(),
35
               new CrowdingComparator()));
36
37
      Variation variation = new GAVariation (
          new SBX(1.0, 25.0),
39
          new PM(1.0 / problem.getNumberOfVariables(), 30.0));
40
41
      nsgaii = new NSGAII(
42
          problem,
43
          population,
44
          null, // no archive
45
          selection,
46
          variation,
47
          initialization);
48
49
      // define our GDE3 instance
50
      gde3 = new GDE3 (problem,
          population,
52
          new ParetoDominanceComparator(),
          new DifferentialEvolutionSelection(),
54
          new DifferentialEvolution (0.1, 0.9),
          initialization);
56
```

```
57
58
     @Override
59
     protected void iterate() {
60
       if (iteration % 2 == 0) {
61
62
         nsgaii.iterate();
       } else {
63
         gde3.iterate();
64
65
       numberOfEvaluations = nsqaii.getNumberOfEvaluations() +
67
           gde3.getNumberOfEvaluations();
68
       iteration++;
70
71
72
```

MOEAFramework/book/chapter5/Hyperheuristic.java

Here, we manually create the NSGA-II and GDE3 instance on lines 33-48 and 51-56, respectively. The primary difference is that we share the same population instance between both algorithm. This way, the algorithms are evolving the same population. In the iterate method on lines 60-71, we iterate between running NSGA-II and GDE3. Note in particular how we must update the numberOfEvaluations variable during each iteration. This is necessary since each algorithm may produce a different number of offspring and we must ensure the number of evaluations is tabulated correctly.

Lets create a custom provider:

```
package chapter5;
2
  import java.util.Properties;
3
  import org.moeaframework.core.Algorithm;
5
  import org.moeaframework.core.Initialization;
  import org.moeaframework.core.NondominatedSortingPopulation;
  import org.moeaframework.core.Problem;
  import org.moeaframework.core.operator.RandomInitialization;
  import org.moeaframework.core.spi.AlgorithmProvider;
  import org.moeaframework.util.TypedProperties;
  public class HyperheuristicProvider extends AlgorithmProvider {
14
    @Override
    public Algorithm getAlgorithm(String name, Properties properties, Problem
16
       problem) {
      if (name.equalsIgnoreCase("hyperheuristic")) {
17
        TypedProperties typedProperties = new TypedProperties(properties);
18
19
        int populationSize = typedProperties.getInt("populationSize", 100);
20
```

```
21
         Initialization initialization = new RandomInitialization(
22
             problem,
23
             populationSize);
24
         Algorithm algorithm = new Hyperheuristic(
26
             problem,
27
             new NondominatedSortingPopulation(),
28
             initialization);
30
         return algorithm;
         else {
         return null;
33
34
35
36
```

MOEAFramework/book/chapter5/HyperheuristicProvider.java

and test our new algorithm:

```
package chapter5;
2
  import org.moeaframework.Executor;
  import org.moeaframework.core.NondominatedPopulation;
  import org.moeaframework.core.spi.AlgorithmFactory;
  import chapter2.SchafferProblem;
  public class RunHyperheuristic {
8
9
    public static void main(String[] args) {
      AlgorithmFactory.getInstance().addProvider(new HyperheuristicProvider());
11
      NondominatedPopulation result = new Executor()
           .withAlgorithm("hyperheuristic")
14
           .withProblemClass(SchafferProblem.class)
15
           .withMaxEvaluations (10000)
16
           .run();
17
18
19
20
```

MOEAFramework/book/chapter5/RunHyperheuristic.java

5.6 Custom Types and Operators

The MOEA Framework provides several representations for decision variables, including real-values, integers, binary strings, permutations, and programs (expression trees). It is a

good practice to reuse these built-in operators when able. However, some situations require defining new types and operators, including:

- 1. Data can not be represented by a composite of built-in types
- 2. Provide better structure (grouping) of variables
- 3. Define custom variation operators

Suppose we needed to represent colors. A color is commonly represented on a computer as a composite of three primary colors: red, green, and blue. The value for each primary color typically ranges between 0 and 255, where 0 indicates absence of the primary color and 255 indicates complete saturation. We typically express a color as a tuple of three values, e.g. (red, green, blue). Some common colors are shown below:

```
• Black - (0, 0, 0)
```

- Red (255, 0, 0)
- Green (0, 255, 0)
- Blue (0, 0, 255)
- Yellow (255, 255, 0)
- White (255, 255, 255)

We could choose to represent colors as three separate real or integer values. However, we can provide better structure and enable the use of custom operators by defining a new type. First, we create the Color type by implementing the Variable interface. A variable requires two methods: copy() and randomize(). The copy() method is responsible for creating an exact copy of a color (which is used to create new offspring solutions) and randomize() creates a new random color (which is used to create the initial population). Our Color class is shown below:

```
package chapter5;
import org.moeaframework.core.PRNG;
import org.moeaframework.core.Variable;

public class Color implements Variable {
   private static final long serialVersionUID = 8461741347578471248L;
   private int r;
   private int g;
```

```
private int b;
14
15
     public Color() {
16
       super();
17
18
19
20
    public Color(int r, int g, int b) {
       this();
21
       this.r = r;
22
       this.g = g;
23
       this.b = b;
24
25
26
    protected int getR() {
27
       return r;
28
29
30
31
    protected void setR(int r) {
       this.r = r;
32
33
34
    protected int getG() {
35
       return g;
36
37
38
    protected void setG(int g) {
39
      this.g = g;
40
41
42
     protected int getB() {
43
      return b;
44
45
46
    protected void setB(int b) {
47
48
       this.b = b;
49
50
     @Override
51
52
    public Variable copy() {
       return new Color(r, g, b);
54
     @Override
56
    public void randomize() {
57
      r = PRNG.nextInt(256);
58
59
      g = PRNG.nextInt(256);
60
      b = PRNG.nextInt(256);
61
62
     @Override
63
    public String toString() {
64
```

```
return "Color [r=" + r + ", g=" + g + ", b=" + b + "]";

66 }

87 }
```

MOEAFramework/book/chapter5/Color.java

Second, we need to define any variation operators for this type. The variation operators describe how to generate new candidate offspring during search. For simplicity, we will define a mutation operator that accepts a single parent, copies the parent, randomly perturbs the color in the copy, and returns the result. Be sure to always create copies of parents to avoid modifying the original parent solution.

```
package chapter5;
  import org.moeaframework.core.PRNG;
3
  import org.moeaframework.core.Solution;
  import org.moeaframework.core.Variation;
  public class ColorMutation implements Variation {
7
8
    @Override
9
    public int getArity() {
       return 1;
13
    @Override
14
    public Solution[] evolve(Solution[] parents) {
       Solution result = parents[0].copy();
16
17
       for (int i = 0; i < result.getNumberOfVariables(); i++) {</pre>
18
         if (result.getVariable(i) instanceof Color) {
19
           Color color = (Color)result.getVariable(i);
20
           color.setR(Math.min(255, Math.max(0, color.getR() + PRNG.nextInt(-5,
21
              5))));
22
           color.setG(Math.min(255, Math.max(0, color.getG() + PRNG.nextInt(-5,
              5))));
           color.setB(Math.min(255, Math.max(0, color.getB() + PRNG.nextInt(-5,
23
              5))));
24
25
26
      return new Solution[] { result };
27
28
29
30
  };
```

MOEAFramework/book/chapter5/ColorMutation.java

Finally, we create an operator provider for this new type. The operator provider is responsible for specifying which variation operators are appropriate for a given problem. In

this example, we check to see if the problem type contains any Color variables, and if so, instruct the MOEA Framework to use our variation operator.

```
package chapter5;
2
  import java.util.Properties;
3
  import org.moeaframework.core.Problem;
  import org.moeaframework.core.Solution;
  import org.moeaframework.core.Variation;
  import org.moeaframework.core.spi.OperatorProvider;
  public class ColorProvider extends OperatorProvider {
10
11
    @Override
    public String getMutationHint(Problem problem) {
13
       Solution solution = problem.newSolution();
14
      for (int i = 0; i < problem.getNumberOfVariables(); i++) {</pre>
16
17
         if (solution.getVariable(i) instanceof Color) {
           return "colormutation";
19
20
      return null;
22
23
24
    @Override
25
    public String getVariationHint(Problem problem) {
26
      return getMutationHint(problem);
27
28
29
    @Override
30
    public Variation getVariation (String name, Properties properties,
31
        Problem problem) {
32
      if (name.equalsIgnoreCase("colormutation")) {
33
         return new ColorMutation();
34
35
36
       return null;
38
39
40
```

MOEAFramework/book/chapter5/ColorProvider.java

At this point, we have defined our new decision variable type and the appropriate operators. We can now define a problem using this type. The problem we will solve is very simple: find N colors that, when mixed, produce a color closest to the target, assuming the background or base color is white (255, 255, 255).

```
package chapter5;
  import org.moeaframework.core.Solution;
  import org.moeaframework.problem.AbstractProblem;
  public class ColorBlenderProblem extends AbstractProblem {
5
6
    private final Color target;
7
    public ColorBlenderProblem(int N, Color target) {
9
      super(N, 1);
      this.target = target;
14
    public Color blend(Color c1, Color c2, double ratio) {
         double invRatio = 1.0 - ratio;
         int r = (int)((c1.getR() * invRatio) + (c2.getR() * ratio));
16
         int g = (int)((c1.getG() * invRatio) + (c2.getG() * ratio));
17
         int b = (int)((c1.getB() * invRatio) + (c2.getB() * ratio));
18
19
         return new Color(r, g, b);
20
21
22
    @Override
23
    public void evaluate(Solution solution) {
24
      Color result = new Color(255, 255, 255);
25
26
       for (int i = 0; i < numberOfVariables; i++) {</pre>
         result = blend(result, (Color)solution.getVariable(i), 0.5);
28
29
30
31
       int diff = Math.abs(result.getR()-target.getR()) +
           Math.abs(result.getG()-target.getG()) +
32
           Math.abs(result.getB()-target.getB());
33
34
      solution.setObjective(0, diff);
35
    }
36
37
    @Override
38
    public Solution newSolution() {
39
       Solution solution = new Solution(numberOfVariables, numberOfObjectives);
40
41
       for (int i = 0; i < numberOfVariables; i++) {</pre>
42
         solution.setVariable(i, new Color());
43
44
45
      return solution;
46
47
48
49
```

MOEAFramework/book/chapter5/ColorBlenderProblem.java

Note how the constructor on line 9 accepts two arguments: N, the number of colors to combine, and target, the target color. This will allow us to run this problem with different inputs, as shown below.

```
package chapter5;
  import org.moeaframework.Executor;
  import org.moeaframework.core.NondominatedPopulation;
  import org.moeaframework.core.Solution;
  import org.moeaframework.core.spi.OperatorFactory;
  public class RunColorBlenderProblem {
7
8
    public static void main(String[] args) {
9
      OperatorFactory.getInstance().addProvider(new ColorProvider());
12
      NondominatedPopulation result = new Executor()
           .withProblemClass(ColorBlenderProblem.class, 3, new Color(127, 127,
           .withAlgorithm("NSGAII")
14
           .withMaxEvaluations(10000)
           .distributeOnAllCores()
16
17
           .run();
18
      for (Solution solution : result) {
19
        System.out.print(solution.getObjective(0));
20
21
        for (int i = 0; i < solution.getNumberOfVariables(); i++) {</pre>
           System.out.print(" ");
           System.out.print(solution.getVariable(i));
25
27
28
29
```

MOEAFramework/book/chapter5/RunColorBlenderProblem.java

As before, we must register our custom operator provider on line 11. Once registered, we can optimize the problem. Also observe on line 13 how we pass the arguments when calling withProblemClass. The result is shown below:

```
0.0 Color [r=148, g=20, b=132] Color [r=158, g=252, b=127] Color [r=76, g=61, b=94]
```

The three colors (148, 20, 132), (158, 252, 127) and (76, 61, 94) combine on a white background to produce the target color (127, 127, 127) exactly. You will likely see different results if running this example since there are many color combinations that produce the same result.

5.7 Learning the API

You may be wondering how we know which classes to implement when defining custom components. We make available online the entire application programming interface (API) for the MOEA Framework. The API describes in detail every class, function, and argument in the code. You can find the API online at http://moeaframework.org/javadoc/index.html.

Chapter 6

The Diagnostic Tool

We'll take a brief break from code and turn our attention to the MOEA Framework's diagnostic tool. The diagnostic tool provides a graphical interface to quickly run and analyze MOEAs on a test problems. We have seen many of the functions provided by the diagnostic tool in the preceding chapters, but the diagnostic tool provides an easy-to-use point-and-click interface for accessing these features.

6.1 Using the Diagnostic Tool

To run the diagnostic tool, navigate to the MOEA Framework folder on your computer and double-click the launch-diagnostic-tool.bat file. You can manually run the diagnostic tool with the following command:

```
java -Djava.ext.dirs=lib org.moeaframework.analysis.diagnostics.LaunchDiagnosticTool
```

Figure 6.1 provides a screenshot of the diagnostic tool window. This window is composed of the following sections:

- 1. The configuration panel. This panel lets you select the algorithm, problem, number of repetitions (seeds), and maximum number of function evaluations (NFE).
- 2. The execution panel. Clicking run will execute the algorithm as configured in the configuration panel. Two progress bars display the individual run progress and the total progress for all seeds. Any in-progress runs can be canceled.
- 3. The displayed results table. This table displays the completed runs. The entries which are selected/highlighted are displayed in the charts. You can click an individual line to show the data for just that entry, click while holding the Alt key to select multiple entries, or click the Select All button to select all entries.

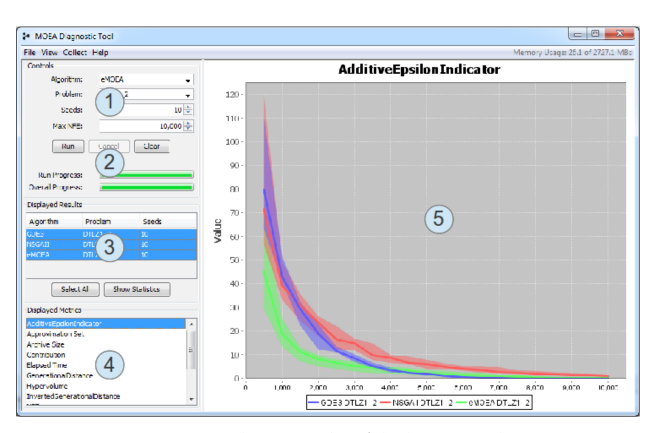


Figure 6.1: The main window of the diagnostic tool.

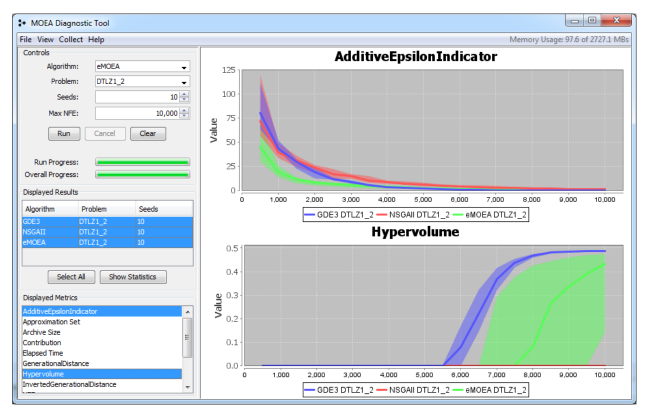


Figure 6.2: Screenshot of the diagnostic tool displaying two side-by-side metrics. You can select as many metrics to display by holding down the Alt key and clicking a row in the displayed metrics table.

- 4. The displayed metrics table. Similar to the displayed results table, the selected metrics are displayed in the charts. You can select one metric or multiple metrics by holding the Alt key while clicking.
- 5. The actual charts. A chart will be generated for each selected metric. Thus, if two metrics are selected, then two charts will be displayed side-by-side. See Figure 6.2 for an example.

Some algorithms do not provide certain metrics. When selecting a specific metric, only those algorithms that provide that metric will be displayed in the chart.

Quantile Plots vs Individual Traces

By default, the charts displayed in the diagnostic tool show the statistical 25%, 50% and 75% quantiles. The 50% quantile is the thick colored line, and the 25% and 75% quantiles are depicted by the colored area. This quantile view allows you to quickly distinguish the performance between multiple algorithms, particularly when there is significant overlap between two or more algorithms.

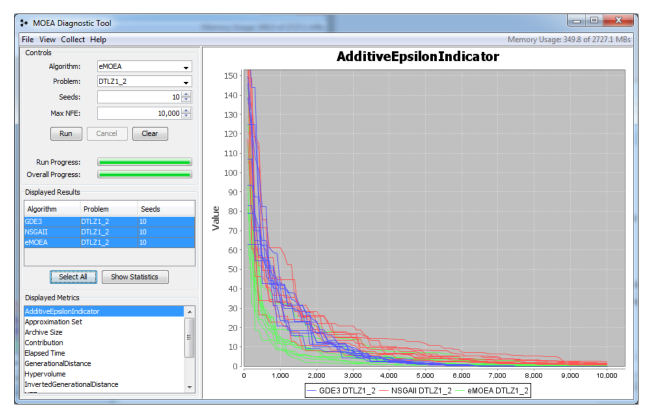


Figure 6.3: Screenshot of the diagnostic tool displaying the individual traces rather than the quantile view. The individual traces provide access to the raw data, but the quantile view is often easier to interpret.

You can alternatively view the raw, individual traces by selecting 'Show Individual Traces' in the View menu. Each colored line represents one seed. Figure 6.3 provides an example of plots showing individual traces. You can always switch back to the quantile view using the View menu.

Viewing Approximation Set Dynamics

Another powerful feature of the diagnostic tool is the visualization of approximation set dynamics. The approximation set dynamics show how the algorithm's result (its approximation set) evolved throughout the run. To view the approximation set dynamics, right-click on one of the rows in the displayed results table. A menu will appear with the option to show the approximation set. A window similar to Figure 6.4 will appear.

This window displays the following items:

1. The approximation set plot. This plot can only show two dimensions. If available, the reference set for the problem will be shown as black points. All other points are the solutions produced by the algorithm. Different seeds are displayed in different colors.

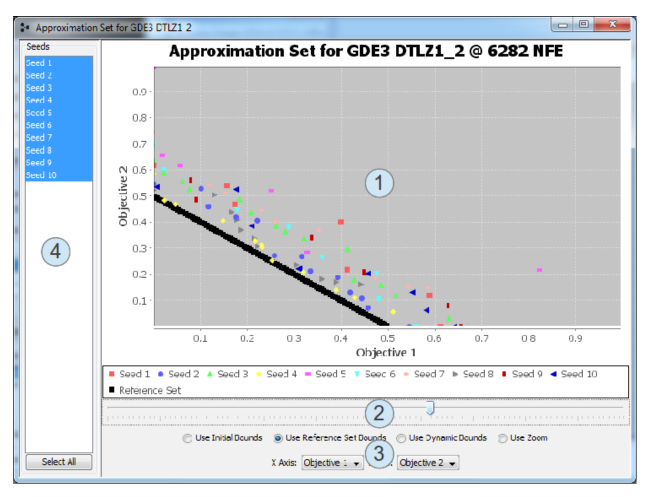


Figure 6.4: Screenshot of the approximation set viewer. This allows you to view the approximation set at any point in the algorithm's execution.

- 2. The evolution slider. Dragging the slider to the left or right will show the approximation set from earlier or later in the evolution.
- 3. The display controls. These controls let you adjust how the data is displayed. Each of the radio buttons switches between different scaling options. The most common option is 'Use Reference Set Bounds', which scales the plot so that the reference set fills most of the displayed region.
- 4. The displayed seeds table. By default, the approximation sets for all seeds are displayed and are distinguished by color. You can also downselect to display one or a selected group of seeds by selecting entries in this table. Multiple entries can be selected by holding the Alt key while clicking.

You can manually zoom to any portion in these plots (both in the approximation set viewer and the plots in the main diagnostic tool window) by positioning the cursor at the top-left corner of the zoom region, pressing and holding down the left-mouse button, dragging the cursor to the bottom-right corner of the zoom region, and releasing the left-mouse button. You can reset the zoom by pressing and holding the left-mouse button, dragging the cursor to the top-left portion of the plot, and releasing the left-mouse button.

Statistical Results

The diagnostic tool also allows you to exercise the statistical testing tools provided by the MOEA Framework with the click of a button. If you have two or more entries selected in the displayed results table, the 'Show Statistics' button will become enabled. The show statistics button also requires each of the selected entries to use the same problem. The button will remain disabled unless this condition is satisfied. If the button is disabled, please ensure you have two or more rows selected and all selected entries are using the same problem. Figure 6.5 shows the example output from clicking this button.

Improving Performance and Memory Efficiency

By default, the diagnostic tool collects and displays all available data. If you know ahead of time that certain pieces of data are not needed for your experiments, you can often increase the performance and memory efficiency of the program by disabling unneeded data. You can enable or disable the collection of data by checking or unchecking the appropriate item in the Collect menu.

6.2 Adding Custom Algorithms

What makes the diagnostic tool particularly powerful is that it can incorporate custom algorithms and problems. Lets take, for example, the RandomWalker and Hyperheuristic algorithms we created in the previous chapter. We can incorporate these algorithms into the diagnostic tool with the code below:

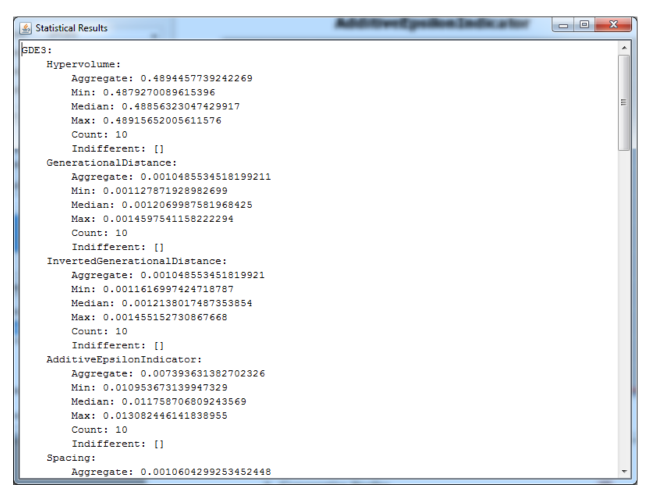


Figure 6.5: Screenshot of the statistics output by the diagnostic tool.

```
package chapter6;
2
3
  import org.moeaframework.analysis.diagnostics.LaunchDiagnosticTool;
  import org.moeaframework.core.Settings;
  import org.moeaframework.core.spi.AlgorithmFactory;
  import chapter5.HyperheuristicProvider;
7
  import chapter5.RandomWalkerProvider;
  public class RunDiagnosticTool {
    public static void main(String[] args) throws Exception {
      AlgorithmFactory.getInstance().addProvider(new RandomWalkerProvider());
14
      AlgorithmFactory.getInstance().addProvider(new HyperheuristicProvider());
      Settings.PROPERTIES.setString(
16
           Settings.KEY_DIAGNOSTIC_TOOL_ALGORITHMS,
17
           "RandomWalker, Hyperheuristic, NSGAII, GDE3");
18
19
      LaunchDiagnosticTool.main(args);
20
21
22
```

MOEAFramework/book/chapter6/RunDiagnosticTool.java

As before, we must register our custom algorithm provides with the MOEA Framework on lines 13 and 14. Lines 16-22 set two properties to configure the diagnostic tool. The first property defines which algorithms appear in the tool. We add our two new algorithms to this list. The second property allows our custom algorithms to be instrumented. Without specifying the allowed packages property, the diagnostic tool will be unable to collect any runtime data for our two custom algorithms. Finally, on line 24, we launch the diagnostic tool window.

Now we want to run each of the four algorithms on a test problem so we can view their plots. We will use the UF1 problem, but feel free to test with other problems. Select each algorithm from the drop-down and click the Run button.

Figure 6.6 shows the resulting plot. Here, we've selected inverted generational distance. As expected, the RandomWalker does not perform very well — after all, it's just taking random steps. The other algorithms perform very similarly. Using the mouse button, we can zoom in, as shown in Figure 6.7. Now we can see a difference between the algorithms. NSGA-II in green and GDE3 and red show differences (the difference is slight, but it is noticeable). Our hyperheuristic, which combines NSGA-II and GDE3, matches the performance of the better individual algorithm, GDE3. Even with our simple hyperheuristic, we can see that the hyperheuristic was able to perform at least as good as its component heuristics.

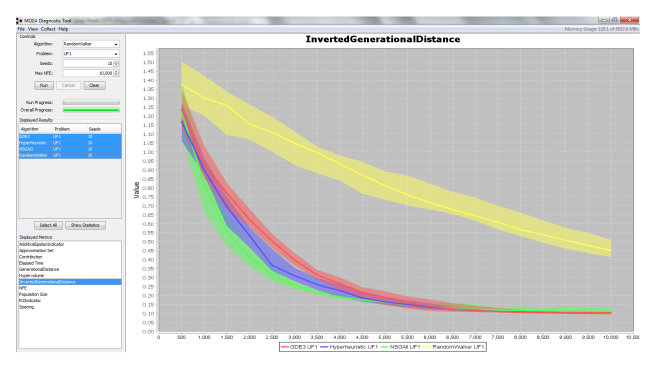


Figure 6.6: Comparison of our two custom algorithms with NSGA-II and GDE3

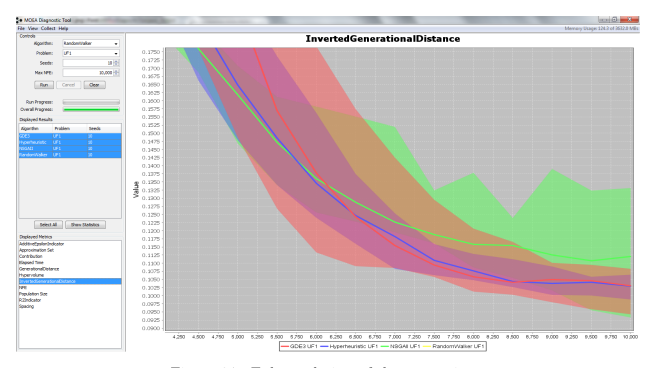


Figure 6.7: Enhanced view of the comparison.

6.3 Adding Custom Problems

Similar to how we added custom algorithms by registering a provider with the MOEA Framework, user-defined problems are defined using the ProblemProvider. For example, below we create a problem provider for the Schaffer problem we created in Chapter 2:

```
package chapter6;
2
  import java.io.File;
3
  import java.io.IOException;
5
  import org.moeaframework.core.FrameworkException;
  import org.moeaframework.core.NondominatedPopulation;
  import org.moeaframework.core.PopulationIO;
  import org.moeaframework.core.Problem;
  import org.moeaframework.core.spi.ProblemProvider;
11
12
  import chapter2.SchafferProblem;
  public class SchafferProblemProvider extends ProblemProvider {
14
    @Override
16
17
    public Problem getProblem(String name) {
      if (name.equalsIgnoreCase("MySchafferProblem")) {
18
         return new SchafferProblem();
19
        else {
20
         return null;
22
    }
23
24
    @Override
25
    public NondominatedPopulation getReferenceSet(String name) {
26
      if (name.equalsIgnoreCase("MySchafferProblem")) {
         try {
28
           return new NondominatedPopulation (
               PopulationIO.readObjectives(new File("./pf/Schaffer.pf")));
30
         } catch (IOException e) {
           throw new FrameworkException(e);
32
33
34
      } else {
35
         return null;
36
37
38
```

MOEAFramework/book/chapter6/SchafferProblemProvider.java

A ProblemProvider must define two methods: getProblem and getReferenceSet. getProblem returns an instance of the problem if the name matches. getReferenceSet returns the reference set for the problem. Both methods

must return **null** if the name does not match any supported problem.

With the problem provider found, we can register it with the MOEA Framework and launch the diagnostic tool:

```
package chapter6;
2
  import org.moeaframework.analysis.diagnostics.LaunchDiagnosticTool;
  import org.moeaframework.core.Settings;
  import org.moeaframework.core.spi.AlgorithmFactory;
  import org.moeaframework.core.spi.ProblemFactory;
  import chapter5.HyperheuristicProvider;
8
  import chapter5.RandomWalkerProvider;
9
  public class CustomProblem {
12
    public static void main(String[] args) throws Exception {
      AlgorithmFactory.getInstance().addProvider(new RandomWalkerProvider());
14
      AlgorithmFactory.getInstance().addProvider(new HyperheuristicProvider());
16
      ProblemFactory.getInstance().addProvider(new SchafferProblemProvider());
17
18
      Settings.PROPERTIES.setString(
19
           Settings.KEY_DIAGNOSTIC_TOOL_ALGORITHMS,
20
           "RandomWalker, Hyperheuristic, NSGAII, GDE3");
22
      Settings.PROPERTIES.setString(
           Settings.KEY_DIAGNOSTIC_TOOL_PROBLEMS,
24
           "MySchafferProblem");
25
26
      LaunchDiagnosticTool.main(args);
28
29
30
```

MOEAFramework/book/chapter6/CustomProblem.java

Chapter 7

Subsets, Permutations, and Programs

We have seen thus far real-valued and bit string encoding problems. The MOEA Framework supports three additional, commonly-used representations: subsets, permutations, and programs. Subsets are similar to bit string encodings, except they represent situations where only a fixed number of items can be selected (whereas bit strings allow a variable number of items). Permutations are often used in problems where ordering is important. Programs are used when one must construct code, rule systems, or decision trees to solve a task. This chapter discusses these representations.

7.1 Subsets

Previously we saw with the bit string representation how we can solve the Knapsack problem. Recall that in the Knapsack problem we are tasked with identifying the items to place in a Knapsack to maximize value constrained only by the capacity of the Knapsack. By using the bit string (binary) representation, we allowed the optimization algorithm to select anywhere from 0 up to N items.

Subsets are good for a similar but slightly different class of problems. Instead of allowing one to select anywhere from 0 up to N items, what if they must select exactly k items. When the subset is a fixed size, we use the Subset class. Subsets are similar to selection without replacement. It will pick k items from a set containing N items without repeating any of the selected items. (Tip: if you need selection with replacement, where items can be selected more than once, just use k integer variables).

To demonstrate subsets, we will use a slight twist on the classic subset-sum problem, which we call the fixed-size subset-sum problem. In the fixed-size subset-sum problem, we are given N integers, both positive and negative. The goal is to find a subset of the integers of size k that sums to 0. For example, if we have the numbers -5, -3, -1, 2, 4, 6 and k = 4, we could pick -5, -3, 2, 6 since -5 + -3 + 2 + 6 = 0.

The full code for the fixed-size subset-sum problem is shown below:

```
import org.moeaframework.core.Solution;
  import org.moeaframework.core.variable.EncodingUtils;
4
  import org.moeaframework.core.variable.Subset;
  import org.moeaframework.problem.AbstractProblem;
  public class FixedSubsetSumProblem extends AbstractProblem {
8
9
    private int k;
    private int[] values;
    public FixedSubsetSumProblem(int k, int[] values) {
14
       super(1, 1);
      this.k = k;
      this.values = values;
17
18
19
    @Override
20
    public void evaluate(Solution solution) {
21
       int[] subset = EncodingUtils.getSubset(solution.getVariable(0));
23
       int sum = 0;
24
       for (int i = 0; i < subset.length; i++) {</pre>
25
         sum += values[subset[i]];
26
27
28
       solution.setObjective(0, Math.abs(sum));
29
30
31
    @Override
32
    public Solution newSolution() {
33
       Solution solution = new Solution(1, 1);
      solution.setVariable(0, new Subset(k, values.length));
35
      return solution;
37
38
39
```

MOEAF ramework/book/chapter 7/Fixed Subset Sum Problem. java

We construct the subset with **new** Subset (k, values.length) to indicate we want to form subsets of size k from a set of size values.length. Note that the subset does not actually store the values. Instead, it stores the indices of the selected items. When evaluating the solution, we must be careful to convert from the indices in the subset to the original values using values[subset[i]].

When our problem implemented, we can then solve it:

```
package chapter7;
import java.util.Arrays;
```

```
import org.moeaframework.Executor;
5
  import org.moeaframework.core.NondominatedPopulation;
6
  import org.moeaframework.core.Solution;
  import org.moeaframework.core.variable.EncodingUtils;
10
  public class SolveFixedSubsetSumProblem {
11
    private static int[] values = { -1, 1, 3, 5, 9, -4, -2, -8, 17, 24, 18, -16,
         -20 };
    public static void main(String[] args) {
14
      NondominatedPopulation result = new Executor()
           .withAlgorithm("NSGAII")
16
           .withProblemClass(FixedSubsetSumProblem.class, 6, values)
17
           .withMaxEvaluations(10000)
           .run();
19
20
21
22
      for (Solution solution : result) {
        int[] subset = EncodingUtils.getSubset(solution.getVariable(0));
24
        System.out.println(Arrays.toString(toValues(subset, values)) +
25
             " => " + solution.getObjective(0));
26
27
28
29
    public static int[] toValues(int[] subset, int[] values) {
30
      int[] result = new int[subset.length];
32
      for (int i = 0; i < subset.length; i++) {</pre>
33
        result[i] = values[subset[i]];
34
36
      return result;
37
38
39
40
```

MOEAFramework/book/chapter7/SolveFixedSubsetSumProblem.java

You should see output similar to the following:

```
[9, 3, -20, -8, 17, -1] \Rightarrow 0.0
```

7.2 Permutations

Permutations are commonly seen in problems involving ordering. For example, job scheduling often uses permutations to specify the scheduling priority for jobs. Or, in the famous travelling salesman problem¹, we seek the shortest route that visits all cities. We could use a permutation to represent the path. We will demonstrate the use of permutations in this section by constructing a simple solver for traveling salesman problems.

The full code for the travelling salesman problem is provided below.

```
package chapter7;
  import org.moeaframework.core.Solution;
3
  import org.moeaframework.core.variable.EncodingUtils;
  import org.moeaframework.problem.AbstractProblem;
  public class TravellingSalesmanProblem extends AbstractProblem {
7
    private double[][] cities;
9
    public TravellingSalesmanProblem(double[][] cities) {
12
      super(1, 1);
      this.cities = cities;
14
    @Override
16
17
    public Solution newSolution() {
      Solution solution = new Solution(1, 1);
18
      solution.setVariable(0, EncodingUtils.newPermutation(cities.length));
19
      return solution;
20
    }
21
    private double distance(int i, int j) {
23
      return Math.sqrt(
24
           Math.pow(cities[i][0]-cities[j][0], 2.0) +
           Math.pow(cities[i][1]-cities[j][1], 2.0));
26
27
2.8
    @Override
29
    public void evaluate(Solution solution) {
30
      int[] path = EncodingUtils.getPermutation(solution.getVariable(0));
31
      double totalDistance = 0.0;
32
33
      for (int i = 0; i < path.length; i++) {</pre>
34
         totalDistance += distance(path[i], path[(i+1) % path.length]);
35
36
37
      solution.setObjective(0, totalDistance);
38
39
```

¹https://en.wikipedia.org/wiki/Travelling_salesman_problem

40

MOEAFramework/book/chapter7/TravellingSalesmanProblem.java

On lines 11-14 we construct the problem. For flexibility, we'll make the list of cities an argument. The cities are stored in a Nx2 array storing the x and y coordinates of N cities. The newSolution method on lines 17-21 shows how to construct a solution. A permutation is stored in a single decision variable. Lastly, we define the evaluate method on lines 30-39. First, we extract the permutation on line 31. Second, on lines 32-36, we sum up the total distance travelled by the salesman. Finally, on line 38, we set the objective value.

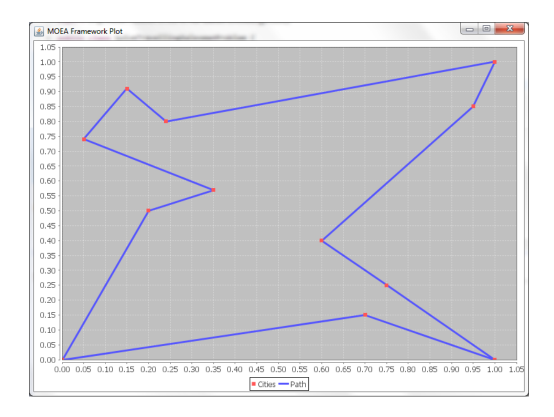
The code for optimizing this problem is shown below.

```
package chapter7;
  import org.moeaframework.Executor;
3
  import org.moeaframework.analysis.plot.Plot;
  import org.moeaframework.core.NondominatedPopulation;
  import org.moeaframework.core.variable.EncodingUtils;
6
  public class SolveTravellingSalesmanProblem {
    public static double[][] CITIES = {
         { 0.0,
                 0.0
                      },
                 1.0
         { 1.0,
                      },
          0.75, 0.25},
13
          1.0,
                 0.0
14
          0.95, 0.85 },
          0.2,
                 0.5
         {
16
17
          0.7,
                 0.15 },
                 0.4
          0.6,
18
          0.15, 0.91},
19
          0.24, 0.8
20
          0.05, 0.74},
21
         { 0.35, 0.57 }
22
23
    };
24
    public static void main(String[] args) {
25
      NondominatedPopulation result = new Executor()
26
           .withAlgorithm("NSGAII")
27
           .withProblemClass(TravellingSalesmanProblem.class, CITIES)
           .withMaxEvaluations(10000)
29
           .run();
      int[] bestPath = EncodingUtils.getPermutation(result.get(0).getVariable(0)
          );
33
      new Plot()
34
           .scatter("Cities", getCoordinate(0), getCoordinate(1))
35
           .line("Path", getCoordinate(0, bestPath), getCoordinate(1, bestPath))
36
```

```
.show();
38
39
    public static double[] getCoordinate(int index) {
40
       double[] result = new double[CITIES.length];
41
42
43
       for (int i = 0; i < CITIES.length; i++) {</pre>
         result[i] = CITIES[i][index];
44
45
46
       return result;
47
48
49
    public static double[] getCoordinate(int index, int[] path) {
50
       double[] result = new double[path.length+1];
51
52
       for (int i = 0; i < path.length; i++) {</pre>
53
         result[i] = CITIES[path[i]][index];
54
55
56
       result[path.length] = CITIES[path[0]][index];
58
       return result;
60
61
62
```

MOEAFramework/book/chapter7/SolveTravellingSalesmanProblem.java

We define two helper methods, both called getCoordinate, for extracting the coordinates of the cities for plotting. The resulting plot looks like:



7.3 Programs

My problems require developing computer code, rule systems, or decision trees. One example would be developing a robot to navigate an environment and avoid hitting obstacles. Available to the robots is input from various sensors which it uses to control steering and speed. We won't be so bold in this book to build such a robot, but we will explore here two problems. The first is a symbolic regression problem, where we attempt to find an expression to fit a set of points. The second is the ant problem, where we program a robotic ant to find food in a virtual world.

7.3.1 Type-based Rule System

To accomplish these tasks, the MOEA Framework contains a simple but extensible LISP-like programming language. The first step towards evolving programs is to define the rules (i.e., the syntax and semantics). When defining the rules, two important properties should be kept in mind: *closure* and *sufficiency*.

The closure property requires all program element to be able to accept as arguments any value and data type that could possibly be returned by any other function or terminal. All programs generated or evolved by the MOEA Framework are strongly typed. No program produced by the MOEA Framework will pass an argument to a function that is an incorrect type. Furthermore, all functions guard against invalid inputs. For example, the log of a

negative number is undefined. Rather then causing an error, the log method will guard itself and return 0.0. This allows the rest of the calculation to continue unabated. With these two behaviors built into the MOEA Framework, the closure property is guaranteed.

The sufficiency property states that the rule set must contain all the necessary functions and terminals necessary to produce a solution to the problem. Ensuring this property holds is more challenging as it will depend on the problem domain. For instance, the operators And, Or and Not are sufficient to produce all boolean expressions. It may not be so obvious in other problem domains which program elements are required to ensure sufficiency. Additionally, it is often helpful to restrict the rule set to those program elements that are sufficient, thus reducing the search space for the evolutionary algorithm.

The MOEA Framework comes packaged with over 45 pre-defined program elements for defining constants, variables, arithmetic operators, control structures, functions, etc. These program elements are listed below.

Operator	Description
Abs	Calculates the absolute value of a number
Acos	Calculates the arc cosine of a number
Acosh	Calculates the hyperbolic arc cosine of a number
Add	Adds two numbers
And	Calculates the logical AND of two boolean values
Asin	Calculates the arc sine of a number
Asinh	Calculates the hyperbolic arc sine of a number
Atan	Calculates the arc tangent of a number
Atanh	Calculates the hyperbolic arc tangent of a number
Call	Calls a named function
Ceil	Calculates the smallest integer value that is greater than or equal
	to a number
Constant	Defines a constant value
Cos	Calculates the trigonometric cosine of an angle specified in radians
Cosh	Calculates the hyperbolic cosine of a number
Define	Defines a callable, named function
Divide	Divides two numbers
Equals	Compares two numbers for equality
Exp	Calculates the result of Euler's number e raised to the power of a number
Floor	Calculates the largest integer value that is less than or equal to a number
For	Executes an expression for a given number of iterations
Get	Reads the value stored in a named variable within the current
300	scope
GreaterThan	Compares if one number is greater than another
Greater Than Or Equal	Compares if a number is greater than or equal to another
STEARCH THAILOTEQUAI	compares it a number is greater than or equal to another

If Else Executes one of two expressions depending on the result of a

boolean expression

Lambda Defines an immutable, anonymous function LessThan Compares if a number is less than another

LessThanOrEqual Compares if a number is less than or equal to another

Log The node for calculating the natural logarithm of a number

Log10 Calculates the base-10 logarithm of a number

Max Calculates the maximum value of the two arguments
Min Calculates the minimum value of the two arguments
Modulus Calculates the modulus, or remainder, of two numbers

Multiply Multiplies two numbers NOP Defines an empty expression

Not Calculates the logical NOT of a boolean value
Or Calculates the logical OR of two boolean values
Power Calculates the power of a base number and exponent

Round Rounds a number to the nearest integer

Sequence Executes two or more expressions in sequence

Set Assigns the value of a named variable within the current scope

Sign Calculates the sign of a number

Sin Calculates the trigonometric sine of an angle specified in radians

Sinh Calculates the hyperbolic sine of a number

Square Calculates the square of a number SquareRoot Calculates the square root of a number

Subtracts two numbers

Tan Calculates the trigonometric tangent of an angle specified in radi-

ans

Tanh Calculates the hyperbolic tangent of a number Truncate Truncates, or bounds, a number within a range

While Repeatedly executes an expression while a condition, a boolean

expression, remains true

Lets consider the problem of symbolic regression. Suppose we have a bunch of data points, say [(0,0),(1,2),(2,4),(3,6),(4,8),(5,10)]. The goal of symbolic regression is to find a mathematical expression that fits the data points. In this example, the function $f(x) = x^2$ would work. There is typically no single, unique function that runs through all points, thus creating issues like overfitting the data. However, this topic is outside the scope of this manual.

For this problem, we could construct a rule set using several arithmetic operators. One terminal is included, the variable x. We will assign this variable later when evaluating the program. The last setting required is the return type of the program. In this case, the program will return a number.

```
rules.add(new Add());
       rules.add(new Multiply());
3
       rules.add(new Subtract());
4
       rules.add(new Divide());
5
       rules.add(new Sin());
6
       rules.add(new Cos());
8
       rules.add(new Exp());
       rules.add(new Log());
9
       rules.add(new Get(Number.class, "x"));
10
       rules.setReturnType(Number.class);
```

7.3.2 Defining the Problem

With the rules defined, we can now define the problem. Programs require only a single decision variable of type Program. For example, the newSolution method would appear as follows:

```
public Solution newSolution() {
    Solution solution = new Solution(1, 1);
    solution.setVariable(0, new Program(rules));
    return solution;
}
```

Inside the evaluate method, we must execute the program. Each program executes inside an environment. The environment holds all of the variables and other identifiers that the program can access throughout its execution. Since we previously defined the variable x (with the Get node), we want to initialize the value of x in the environment. Once the environment is initialized, we can evaluate the program. Since we set the return type to be a number in the rule set, we cast the output from the program's evaluation to a number.

```
public void evaluate(Solution solution) {
    Program program = (Program)solution.getVariable(0);

Environment environment = new Environment();
    environment.set("x", 15);

double result = (Number)program.evaluate(environment)).doubleValue();
}
```

Returning to our symbolic regression example, we can now create the problem definition. Our rules will be a set of mathematical operators. Our evaluate method will compare the original function to the approximated function. We will compare the two functions by sampling the values at 100 different locations and compute the mean square error. By

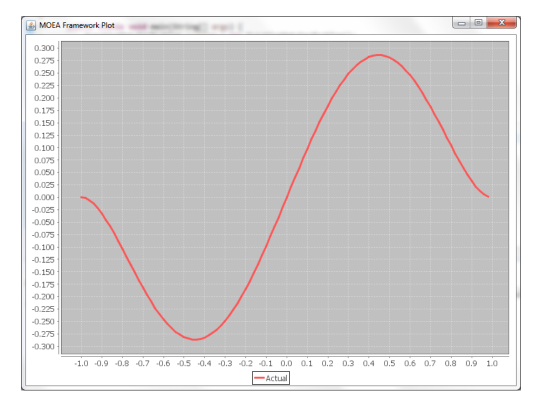
minimizing the mean square error, we minimize the difference between the two functions at the sampled points. The full problem class is shown below:

```
package chapter7;
2
  import org.moeaframework.core.Solution;
3
  import org.moeaframework.core.variable.Program;
  import org.moeaframework.problem.AbstractProblem;
  import org.moeaframework.util.tree.Add;
  import org.moeaframework.util.tree.Cos;
  import org.moeaframework.util.tree.Divide;
8
  import org.moeaframework.util.tree.Environment;
  import org.moeaframework.util.tree.Exp;
  import org.moeaframework.util.tree.Get;
11
  import org.moeaframework.util.tree.Log;
  import org.moeaframework.util.tree.Multiply;
  import org.moeaframework.util.tree.Rules;
14
  import org.moeaframework.util.tree.Sin;
  import org.moeaframework.util.tree.Subtract;
16
17
  public class FunctionMatcherProblem extends AbstractProblem {
18
19
    public double[] x;
20
    public double[] y;
22
23
    private Rules rules;
24
25
    public FunctionMatcherProblem() {
26
      super(1, 1);
27
28
      rules = new Rules();
29
       rules.add(new Add());
30
       rules.add(new Multiply());
31
       rules.add(new Subtract());
32
       rules.add(new Divide());
33
       rules.add(new Sin());
34
       rules.add(new Cos());
35
       rules.add(new Exp());
36
37
       rules.add(new Log());
       rules.add(new Get(Number.class, "x"));
38
       rules.setReturnType(Number.class);
39
      rules.setMaxVariationDepth(10);
40
      x = new double[100];
42
43
       y = new double[100];
44
       for (int i = 0; i < 100; i++) {</pre>
45
        x[i] = 2.0*(i / 100.0) - 1.0; // range from -1 to 1
46
        y[i] = Math.pow(x[i], 5) - 2.0*Math.pow(x[i], 3) + x[i];
47
48
```

```
49
50
    @Override
51
    public Solution newSolution() {
      Solution solution = new Solution(1, 1);
53
      solution.setVariable(0, new Program(rules));
54
      return solution;
56
57
    public double[] calculate(Program program) {
      double[] approximatedY = new double[x.length];
60
       for (int i = 0; i < x.length; i++) {</pre>
61
         Environment environment = new Environment();
         environment.set("x", x[i]);
         approximatedY[i] = ((Number)program.evaluate(environment)).doubleValue()
66
       return approximatedY;
67
    }
68
69
    @Override
70
    public void evaluate(Solution solution) {
71
      Program program = (Program) solution.getVariable(0);
72
       // calculate the difference between the approximation and actual
74
       double[] approximatedY = calculate(program);
75
       double difference = 0.0;
76
77
       for (int i = 0; i < x.length; i++) {</pre>
         difference += Math.pow(Math.abs(y[i] - approximatedY[i]), 2.0);
79
81
      difference = Math.sqrt(difference);
82
83
       // protect against NaN
84
       if (Double.isNaN(difference)) {
85
         difference = Double.POSITIVE_INFINITY;
86
87
88
      solution.setObjective(0, difference);
89
90
91
```

MOEAF ramework/book/chapter 7/Function Matcher Problem. java

The target function is defined on lines 45-48. For this example, we are trying to match the function $x^5 - 2x^3 + x$, plotted below.



Use the following code to run this example:

```
package chapter7;
2
  import org.moeaframework.Executor;
  import org.moeaframework.analysis.plot.Plot;
  import org.moeaframework.core.NondominatedPopulation;
  import org.moeaframework.core.variable.Program;
6
7
  public class RunFunctionMatcherProblem {
8
9
    public static void main(String[] args) {
10
      FunctionMatcherProblem problem = new FunctionMatcherProblem();
12
      NondominatedPopulation result = new Executor()
           .withAlgorithm("NSGAII")
14
           .withProblemClass(FunctionMatcherProblem.class)
           .withMaxEvaluations(10000)
16
17
           .run();
18
       // get the resulting function
19
      Program program = (Program) result.get(0).getVariable(0);
20
21
       // print the function
22
      System.out.println(program);
23
       System.out.println("Distance: " + result.get(0).getObjective(0));
24
25
```

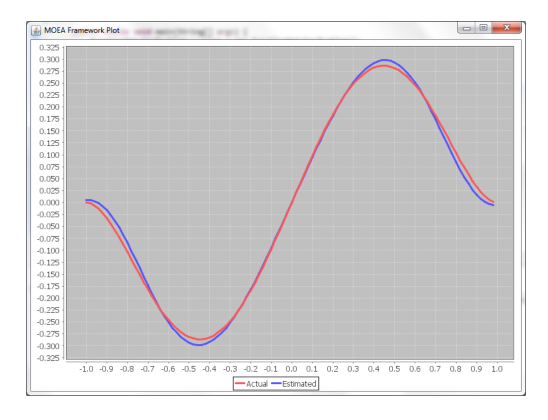
```
// display a plot comparing the two functions
new Plot()
    .line("Actual", problem.x, problem.y)
    .line("Estimated", problem.x, problem.calculate(program))
    .show();
}
```

MOEAFramework/book/chapter7/RunFunctionMatcherProblem.java

The result of running this code is shown below (you may see different results each time you run this program). The program whose output best matches the target function is:

```
(Program (Subtract (Sin x) (Multiply (Sin x) (Sin (Multiply (Sin x) (Add (Multiply x (Sin (Divide x (Sin x)))) (Multiply x (Sin (Divide x (Sin x)))))))))
```

which results in a mean square error of only 0.0275. Not too bad. The corresponding plot of the actual and estimated functions is shown below:



You may observe that the resulting function is more complicated as the original function. It is certainly possible for the optimization algorithm to find the original function, but it may require an extensive amount of time. In reality, we wouldn't know the original function, but would instead only have a bunch of points sampled at different locations.

7.3.3 Custom Operators

In addition to the built-in operators, you can also define your own custom operators. Suppose we want to define an operator to compute the hypotenuse. The operator accepts two inputs, x and y, and returns $\sqrt{x^2 + y^2}$. Our new operator, called Hypot, must extend the Node class, as shown below.

```
package chapter7;
  import org.moeaframework.util.tree.Environment;
3
  import org.moeaframework.util.tree.Node;
  public class Hypot extends Node {
6
7
    public Hypot() {
8
      super(Number.class, Number.class);
9
10
11
    @Override
    public Node copyNode() {
13
      return new Hypot();
14
16
    @Override
17
    public Object evaluate(Environment environment) {
18
      Number x = (Number) getArgument(0).evaluate(environment);
19
      Number y = (Number)getArgument(1).evaluate(environment);
20
21
      return Math.hypot(x.doubleValue(), y.doubleValue());
22
23
24
```

MOEAFramework/book/chapter7/Hypot.java

Three methods are required. The constructor, on lines 8-10, at a minimum must call the constructor of the Node class by calling **super**. The first argument to **super** is the output/return type, and the remaining arguments are the input types. Thus, by calling **super**(Number.class, Number.class, Number.class), we are defining an operator with two numeric inputs. The copyNode method on lines 13-15 returns a duplicate copy of the node. Finally, on lines 18-23, the evaluate method reads the arguments, computes and returns the result.

The standard way to read arguments is shown on lines 19 and 20 and involves three steps. Step 1 is reading the argument by calling getArgument. The argument will be another Node, which itself could be an constant value or a more complex expression. Step 2 recursively evaluates the arguments in the given environment. Step 3 casts the result to the appropriate type. This should be the same type we defined in the constructor.

For flexibility, we recommend using the abstract Number class to represent numeric values rather than Integer or Double. Doing so allows the operators to work on any numeric inputs. The operator should automatically convert integers to doubles if required.

After defining our Hypot class, we add it to the rules:

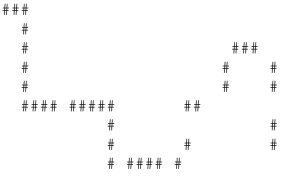
```
rules.add(new Hypot());
```

7.3.4 Ant Problem

A good example of defining custom operators is the Ant problem. The ant problem defines a world containing food at certain positions, and aims to design a robotic ant to navigate through the world and eat the food. This robotic ant is controlled by a program consisting of several operations:

- 1. if-else condition,
- 2. turn left,
- 3. turn right,
- 4. move forward, and
- 5. check if food is ahead.

We want to use the MOEA Framework to develop a program consisting of these operations, allowing the robotic ant to eat all the food. The complete code for the ant problem can be found in the <code>examples/org/moeaframework/examples/gp/ant</code> folder. Each of the operations is defined in a separate Java class, such as <code>IsFoodAhead.java</code> and <code>IurnLeft.java</code>. These operators act on a special world environment defined in <code>Morld.java</code>. <code>Morld.java</code> defines the size of the world, the current position of the ant, and the location of food. For example, the world initially starts off empty with some locations containing food, as indicated by the <code>#</code> symbol. For example, the world could appear as follows:



The ant starts at location (0, 0) and, using the program, tries to follow the path of food. As it encounters food, it changes the symbol to @ to indicate the food has been eaten. For example, a successful ant would results in the state:

```
Moves: 78 / 500
Food: 36 / 36
 999
    @
    (a
                                 @@@
    (a
                                a
                                a
                                       @
    0000 00000
                          (a (a
                                      9
                          (a
                                      (a
                  0 0 0 0
```

The rules for the ant program are shown below:

```
rules = new Rules();
rules.add(new TurnLeft());
rules.add(new TurnRight());
rules.add(new MoveForward());
rules.add(new IsFoodAhead());
rules.add(new IfElse(Void.class));
rules.add(new Sequence(Void.class, Void.class));
rules.setReturnType(Void.class);
```

So how does the MOEA Framework know how to combine these operations? The type system. First lets consider IfElse. By passing in the type Void.class, we construct an if-else structure with three inputs — (condition, if-case, else-case) — with types (Boolean.class, Void.class, Void.class) and a return type of Void.class. The condition argument is always type Boolean.java, which implies any operation that returns type Boolean.class can fit. In this example, only one operation matches: IsFoodAhead. The other operations — TurnLeft, TurnRight, and MoveForward — all have a return type of Void.class. The type Void.class indicates that the expression does not return a value.

IfElse and Sequence are special structures. You can customize the type of the operator. For example, calling **new** IfElse(Void.class) says that any expression in the if-case or else-case must return type Void.class. Likewise, calling **new** Sequence(Void.class, Void.class) allows sequencing together two expressions with return type Void.class. This means that the if-else cases and sequences can only contain operators TurnLeft, TurnRight, MoveForward, and nested IfElse statements or Sequences. Finally, the overall program has a return type of Void.class Note the return type is Void.class, which indicates the program should not return a value.

Within examples/org/moeaframework/examples/gp/ant are two larger examples which can be executed by running SantaFeExample.java and LosAltosExample.java. You may observe that different programs are generated for these two examples. Since we are evolving a program to solve a single example, the program tends to be "overfit". In other words, the program works well on the example it was designed for, but will perform poorly on other examples. One way to combat overfitting is to evaluate the program on multiple example. The results from each individual evaluation could be averaged. It is also common to use a minimax approach, where you try to minimize the maximum value (or maximimize the minimum value). In this way, you are searching for a program with the best worst-case performance.

Chapter 8

Integers and Mixed Integer Programming

Another discrete type commonly seen in optimization is integers. We devote a chapter to integers because there are actually two ways to represent integers, as we will discuss below. Additionally, it is commonly to see problems with integers and real-values mixed together, forming what is called mixed integer programming.

8.1 Two Representations

The MOEA Framework supports two representations for integers. One is backed by real-values and the other by bit strings. The real-value representation is straightforward. Simply encode your integer as a real-value and cast/truncate to an integer. The EncodingUtils.newInt(a, b) and EncodingUtils.getInt(...) methods conveniently perform this function. The bit string representation on the other hand is similar to the binary representation of numbers in a computer. There are caveats to using both methods, which must be carefully understood to avoid potential pitfalls.

Real-Valued Representation for Integers

When using EncodingUtils.newInt(a, b), the underlying variable is actually a RealVariable bounded by [a,b+1). When reading the value using EncodingUtils. getInt(...), the real-value is truncated using the floor operation to convert it to an integer.

Advantages:

- Works with all algorithms and operators for real-values.
- Can be combined with other real-valued decision variables without changing operators.

Disadvantages:

• Small changes to the underlying real value will not cause any changes when truncated to an integer.

Bit String Representation for Integers

When using EncodingUtils.newBinaryInt(a, b), the underlying variable is actually a BinaryVariable. The length of the bit string is chosen to encompass the bounds of the integer. The corresponding value is read using EncodingUtils.getInt(...). There are two methods to encode the bit string. First is the traditional binary encoding used by computers. The disadvantage of binary encoding is that a single bit flip can cause a substantial change to the resulting integer. Alternatively, you can use gray coding, which is designed so that there exists a single bit flip to move between adjacent integers. This enables smoother transitions between values, but large jumps can still occur.

Advantages:

- Works with all algorithms and operators for bit strings (binary).
- Mutation operators are guaranteed to change the value of the integer.

Disadvantages:

- If $b-a < 2^n$, then some values will occur with higher probability (since more than one bit pattern maps to the integer).
- Large changes to the integer value can be caused by small changes to the underlying bit string.

Gray coding is used by default. Binary coding can be enabled by calling the constructor **new** BinaryIntegerVariable(a, b, **false**).

Demonstration

Lets look at how both representations work on a problem. We will modify the Schaffer problem from Chapter 2 to use integer values instead of real values. For example, here is the version using the real-valued representation for integers:

```
package chapter8;

import org.moeaframework.core.Solution;
import org.moeaframework.core.variable.EncodingUtils;
import org.moeaframework.problem.AbstractProblem;

public class RealIntegerSchafferProblem extends AbstractProblem {
```

```
8
    public RealIntegerSchafferProblem() {
9
       super(1, 2);
10
11
    @Override
    public void evaluate(Solution solution) {
14
      double x = EncodingUtils.getInt(solution.getVariable(0));
15
16
      solution.setObjective(0, Math.pow(x, 2.0));
       solution.setObjective(1, Math.pow(x - 2.0, 2.0));
18
19
    }
20
    @Override
21
    public Solution newSolution() {
22
      Solution solution = new Solution(1, 2);
23
      solution.setVariable(0, EncodingUtils.newInt(-10, 10));
24
      return solution;
26
27
28
```

MOEAFramework/book/chapter8/RealIntegerSchafferProblem.java Alternatively, here is the same class using the bit string representation:

```
package chapter8;
  import org.moeaframework.core.Solution;
3
  import org.moeaframework.core.variable.EncodingUtils;
  import org.moeaframework.problem.AbstractProblem;
7
  public class BinaryIntegerSchafferProblem extends AbstractProblem {
    public BinaryIntegerSchafferProblem() {
9
      super(1, 2);
10
11
12
    @Override
13
    public void evaluate(Solution solution) {
14
      double x = EncodingUtils.getInt(solution.getVariable(0));
16
      solution.setObjective(0, Math.pow(x, 2.0));
      solution.setObjective(1, Math.pow(x - 2.0, 2.0));
18
19
20
    @Override
21
    public Solution newSolution() {
      Solution solution = new Solution(1, 2);
23
      solution.setVariable(0, EncodingUtils.newBinaryInt(-10, 10));
24
      return solution;
25
26
```

27

MOEAFramework/book/chapter8/BinaryIntegerSchafferProblem.java

Observe that only line 24 changes. Since both representations are read using EncodingUtils.getInt(...), the evaluate method remains unchanged. We can then run both versions:

```
package chapter8;
2
  import org.moeaframework.Executor;
3
  import org.moeaframework.core.NondominatedPopulation;
  import org.moeaframework.core.Solution;
6
  import org.moeaframework.core.variable.EncodingUtils;
  public class RunIntegerSchafferProblems {
8
9
    public static void main(String[] args) {
      NondominatedPopulation result1 = new Executor()
11
           .withAlgorithm("NSGAII")
           .withProblemClass (BinaryIntegerSchafferProblem.class)
13
           .withMaxEvaluations(10000)
14
           .run();
16
      NondominatedPopulation result2 = new Executor()
17
           .withAlgorithm("NSGAII")
18
           .withProblemClass(RealIntegerSchafferProblem.class)
19
           .withMaxEvaluations(10000)
20
           .run();
22
      System.out.println("Binary Integer Encoding:");
23
      for (Solution solution : result1) {
24
        System.out.printf(" d = 0.5f, 0.5f",
25
             EncodingUtils.getInt(solution.getVariable(0)),
26
27
             solution.getObjective(0),
             solution.getObjective(1));
28
29
30
      System.out.println();
31
      System.out.println("Real Integer Encoding:");
      for (Solution solution : result2) {
33
        System.out.printf(" d = 0.5f, .5fn",
             EncodingUtils.getInt(solution.getVariable(0)),
35
             solution.getObjective(0),
36
             solution.getObjective(1));
37
38
39
40
41 }
```

MOEAF ramework/book/chapter 8/RunInteger Schaffer Problems. java

Which produces the following output:

```
Binary Integer Encoding:

0 => 0.00000, 4.00000

1 => 1.00000, 1.00000

2 => 4.00000, 0.00000

Real Integer Encoding:

0 => 0.00000, 4.00000

1 => 1.00000, 1.00000

2 => 4.00000, 0.00000
```

On a simple problem such as this, both representations produce identical results. On more complex problems, you may observe some differences in performance based on which version is used. Experimentation is a good way to determine which version is appropriate for your problem.

8.2 Mixed Integer Programming

Real-world problems tend to not exclusively use only real-values or integers, but instead are formulated as a combination of decision variable types. When mixing real values and integers, one must be mindful of the underlying representation for integers. For example, lets modify the Srinivas problem from Chapter 3 to include one integer and one real-valued decision variable:

```
package chapter8;
  import org.moeaframework.core.Solution;
  import org.moeaframework.core.variable.EncodingUtils;
  import org.moeaframework.problem.AbstractProblem;
  public class MixedIntegerSrinivasProblem extends AbstractProblem {
8
    public MixedIntegerSrinivasProblem() {
9
      super(2, 2, 2);
    @Override
    public void evaluate(Solution solution) {
14
      int x = EncodingUtils.getInt(solution.getVariable(0));
      double y = EncodingUtils.getReal(solution.getVariable(1));
16
      double f1 = Math.pow(x - 2.0, 2.0) + Math.pow(y - 1.0, 2.0) + 2.0;
17
      double f2 = 9.0 * x - Math.pow(y - 1.0, 2.0);
18
```

```
19
       double c1 = Math.pow(x, 2.0) + Math.pow(y, 2.0) - 225.0;
       double c2 = x - 3.0*y + 10.0;
20
21
       solution.setObjective(0, f1);
22
       solution.setObjective(1, f2);
23
       solution.setConstraint(0, c1 <= 0.0 ? 0.0 : c1);</pre>
24
       solution.setConstraint(1, c2 \le 0.0 ? 0.0 : c2);
25
26
27
    @Override
28
    public Solution newSolution() {
       Solution solution = new Solution(2, 2, 2);
30
31
       solution.setVariable(0, EncodingUtils.newBinaryInt(-20, 20));
32
       solution.setVariable(1, EncodingUtils.newReal(-20.0, 20.0));
34
      return solution;
35
36
37
```

MOEAFramework/book/chapter8/MixedIntegerSrinivasProblem.java

If using the real-valued representation for integers, then the same operators can be shared for both real-values and integers. However, if you try to combine the bit string representation with other real-valued variables, you will see the following error:

```
Exception in thread "main" org.moeaframework.core.spi.
   ProviderNotFoundException: no provider for NSGAII
 at org.moeaframework.algorithm.StandardAlgorithms.getAlgorithm(
     StandardAlgorithms.java:248)
 at org.moeaframework.core.spi.AlgorithmFactory.instantiateAlgorithm(
     AlgorithmFactory.java:175)
 at org.moeaframework.core.spi.AlgorithmFactory.getAlgorithm(AlgorithmFactory
     .java:137)
 at org.moeaframework.Executor.runSingleSeed(Executor.java:773)
 at org.moeaframework.Executor.run(Executor.java:730)
 at chapter8.RunMixedIntegerSrinivasProblem.main(
     RunMixedIntegerSrinivasProblem.java:16)
Caused by: org.moeaframework.core.spi.ProviderLookupException: unable to find
   suitable variation operator
 at org.moeaframework.core.spi.OperatorFactory.lookupVariationHint(
     OperatorFactory.java:347)
 at org.moeaframework.core.spi.OperatorFactory.getVariation(OperatorFactory.
     java:182)
 at org.moeaframework.core.spi.OperatorFactory.getVariation(OperatorFactory.
     java:160)
 at org.moeaframework.algorithm.StandardAlgorithms.newNSGAII(
     StandardAlgorithms.java:329)
 at org.moeaframework.algorithm.StandardAlgorithms.getAlgorithm(
     StandardAlgorithms.java:207)
```

... 5 more

Note the line that says "unable to find suitable variation operator". This is an indication that no operator is compatible with the decision variables in the problem. The MOEA Framework automatically determines the appropriate operators given the decision variable types. However, this only works when all decision variables are the same type. If you try to mix the bit string representation of integers with other real-valued variables, you will run into this error. However, in most cases we can address this error by explicitly specifying the operators to use:

```
package chapter8;
2
3
  import org.moeaframework.Executor;
  import org.moeaframework.core.NondominatedPopulation;
  import org.moeaframework.core.Solution;
  import org.moeaframework.core.variable.EncodingUtils;
  public class RunMixedIntegerSrinivasProblem {
8
9
    public static void main(String[] args) {
      NondominatedPopulation result = new Executor()
           .withAlgorithm("NSGAII")
12
           .withProblemClass (MixedIntegerSrinivasProblem.class)
13
           .withMaxEvaluations(10000)
14
           .withProperty("operator", "sbx+hux+pm+bf")
           .run();
16
17
      for (Solution solution : result) {
18
        if (!solution.violatesConstraints()) {
19
           System.out.format("3d \%7.3f => \%7.3f \%7.3f\%n",
20
               EncodingUtils.getInt(solution.getVariable(0)),
21
               EncodingUtils.getReal(solution.getVariable(1)),
23
               solution.getObjective(0),
               solution.getObjective(1));
24
25
26
27
28
```

MOEAFramework/book/chapter8/RunMixedIntegerSrinivasProblem.java

Here, we explicitly specify the operators to use on line 15. Since we are combining bit string and real-valued variables, we must include operators for both types using the operator string "sbx+hux+pm+bf". Each operator is applied in the order they appear in the string. Furthermore, we want crossover operators to appear at the beginning and mutation operators at the end. All operators are type-safe, meaning they will only affect decision variables of the supported type. For example, the "hux" and "bf" (bit flip) operators will only affect

bit strings, while "sbx" and "pm" will only evolve real-values.

Such overloading of operators is also necessary when mixing other decision variable types. Again, just follow the rules above for setting the operator string. In the event that you provide an invalid combination of operators, then you will be given an error message describing the problem.

Chapter 9

I/O Basics

This chapter talks all about I/O: input and output. We'll walk through several use cases and provides example code.

9.1 Printing Solutions

The first thing you'll likely want to do after solving a problem with the MOEA Framework is viewing your solutions. Suppose we want to print the decision variables and objectives for all solutions in a result set:

```
package chapter9;
  import java.io.IOException;
  import org.moeaframework.Executor;
  import org.moeaframework.core.NondominatedPopulation;
  import org.moeaframework.core.Solution;
  import chapter2.SchafferProblem;
9
10
  public class PrintSolutions {
11
    public static void main(String[] args) throws IOException {
13
      NondominatedPopulation result = new Executor()
14
           .withAlgorithm("NSGAII")
           .withProblemClass(SchafferProblem.class)
16
           .withMaxEvaluations(10000)
17
           .run();
18
19
      for (int i = 0; i < result.size(); i++) {</pre>
20
         Solution solution = result.get(i);
21
         System.out.print("Solution " + i + ":");
22
         for (int j = 0; j < solution.getNumberOfVariables(); j++) {</pre>
24
```

```
System.out.print(" ");
           System.out.print(solution.getVariable(j));
26
27
2.8
         System.out.print(" =>");
29
30
         for (int j = 0; j < solution.getNumberOfObjectives(); j++) {</pre>
           System.out.print(" ");
32
           System.out.print(solution.getObjective(j));
35
         System.out.println();
37
38
39
40
```

MOEAFramework/book/chapter9/PrintSolutions.java

The following output is produced:

```
Solution 0: 2.0004392968937066 => 4.001757380556588 1.929817608202561E-7 Solution 1: -3.7964704972906493E-4 => 1.441318823679831E-7 4.001518732330799 Solution 2: 0.06057459075229121 => 0.0036692810448075643 3.7613709180356425 Solution 3: 1.657358848462215 => 2.7468383525759994 0.11740295872713932 Solution 4: 0.9436776287650016 => 0.8905274670315362 1.11581695197153 Solution 5: 1.8601691079468476 => 3.460229110159771 0.019552678372380346 Solution 6: 1.4155832357245992 => 2.003875897264526 0.34154295436612936 ...
```

We could clean up this output in several ways. First, the output shows way too many decimal digits. We can print formatted numbers as follows:

```
for (int i = 0; i < result.size(); i++) {</pre>
         Solution solution = result.get(i);
2
3
         System.out.print("Solution " + i + ":");
         for (int j = 0; j < solution.getNumberOfVariables(); j++) {</pre>
           System.out.printf(" %.3f", EncodingUtils.getReal(solution.getVariable(
               j)));
         System.out.print(" =>");
9
         for (int j = 0; j < solution.getNumberOfObjectives(); j++) {</pre>
11
           System.out.printf(" %.3f", solution.getObjective(j));
13
14
         System.out.println();
16
```

7

MOEAFramework/book/chapter9/PrintFormattedSolutions.java

This is better:

```
Solution 0: 0.000 => 0.000 4.001

Solution 1: 2.000 => 4.001 0.000

Solution 2: 1.048 => 1.099 0.906

Solution 3: 0.374 => 0.140 2.643

Solution 4: 0.335 => 0.113 2.771

Solution 5: 0.418 => 0.175 2.503

Solution 6: 1.947 => 3.790 0.003

...
```

A second improvement would be to order the solutions. Right now, the solutions appear in a random order. However, we could sort them lexicographically by their objective values.

```
result.sort(new LexicographicalComparator());
```

MOEAFramework/book/chapter9/PrintLexicographicalOrdering.java

```
Solution 0: 0.000 => 0.000 4.000

Solution 1: 0.020 => 0.000 3.919

Solution 2: 0.024 => 0.001 3.903

Solution 3: 0.043 => 0.002 3.831

Solution 4: 0.047 => 0.002 3.815

Solution 5: 0.075 => 0.006 3.707

Solution 6: 0.103 => 0.011 3.599

...
```

Great. Now we can clearly see an inverse relationship between the two objectives. As the first objective increases, the second objective decreases.

Note: When printing result sets, always remember to check solution. violatesConstraints() if the problem has constraints. Otherwise, you may display infeasible solutions in the output.

9.2 Files

Instead of printing the information to the console, we can save the solutions directly to a file. There are advantages and disadvantages of using files. The primary advantage is that you can load the saved data. For example, you could save the results from several runs, then load them in later to generate plots. The primary disadvantage is that the generated file is not human-readable. You can't open the saved file in a text editor to see its contents. It can only be reloaded into the MOEA Framework.

We use the PopulationIO class to read and write files. For example, here we save the result to a file:

```
try {
    PopulationIO.write(new File("solutions.dat"), result);
    System.out.println("Saved " + result.size() + " solutions!");
} catch (IOException e) {
    e.printStackTrace();
}
```

MOEAFramework/book/chapter9/SaveToFile.java

The file can then be loaded back into the MOEA Framework:

```
try {
    Population result = PopulationIO.read(new File("solutions.dat"));

System.out.println("Read " + result.size() + " solutions!");

catch (IOException e) {
    e.printStackTrace();
}
```

MOEAFramework/book/chapter9/LoadFromFile.java

If you need to save the solutions in a human-readable format, use the printing commands from the previous section.

9.3 Checkpoints

If checkpoints are enabled, a running algorithm will periodically save checkpoint files. The checkpoint file stores the current state of the algorithm. If the run is interrupted, such as during a power outage, the run can be resumed at the last saved checkpoint. Checkpointing can be enabled when using the Executor to run algorithms. Calling setCheckpointFile sets the file location for the checkpoint file, and checkpointEveryIteration or setCheckpointFrequency control how frequently the checkpoint file is saved.

Resuming a run from a checkpoint occurs automatically. If the checkpoint file does not exist, a run starts from the beginning. However, if the checkpoint file exists, then the run is automatically resumed at that checkpoint. For this reason, care must be taken when using checkpoints as they can be a source of confusion for new users. For instance, using the same checkpoint file from an unrelated run can cause unexpected behavior or an error. For this reason, checkpoints are recommended only when solving time-consuming problems.

```
package chapter9;
import java.io.File;
import org.moeaframework.Executor;
```

```
import org.moeaframework.core.NondominatedPopulation;
  import chapter2.SchafferProblem;
8
9
  public class Checkpoints {
    public static void main(String[] args) {
      File checkpointFile = new File("checkpoint.dat");
13
      long start = System.currentTimeMillis();
14
      if (checkpointFile.exists()) {
16
        System.out.println("Checkpoint file exists, will resume from prior run!"
17
            );
18
20
      NondominatedPopulation result = new Executor()
           .withAlgorithm("NSGAII")
21
           .withProblemClass (SchafferProblem.class)
22
           .withMaxEvaluations(1000000)
           .withCheckpointFrequency(10000)
24
           .withCheckpointFile(checkpointFile)
26
27
      System.out.println("Elapsed time: " + (System.currentTimeMillis() - start)
28
           / 1000 + "s");
29
30
31
```

MOEAFramework/book/chapter9/Checkpoints.java

The first time you run this code, it'll take several seconds to produce the result. If you run the code a second time, it should terminate immediately since the checkpoint file exists. Note that checkpoint files are never deleted by the MOEA Framework. Each time you run this example, it will resume from its last save point. If you want to run this example from the beginning, you must delete the checkpoint file manually.

9.4 Creating Reference Sets

All of the test problems provided by the MOEA Framework are accompanied by a predefined reference set. The reference set contains the Pareto optimal solutions to a problem, or if the set is too large, an approximation of the Pareto optimal solutions. If you are working on a new problem, you may not have a reference set available. A common method of generating reference sets, particularly for real-world problems, is to solve the problem using many different algorithms repetitively. The combined result from all runs becomes your reference set.

Lets start by generating the reference set for the Schaffer problem:

```
package chapter9;
  import java.io.File;
3
  import java.io.IOException;
  import org.moeaframework.Executor;
  import org.moeaframework.core.NondominatedPopulation;
  import org.moeaframework.core.PopulationIO;
  import chapter2.SchafferProblem;
10
  public class CreatingReferenceSet {
    public static void main(String[] args) throws IOException {
14
      int nseeds = 25;
      String[] algorithms = new String[] { "NSGAII", "GDE3", "OMOPSO" };
16
17
      NondominatedPopulation referenceSet = new NondominatedPopulation();
18
19
      Executor executor = new Executor()
20
           .withProblemClass(SchafferProblem.class)
22
           .withMaxEvaluations(10000);
23
      for (String algorithm : algorithms) {
24
        executor.withAlgorithm(algorithm);
25
26
        for (int i = 0; i < nseeds; i++) {</pre>
           referenceSet.addAll(executor.run());
28
29
30
31
      PopulationIO.writeObjectives(new File("Schaffer.pf"), referenceSet);
32
33
34
```

MOEAFramework/book/chapter9/CreatingReferenceSet.java

On line 18 we create the NondominatedPopulation that will store the combined Pareto solutions from all runs. The loop on line 24 runs each of our test algorithms for 25 repetitions. Finally, we save the reference set to a file on line 32.

When using a NondominatedPopulation, an issue one frequently encounters is that the sets can become arbitrarily large. There is no bound on the size of the set. The only restriction is that all solutions must be non-dominated. After running the above code, we observed that the set contained 9700 solutions. This can become cumbersome to work with in practice!

A common technique for reducing the size of a reference set is to use ϵ -dominance (Laumanns et al., 2002). ϵ -dominance is similar to Pareto dominance, except the ϵ defines a minimum resolution of solutions. Larger ϵ s produce smaller sets. With a small tweak to the previous code, we can define ϵ s:

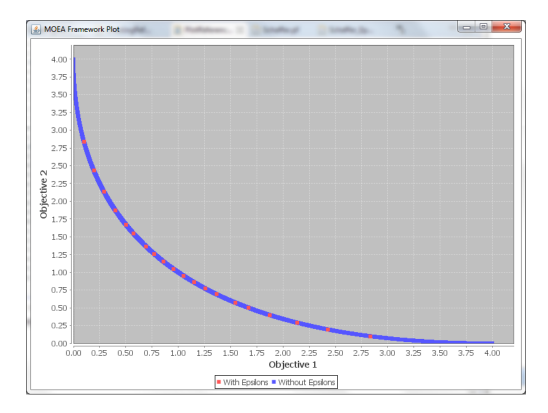
```
NondominatedPopulation referenceSet = new EpsilonBoxDominanceArchive(
new double[] { 0.1, 0.1 });
```

MOEAFramework/book/chapter9/CreatingReferenceSetWithEpsilons.java

With ϵ s, the resulting reference set contains only 20 solutions. Plotting the two sets shows the drastic difference:

```
1
  package chapter9;
2
  import java.io.File;
  import java.io.IOException;
  import org.moeaframework.analysis.plot.Plot;
  import org.moeaframework.core.Population;
  import org.moeaframework.core.PopulationIO;
9
  public class PlotReferenceSet {
10
    public static void main(String[] args) throws IOException {
      Population with Epsilons = Population IO. read Objectives (
           new File("Schaffer_Epsilon.pf"));
14
      Population withoutEpsilons = PopulationIO.readObjectives(
16
           new File("Schaffer.pf"));
17
18
      new Plot()
19
           .add("With Epsilons", withEpsilons)
20
           .add("Without Epsilons", withoutEpsilons)
21
22
           .show();
23
24
```

MOEAFramework/book/chapter9/PlotReferenceSet.java



Observe that the red and blue points lie on the same curve, except we approximate the curve with significantly fewer red points. If using this reference set in practice, we would probably want to pick slightly smaller ϵ s, since 20 points may be too small. For this problem, 100 would be ideal.

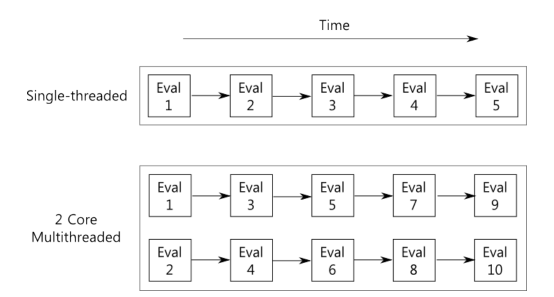
Chapter 10

Performance Enhancements

This chapter details two ways to improve the performance of your optimization. The first utilizes multithreading on your computer to utilize all available computing cores. The second rewrites the problem definition in another language. We consider compiling to a native library using C/C++ and writing functions in Python. Other languages can be supported following similar techniques.

10.1 Multithreading

When you typically optimize a program, you evaluate one solution at a time. This is a single-threaded program, where each step is performed sequentially. However, most modern day computers have processors with multiple computing cores. Suppose a computer has 4 cores. A single-threaded program would only utilize 1 of the 4 cores, or 25% of the available resources. Multithreaded programs are designed to split the computations into multiple execution threads, each of which can be executed on a separate core simultaneously. In the context of an MOEA, as shown below, the objective function evaluations can be multithreaded.



This diagram depicts the 2 core multithreaded program achieving twice as many evaluations as the single threaded program. In reality, it is a bit more nuanced. First, not all MOEAs can support multithreading. Some are inherently single threaded. The MOEA Framework uses special "future" objects to automatically detect if an MOEA support multithreading. Second, there is often additional overhead encountered when using multithreading or any type of parallelization. This overhead comes from a variety of sources, but perhaps the most common source of overhead is from communicating data between the two cores in memory. Memory access tends to be significantly slower than CPU processing. If the problem consists only of a few lines of code, then likely the evaluation time is less than the communication overhead. Enabling multithreading would slow down the program in this case. Third, if you attempt to utilize all cores on a computer, particularly one with iterative user sessions such as Windows, you will have to compete with other programs for time on the CPU.

To demonstrate this, we will modify our Schaffer problem. The original Schaffer problem is very simple and very fast to compute. To slow down the evaluations, lets add a loop to add some additional computing time to each function evaluation:

```
double sum = 0.0;

for (int i = 0; i < 100000; i++) {
    sum += i;
}</pre>
```

On a modern computer, this adds approximately 0.1 milliseconds per evaluation. Thus, our new Schaffer problem, which we call ExpensiveSchafferProblem, would appear as follows:

```
package chapter10;
```

```
import org.moeaframework.core.Solution;
  import org.moeaframework.core.variable.EncodingUtils;
4
  import org.moeaframework.problem.AbstractProblem;
  public class ExpensiveSchafferProblem extends AbstractProblem {
8
    public ExpensiveSchafferProblem() {
9
       super(1, 2);
    @Override
    public void evaluate(Solution solution) {
14
      double x = EncodingUtils.getReal(solution.getVariable(0));
       // perform some expensive calculation
17
      double sum = 0.0;
18
19
       for (int i = 0; i < 100000; i++) {</pre>
20
         sum += i;
21
23
      solution.setObjective(0, Math.pow(x, 2.0));
24
       solution.setObjective(1, Math.pow(x - 2.0, 2.0));
25
26
27
    @Override
28
    public Solution newSolution() {
29
       Solution solution = new Solution(1, 2);
30
       solution.setVariable(0, EncodingUtils.newReal(-10.0, 10.0));
31
      return solution;
32
33
34
35
```

MOEAFramework/book/chapter10/ExpensiveSchafferProblem.java

Enabling multithreading in the MOEA Framework is straightforward: when using the Executor simply call distributeOnAllCores() to run on all available cores, or call distributeOn(N) to run on N cores. Any optimization algorithm that supports multithreading will automatically distribute evaluations on the requested number of cores on your computer. In the following code, we compare the runtime between the single threaded version and a multithreaded version:

```
package chapter10;

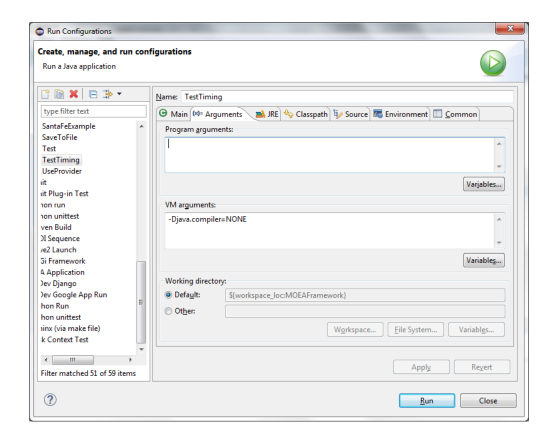
import org.moeaframework.Executor;

public class TestTiming {
   public static void main(String[] args) {
```

```
long startTime;
8
9
       // run without multithreading
10
      startTime = System.currentTimeMillis();
      new Executor()
           .withAlgorithm("NSGAII")
14
           .withProblemClass(ExpensiveSchafferProblem.class)
15
           .withMaxEvaluations(10000)
16
           .run();
18
19
       System.out.println("Single threaded: " + (System.currentTimeMillis()-
          startTime));
20
       // run with mulithreading
21
       startTime = System.currentTimeMillis();
22
      new Executor()
24
           .withAlgorithm("NSGAII")
25
           .withProblemClass (ExpensiveSchafferProblem.class)
26
           .withMaxEvaluations(10000)
27
           .distributeOnAllCores()
28
           .run();
29
30
      System.out.println("Multithreading: " + (System.currentTimeMillis() -
31
          startTime));
32
33
```

MOEAFramework/book/chapter10/TestTiming.java

Prior to running the example, it is important to set the <code>-Djava.compiler=NONE</code> Java VM option. Without this option, Java will attempt to optimize our program and will remove the for loop. In practice, this optimization is very useful, but for this demonstration we must temporarily disable it. In Eclipse, you set it in the run configuration window, as shown below.



Running this program produces the output below:

Single threaded: 14339 Multithreading: 4679

It took 14.3 seconds for the single threaded program and 4.6 seconds for the multithreaded program on an 4 core machine. We would expect the multithreaded program to run in 14.3/4 = 3.575 seconds, but due to overhead it takes much longer. When comparing single threaded and multithreaded programs, there are two useful calculations. The first is speedup:

$$S = T_s/T_p, (10.1)$$

where S is the parallel speedup, T_s is the total runtime for the single threaded (or serial) version, and T_p is the total runtime for the multithreaded (or parallel) version. In our example, the speedup is 14.3/4.6 = 3.1, meaning that our multithreaded version runs approximately three times faster. The second calculation is efficiency:

$$E = S/p, (10.2)$$

where E is the parallel efficiency, S is the parallel speedup, and p is the number of processors or cores. Thus, in our example, the efficiency is 3.1/4 = 0.78 or 78%. In other words, we

utilized 78% of the available computing resources. The remaining 22% is lost due to overhead. Larger overheads require longer evaluation times in order to achieve higher efficiency. Conversely, smaller overheads make multithreading and parallelization more efficient. If the overhead is quite large or the evaluation time is too short, it is possible for the multithreaded version to take longer to run! If considering the use of multithreading, it is good practice to test the speedup and efficiency on a small run.

10.2 Termination Conditions

In all of the examples seen thus far, we have run the algorithm for a fixed number of evaluations using withMaxEvaluations (...). Running for a fixed number of evaluations is commonly used when comparing different algorithms, since each algorithm will be given the same number of evaluations. However, in practical situations, one is often limited by computing resources. For example, management may require an engineering team to devise a feasible solution within a day or just a few hours. This can be achieved by replacing withMaxEvaluations (...) with withMaxTime (...).

```
new Executor()
    .withAlgorithm("NSGAII")
    .withProblemClass(ExpensiveSchafferProblem.class)
    .withMaxTime(30000) // 30 seconds
    .run();
```

MOEAFramework/book/chapter10/TerminationCondition.java

In this example, we are running the algorithm for 30 seconds, as indicated by withMaxTime (30000). Note that the time is specified in milliseconds (1 second = 1000 milliseconds).

You may notice that the algorithm takes slightly longer than 30 seconds to run. This is because the algorithm is allowed to finish the current iteration before terminating. For problems with longer evaluation times, this can become more noticeable. Be aware of this when scheduling jobs on computing resources that have hard time limits.

10.3 Native Compilation

While Java itself is a fast language compared to interpreted languages like Python, it still remains around 2-3 times slower than C, C++, and Fortran. If your evaluation contains a non-trivial amount of code, you could experience speedup by writing the evaluation code in one of these languages and compiling the code natively. Java Native Access (JNA)¹ is an open source library that lets a Java program invoke natively compiled code. If your problem is already written in one of these native languages, you can also use JNA to call the functions directly.

¹https://github.com/java-native-access/jna

Lets continue with the Schaffer problem. Suppose we wanted to re-write the Schaffer problem in C. We would need a C function with two arguments: an array of decision variables and an array of the objectives. If the problem has constraints, we would include a third array. Our function, called schaffer, would look like:

```
#include <math.h>

void schaffer(double* vars, double* objs) {
  objs[0] = pow(vars[0], 2.0);
  objs[1] = pow(vars[0] - 2.0, 2.0);
}
```

MOEAFramework/book/chapter10/schaffer.c

We can then compile this code to a native shared library as shown below.

```
gcc -03 -shared -o schaffer.dll schaffer.c -lm
```

When compiling code natively, there are many issues that can arise. First, you must determine if your computer is 32-bit or 64-bit. If you have a 64-bit computer, you will want to ensure that your installed version of Java is 64-bit. You will likely also need to include the -m64 option when compiling the example code, as shown below. Failing to do so will produce an UnsatisfiedLinkException error message when running this example.

```
gcc -m64 -03 -shared -o schaffer.dll schaffer.c -lm
```

Second, you must install an appropriate compiler for your system. If using a 64-bit machine, be sure to get a 64-bit compatible compiler. On Windows, we recommend MinGW². Third, you will need to use an appropriate shared library extension for your operating system. As shown in these examples, we use the .dll extension on Windows. Following a Unix/Linux naming convention, you would call the library libschaffer.so.

Next, we need to write a Java class that interfaces with the native library we just compiled, as shown below:

```
package chapter10;

import org.moeaframework.core.Solution;
import org.moeaframework.core.variable.EncodingUtils;
import org.moeaframework.problem.AbstractProblem;

import com.sun.jna.Native;

public class JNASchafferProblem extends AbstractProblem {
```

²http://www.mingw.org/

```
public static native double schaffer(double[] vars, double[] objs);
11
12
    static {
13
       System.setProperty("jna.library.path", "./book/chapter10");
14
      Native.register("schaffer");
16
17
    public JNASchafferProblem() {
18
       super(1, 2);
19
20
21
    @Override
22
    public void evaluate(Solution solution) {
23
       double[] vars = EncodingUtils.getReal(solution);
24
       double[] objs = new double[solution.getNumberOfObjectives()];
25
26
       schaffer(vars, objs);
27
2.8
       solution.setObjectives(objs);
29
    }
30
    @Override
32
    public Solution newSolution() {
33
       Solution solution = new Solution(1, 2);
34
       solution.setVariable(0, EncodingUtils.newReal(-10.0, 10.0));
35
       return solution;
36
37
38
```

MOEAFramework/book/chapter10/JNASchafferProblem.java

As with previous problems, we extend the AbstractProblem. However, unlike previous examples, we need to connect this code to the native library. First, on line 11, we create a "native" version of the schaffer function. The name and arguments to this function must match the name and arguments of the C function. In this case, a double array (double[]) maps to a double pointer in C (double*).

Second, on lines 13-16, we connect our Java class with the native library using JNA. We first set the jna.library.path system property on line 14 to include the path to the compiled library (i.e., schaffer.dll). This is necessary for JNA to locate the library. Then, on line 15, we call Native.register("schaffer") register the code with JNA. JNA searches its library path for the shared library, called schaffer.dll (on Windows) or libschaffer.so (on Linux). Upon finding the library, it loads it into memory and configures Java so that calls to the schaffer method in Java are forwarded to the native library.

Finally, in the evaluate method on lines 23-30, we call the schaffer method to evaluate the problem. On line 24, we create a double array with the decision variables, and on line 25 we create an empty array to store the resulting objectives. The schaffer

method is invoked on line 27 to call the underlying native code. Lastly, we set the objectives for the solution. One would handle constraints similarly.

We can then optimize the problem as we have done in the past:

```
NondominatedPopulation result = new Executor()
.withAlgorithm("NSGAII")
.withProblemClass(JNASchafferProblem.class)
.withMaxEvaluations(10000)
.run();
```

MOEAFramework/book/chapter10/ComparingCInterface.java

10.4 Standard I/O

If native compilation is not an option, the MOEA Framework also supports running models via standard I/O or network sockets with the ExternalProblem class. Here, we will discuss standard I/O using Python.

The ExternalProblem class defines a very simple interface for communicating with models over standard input and output (I/O). When the optimization starts, the MOEA Framework launches the process for evaluating solutions. For each solution, it sends a single line to the process containing the decision variables using whitespace to separate values. For example, it would pass five real-valued decision variables as:

```
0.109 0.912 0.3 0.291 0.154
```

The process parses the input line, evaluates the problem, and writes the objectives and constraints to its standard output, again separated by whitespace. For example, a problem with two objectives would write:

```
1.05 0.9
```

Any constraints would appear on the same line following the objective values. If the process receives a blank line, it terminates as the optimization has finished. For example, our Schaffer problem could be written in Python as follows:

```
from sys import *
from math import *

while True:
    # Read the next line from standard input
line = raw_input()

# If line is empty, stop
```

```
if line == "":
           break
11
       # Parse the decision variables from the input
       vars = map(float, line.split())
13
14
15
       # Evaluate the Schaffer problem
       objs = (vars[0]**2, (vars[0] - 2)**2)
16
17
       # Print objectives to standard output, flush to write immediately
18
      print "%f %f" % objs
19
       stdout.flush()
20
```

MOEAFramework/book/chapter10/schaffer.py

Line 4 starts an infinite loop that reads each line of input (line 6) and terminates only if a blank line is encountered (line 9). Line 13 parses each value from the input into the array vars. Next, the objectives are calculated on line 16 and written to the process' output on line 19. Always flush the output to immediately print the objectives as shown on line 20, otherwise the program may stall.

In addition to the Python code, we must also create a Java class to represent the problem. Unlike the previous examples, when connecting to an external process in this manner, we want to extend the ExternalProblem class, as shown below:

```
package chapter10;
  import java.io.IOException;
3
  import org.moeaframework.core.Solution;
  import org.moeaframework.core.variable.EncodingUtils;
  import org.moeaframework.problem.ExternalProblem;
  public class StdioSchafferProblem extends ExternalProblem {
9
10
    public StdioSchafferProblem() throws IOException {
11
12
      super("python", "book/chapter10/schaffer.py");
13
14
    @Override
    public String getName() {
16
      return "ExternalSchafferProblem";
18
19
    @Override
20
    public int getNumberOfConstraints() {
21
      return 0;
23
24
    @Override
    public int getNumberOfObjectives() {
26
```

```
return 2;
    }
28
29
    @Override
30
    public int getNumberOfVariables() {
31
32
       return 1;
33
34
    @Override
35
    public Solution newSolution() {
36
       Solution solution = new Solution(1, 2);
37
       solution.setVariable(0, EncodingUtils.newReal(-10.0, 10.0));
38
       return solution;
39
40
41
42
```

MOEAFramework/book/chapter10/StdioSchafferProblem.java

When extending the ExternalProblem class, we need to specify in the constructor the program to execute. In this case, we are running a Python script, so we need to invoke the Python interpreter on our schaffer.py file (line 12). However, once the Java class is defined, we optimize it like any other problem:

MOEAFramework/book/chapter10/ComparingCInterface.java

Lets take a quick look at how these interface options compare. Below we show the runtime for the Schaffer problem when it is implemented completely in Java, when we connect to a native library using JNA, and when we communicate via standard I/O.

```
Pure Java: 262
Via JNA: 377
Via Standard I/O: 532
```

Due to the simplicity of our problem, the pure Java option is fastest. For more complex problems, we would expect the C version to run faster. The JNA version adds an overhead of approximately 0.0115 milliseconds per evaluation, whereas the standard I/O version adds an overhead of approximately 0.027 milliseconds per evaluation.

10.5 A Note on Concurrency

We demonstrated two ways to interface with problems written in different languages: JNA and Standard I/O. One of the advantages of JNA is that it can be combined with multi-threaded evaluations. This requires that the native code you write in C or other language be reentrant. The term reentrant means that the function can be called multiple times, before prior calls have exited, and produce correct results. In order for a function to be reentrant, several rules should be followed³:

- The function must to access any static or global data that is not a constant.
- The function must not modify its own code
- The function must not call other non-reentrant functions

In our example C code, the schaffer function is entirely self-contained. It only reads the decision variables provided as an input to the function and returns the objectives. Thus, this function is reentrant.

If you have a problem that is non-reentrant, it is good practice to make Java's evaluate method synchronized. In you problem definition, you would define the evaluate method as follows:

```
public synchronized void evaluate(Solution solution) {
   // body of method
}
```

The **synchronized** keyword ensures that the evaluate method can be invoked one at a time. Thus, the synchronize keyword will prevent someone from accidentally parallelizing your problem.

³https://en.wikipedia.org/wiki/Reentrancy_(computing)

Chapter 11

Single Objective Optimization

The MOEA Framework, given its name, focuses on solving multiobjective problems. However, it is a general-purpose framework and can be used for single-objective optimization. This is particularly useful for comparing the efficacy of multiobjective algorithms against their single-objective counterparts. This chapter details using single-objective algorithms within the MOEA Framework.

11.1 Single Objective Problems

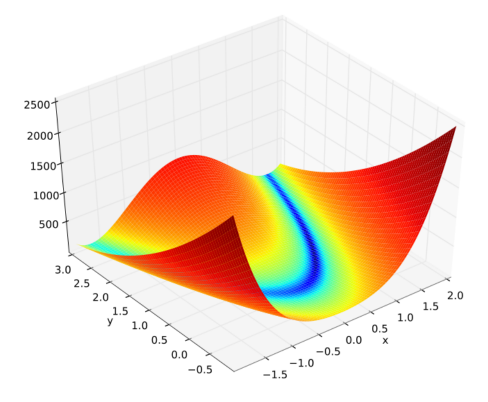
The MOEA Framework contains several basic single-objective optimizers, including a genetic algorithm with elitism (GA), differential evolution (DE), and evolution strategies (ES). These optimizers will work on both single and multi-objective problems. For single-objective problems with one objective, which is typically called the fitness function, these optimizers work as described in the EA literature.

We will demonstrate solving a single-objective problem using the MOEA Framework using the classic Rosenbrock function. The Rosenbrock function is defined as:

$$f(x,y) = (1-x)^2 + 100(y-x^2)^2$$
(11.1)

with an optimum of f(x,y) = 0 at x,y = 1. This function is pictured below ¹:

¹Image courtesy of Wikipedia - https://en.wikipedia.org/wiki/Rosenbrock_function



We will demonstrate solving this function with a genetic algorithm and differential evolution. The process is identical to constructing a multiobjective problem. First, we define the Rosenbrock function by extending the AbstractProblem class. We define this problem to have two decision variables and one objective, as shown below:

```
public class RosenbrockFunction extends AbstractProblem {
2
    public RosenbrockFunction() {
3
      super(2, 1);
4
5
6
7
    @Override
    public void evaluate(Solution solution) {
      double x = EncodingUtils.getReal(solution.getVariable(0));
9
      double y = EncodingUtils.getReal(solution.getVariable(1));
10
      double f = Math.pow(1 - x, 2) + 100*Math.pow(y - Math.pow(x, 2), 2);
      solution.setObjective(0, f);
14
16
    @Override
17
    public Solution newSolution() {
18
      Solution solution = new Solution(2, 1);
19
      solution.setVariable(0, EncodingUtils.newReal(-10.0, 10.0));
20
      solution.setVariable(1, EncodingUtils.newReal(-10.0, 10.0));
21
```

```
22     return solution;
23     }
24     25 }
```

MOEAFramework/book/chapter11/RosenbrockFunction.java

Next, we use the Executor to solve this function using a genetic algorithm with elitism:

```
NondominatedPopulation result = new Executor()
2
           .withAlgorithm("GA")
           .withProblemClass (RosenbrockFunction.class)
3
           .withMaxEvaluations (10000)
           .run();
6
      Solution solution = result.get(0);
8
      System.out.println("x = " +
9
          EncodingUtils.getReal(solution.getVariable(0)));
      System.out.println("y = " +
          EncodingUtils.getReal(solution.getVariable(1)));
12
      System.out.println("f(x,y) = " +
13
           solution.getObjective(0));
14
```

MOEAFramework/book/chapter11/RunBasicGA.java

Since there is a single objective, there is typically a single optimum or several optima with identical fitness values. The NondominatedPopulation will contain only these optimal solutions. Running this example, we get the result:

```
x = 0.9424234853106078

y = 0.8864900554490501

f(x,y) = 0.0035946034840779557
```

Not too bad. However, through experience, differential evolution tends to work much better on continuous, smooth, real-valued problems. So lets give it a shot. Switching to a different optimizer is as simple as specifying a different algorithm:

```
NondominatedPopulation result = new Executor()
.withAlgorithm("DE")
.withProblemClass(RosenbrockFunction.class)
.withMaxEvaluations(10000)
.run();
```

MOEAFramework/book/chapter11/RunBasicDE.java

Our result has improved by an order of magnitude:

```
x = 1.0163380470706236

y = 1.0344827454135896
```

11.2 Multiobjective Problems with Weights

All single-objective optimizers are also designed to work with multiobjective problems with no additional changes. This is accomplished by applying a scalarization function, which converts the multiple objectives into a single fitness value that can be optimized using a single-objective optimizer. One standard approach is with linear weights. Given weights $w_1, w_2, ..., w_M$ for a M-objective problem, the linear fitness is defined as:

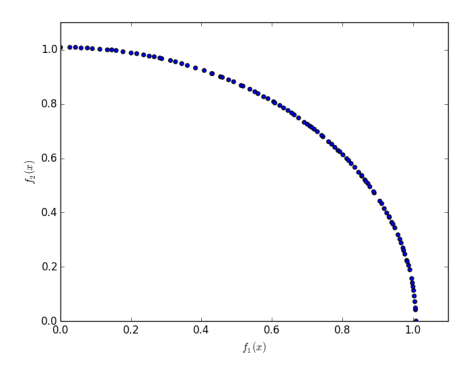
$$f_{linear}(y) = w_1 y_1 + w_2 y_2 + \dots + w_M y_M \tag{11.2}$$

where y_i is the *i*-th objective value. One limitation of using linear weights is the inability to generate concave regions of the Pareto front. An alternative scalarization function that avoids this limitation, called min-max or the Chebyshev function, is defined as:

$$f_{min-max}(y) = \max_{i=1}^{M} (w_i y_i)$$
 (11.3)

That is to say, the fitness of a solution is its largest objective value. Using an optimizer, we then aim to minimize the maximum objective value.

We will now demonstrate solving the 2-objective DTLZ2 problem using various weights. The Pareto front for DTLZ2 is shown below. Note how it forms a concave surface.



We can use the Executor as before. Two optional properties are available: "weights" defines the weights for the scalarizing function, and "method" can be "linear" or "min -max". If no weights are given, the default is (1/M, 1/M, ..., 1/M). Here is an example:

```
new Executor()
.withAlgorithm("GA")
.withProblemClass(DTLZ2.class, 2)
.withMaxEvaluations(10000)
.withProperty("weights", new double[] { 1.0, 0.0 })
.withProperty("method", "min-max")
.run()
```

Lets try different weights to see the effect they have.

```
Executor executor = new Executor()
           .withAlgorithm("GA")
2
           .withProblemClass(DTLZ2.class, 2)
3
           .withMaxEvaluations(10000);
      Solution solution1 = executor
6
           .withProperty("weights", new double[] { 1.0, 0.0 })
           .run()
9
           .get(0);
      Solution solution2 = executor
           .withProperty("weights", new double[] { 0.0, 1.0 })
           .run()
14
           .get(0);
      Solution solution3 = executor
16
           .withProperty("weights", new double[] { 0.5, 0.5 })
17
           .run()
18
           .get(0);
19
20
      Solution solution4 = executor
21
           .withProperty("weights", new double[] { 0.5, 0.5 })
22
           .withProperty("method", "min-max")
23
           .run()
24
           .get(0);
26
      System.out.printf("Solution 1: %.3f %.3f\n",
27
           solution1.getObjective(0), solution1.getObjective(1));
28
      System.out.printf("Solution 2: %.3f %.3f\n",
29
           solution2.getObjective(0), solution2.getObjective(1));
30
      System.out.printf("Solution 3: %.3f %.3f\n",
31
           solution3.getObjective(0), solution3.getObjective(1));
32
      System.out.printf("Solution 4: %.3f %.3f\n",
33
           solution4.getObjective(0), solution4.getObjective(1));
34
```

MOEAFramework/book/chapter11/RunWeightedGA.java

Running this example will produce results similar to:

```
Solution 1: 0.000 1.000
Solution 2: 1.001 0.000
Solution 3: 0.000 1.000
Solution 4: 0.707 0.707
```

Solutions 1, 2, and 3 all use linear weights. Solution 1, which uses the weight (1,0), tries to minimize the first objective. This locates one of the boundary solutions on the Pareto front. Solution 2, which uses the weight (0,1), tries to minimize the second objective, finding the opposite boundary. However, when we supply an intermediate weight, such as (0.5,0.5) for solution 3, we see it produces the same result as solution 1. As discussed earlier, linear weights are unable to identify points within a concave region. On a problem with a concave front, such as DTLZ2, we need to use the min-max method instead. This is shown by solution 4, which identifies the central point on the Pareto front.

11.3 Repeated Single Objectives

As we just saw, we can use a single objective optimizer to find arbitrary points on the Pareto front by supplying different weights. We can expand this process to enumerate the entire Pareto front by supplying many different weights. This process is encapsulated by the RSO, or repeated single objectives, algorithm. Simply put, RSO solves a multiobjective problem by running several single-objective optimizers. Each single-objective optimizer is supplied different weights and run independently from the rest. The solutions found by each single-objective optimizer are finally combined to form the Pareto front.

The code snippet below shows how one would use RSO to solve the DTLZ2 problem.

```
NondominatedPopulation result = new Executor()
           .withAlgorithm("RSO")
2
           .withProblemClass(DTLZ2.class, 2)
3
           .withMaxEvaluations(500000)
           .withProperty("algorithm", "GA")
           .withProperty("instances", 50)
7
           .withProperty("method", "min-max")
           .run();
9
      result.sort(new ObjectiveComparator(0));
      for (Solution solution : result) {
        System.out.printf("%.5f, %.5f\n",
             solution.getObjective(0),
14
             solution.getObjective(1));
15
```

3

MOEAFramework/book/chapter11/RunRSO.java

Note how we can control which single-objective optimizer is utilized with the "algorithm" property. The "instances" property controls how many independent single-objective optimizers are utilized. The max evaluations are spready evenly across all instances. Thus, since we specify a max evaluations of 500,000 and 50 instances, each instance will run for 10,000 function evaluations. The output is shown below:

```
0.00000, 1.00005

0.03993, 0.99931

0.06460, 0.99816

0.08672, 0.99650

0.11275, 0.99384

0.13737, 0.99061

0.16442, 0.98662

0.19144, 0.98168

...

0.99053, 0.13805

0.99376, 0.11273

0.99629, 0.08836

0.99806, 0.06468

0.99931, 0.03875

1.00005, 0.01338

1.00008, 0.00000
```

Now we can ask the question: is it better to run a single-objective optimizer many times or a multiobjective optimizer one time? With the MOEA Framework, we have all the necessary tools to quickly answer this question. For this exercise, we will compare RSO to NSGA-II. Both algorithms will be given the same number of function evaluations and compared using our Analyzer class.

```
Analyzer analyzer = new Analyzer()
           .withProblemClass(DTLZ2.class, 2)
2
           .includeHypervolume();
3
      analyzer.addAll("RSO", new Executor()
           .withSameProblemAs(analyzer)
6
7
           .withAlgorithm("RSO")
           .withMaxEvaluations (50000)
           .withProperty("algorithm", "GA")
9
           .withProperty("instances", 50)
           .withProperty("method", "min-max")
           .runSeeds(50));
      analyzer.addAll("NSGAII", new Executor()
14
           .withSameProblemAs(analyzer)
15
```

```
.withAlgorithm("NSGAII")
.withMaxEvaluations(50000)
.runSeeds(50));

analyzer.printAnalysis();
```

MOEAFramework/book/chapter11/RunRSOvsMOEA.java

The results are quite surprising:

```
NSGAII:
    Hypervolume:
        Min: 0.20864366664800452
        Median: 0.2093293153108441
        Max: 0.20973701273048048
        Count: 50

RSO:
    Hypervolume:
        Min: 0.12487329121903454
        Median: 0.13471352611896967
        Max: 0.14387703753381137
        Count: 50
```

In this example, running an MOEA a single time produces vastly superior results than running many single-objective optimizers. While this quickly demonstrates the power of MOEAs over single-objective optimizers when solving multiobjective problems, a more detailed analysis can be found in (Hughes, 2005).

Appendix A

List of Algorithms

This appendix lists the optimization algorithms provided natively by the MOEA Framework, a short description of the distinct features of the algorithm, and a list of the parameters and default values. Most of these algorithms support a variety of crossover and/or mutation operators. In those cases, refer to Appendix C for a list of the operators and their parameters. Table 4.1 shows a list of these algorithms and their supported types.

In addition to the algorithms listed here, the MOEA Framework also supports third-party libraries via a plugin mechanism. Currently, the MOEA Framework supports optimization algorithms from JMetal and PISA. See Appendix B for more details on the available third-party algorithms. Note that while you can execute these third-party algorithms, some functionality may not be supported. In particular, runtime dynamics can not be collected from JMetal algorithms.

A.1 Multiobjective Optimizers

A.1.1 CMA-ES

CMA-ES is a sophisticated covariance matrix adaptation evolution strategy algorithm for real-valued global optimization (Hansen and Kern, 2004; Igel et al., 2007). CMA-ES produces offspring by sampling a distribution formed by a covariance matrix, hence the name, and updating the covariance matrix based on the surviving offspring. Single and multi-objective variants exist in the literature and both are supported by the MOEA Framework.

Parameters	Description	Default Value
lambda	The offspring population size	100
cc	The cumulation parameter	1
cs	The step size of the cumulation parameter	1
damps	The damping factor for the step size	1
ccov	The learning rate	1
ccovsep	The learning rate when in diagonal-only mode	1
sigma	The initial standard deviation	0.5
diagonalIterations	The number of iterations in which only the covariance diagonal is used	0
indicator	Either "hypervolume" or "epsilon" to specify the use of the hypervolume or additive-epsilon indicator. If unset, crowding distance is used	Unset
initialSearchPoint	Initial guess at the starting location (comma- separated values). If unset, a random initial guess is used	Unset

A.1.2 DBEA

DBEA, or I-DBEA, is the Improved Decomposition-Based Evolutionary Algorithm. DBEA uses the same systematic sampling of reference points as NSGA-III, but utilizes distance along each reference vector to measure convergence and the perpendicular distance to reference vectors to measure diversity (Asafuddoula et al., 2015). DBEA also proposes corner-sort as a means to identify exteme points for normalization. For an M-objective problem, the number of reference points is:

$$H = \begin{pmatrix} M + divisions - 1 \\ divisions \end{pmatrix} \tag{A.1}$$

To use the two-layer approach also used by NSGA-III, replace the divisions parameter with divisionsOuter and divisionsInner.

Parameter	Description	Default Value
divisions	The number of divisions	Problem dependent

A.1.3 ϵ -MOEA

 ϵ -MOEA is a steady-state MOEA that uses ϵ -dominance archiving to record a diverse set of Pareto optimal solutions Deb et al. (2003). The term steady-state means that the algorithm evolves one solution at a time. This is in contrast to generational algorithms, which evolve the entire population every iteration. ϵ -dominance archives are useful since they ensure convergence and diversity throughout search Laumanns et al. (2002). However,

¹Parameter value is derived from other settings. See Igel et al. (2007) for details.

the algorithm requires an additional ϵ parameter which is problem dependent. The ϵ parameter controls the granularity or resolution of the solutions in objective space. Smaller values produce larger, more dense sets while larger values produce smaller sets. In general, the ϵ values should be chosen to yield a moderately-sized Pareto approximate set.

Parameter	Description	Default Value
populationSize	The size of the population	100
epsilon	The ϵ values used by the ϵ -dominance archive,	Problem dependent
	which can either be a single value or a comma-	
	separated array	

A.1.4 ϵ -NSGA-II

 ϵ -NSGA-II combines the generational search of NSGA-II with the guaranteed convergence provided by an ϵ -dominance archive Kollat and Reed (2006). It also features randomized restarts to enhance search and find a diverse set of Pareto optimal solutions. During a random restart, the algorithm empties the current population and fills it with new, randomly-generated solutions.

Parameter	Description	Default Values
populationSize	The size of the population	100
epsilon	The ϵ values used by the ϵ -dominance	Problem depen-
	archive, which can either be a single value	dent
	or a comma-separated array	
injectionRate	Controls the percentage of the population	0.25
	after a restart this is "injected", or copied,	
	from the ϵ -dominance archive	
windowSize	Frequency of checking if a randomized	100
	restart should be triggered (number of it-	
	erations)	
maxWindowSize	The maximum number of iterations be-	100
	tween successive randomized restarts	
${\rm minimum Population Size}$	The smallest possible population size	100
	when injecting new solutions after a ran-	
	domized restart	
maximum Population Size	The largest possible population size when	10000
	injecting new solutions after a randomized	
	restart	

A.1.5 GDE3

GDE3 is the third version of the generalized differential evolution algorithm Kukkonen and Lampinen (2005). The name differential evolution comes from how the algorithm evolves

offspring. It randomly selects three parents. Next, it computes the difference (the differential) between two of the parents. Finally, it offsets the remaining parent by this differential.

Parameter	Description	Default Values
populationSize	The size of the population	100
de.crossoverRate	The crossover rate for differential evolution	0.1
de.stepSize	Control the size of each step taken by differential	0.5
	evolution	

A.1.6 IBEA

IBEA is a indicator-based MOEA that uses the hypervolume performance indicator as a means to rank solutions Zitzler and Künzli (2004). Indicator-based algorithms are based on the idea that a performance indicator, such as hypervolume or additive ϵ -indicator, highlight solutions with desirable qualities. The primary disadvantage of indicator-based methods is that the calculation of the performance indicator can become computationally expensive, particularly as the number of objectives increases.

Parameter	Description	Default Value
populationSize	The size of the population	100
indicator	The indicator function (e.g., "hypervolume", "	"hypervolume"
	epsilon")	

A.1.7 MOEA/D

MOEA/D is a relatively new optimization algorithm based on the concept of decomposing the problem into many single-objective formulations. Several version of MOEA/D exist in the literature. The most common variant seen in the literature, MOEA/D-DE (Li and Zhang, 2009), is the default implementation in the MOEA Framework.

An extension to MOEA/D-DE variant called MOEA/D-DRA introduced a utility function that aimed to reduce the amount of "wasted" effort by the algorithm (Zhang et al., 2009). This variant is enabled by setting the updateUtility parameter to a non-zero value.

Parameter	Description	Default
		Value
populationSize	The size of the population	100
de.crossoverRate	The crossover rate for differential evolution	0.1
de.stepSize	Control the size of each step taken by differential evolution	0.5
pm.rate	The mutation rate for polynomial mutation	1/N
pm. distribution Index	The distribution index for polynomial mutation	20.0
neighborhood Size	The size of the neighborhood used for mating,	0.1
	given as a percentage of the population size	
delta	The probability of mating with an individual from	0.9
	the neighborhood versus the entire population	
eta	The maximum number of spots in the population	0.01
	that an offspring can replace, given as a percentage	
	of the population size	
updateUtility	The frequency, in generations, at which utility val-	Unset
	ues are updated. If set, this uses the MOEA/D-	
	DRA variant; if unset, then then MOEA/D-DE	
	variant is used	

A.1.8 MSOPS

MSOPS is the Multiple Single-Objective Pareto Search algorithm (Hughes, 2003). MSOPS works by enumerating k reference vectors and applying a rank ordering based on two aggregate functions: weighted min-max and vector angle distance scaling (VADS). Solutions with higher rankings with respect to both metrics are preferred. MSOPS only supports real-valued solutions using differential evolution.

Parameter	Description	Default Value
populationSize	The size of the population	100
numberOfWeights	The number of weight vectors	50
de.crossoverRate	The crossover rate for differential evolution	0.1
de.stepSize	Control the size of each step taken by differential	0.5
	evolution	

A.1.9 NSGA-II

NSGA-II is one of the first and most widely used MOEAs (Deb et al., 2000). It enhanced it predecessor, NSGA, by introducing fast non-dominated sorting and using the more computationally efficient crowding distance metric during survival selection.

Parameter	Description	Default Value
populationSize	The size of the population	100
withReplacement	Uses binary tournament selection with (true) or	true
	without (false) replacement	

A.1.10 NSGA-III

NSGA-III is the many-objective successor to NSGA-II, using reference points to direct solutions towards a diverse set (Deb and Jain, 2014). The number of reference points is controlled by the number of objectives and the divisions parameter. For an *M*-objective problem, the number of reference points is:

$$H = \begin{pmatrix} M + divisions - 1 \\ divisions \end{pmatrix} \tag{A.2}$$

The authors also propose a two-layer approach for divisions for many-objective problems where an outer and inner division number is specified. To use the two-layer approach, replace the divisions parameter with divisionsOuter and divisionsInner.

Parameter	Description	Default Value
populationSize	The size of the population. If unset, the popula-	Unset
	tion size is equal to the number of reference points	
divisions	The number of divisions	Problem dependent

A.1.11 OMOPSO

OMOPSO is a multiobjective particle swarm optimization algorithm that includes an ϵ -dominance archive to discover a diverse set of Pareto optimal solutions (Sierra and Coello Coello, 2005). This implementation of OMOPSO differs slightly from the original author's implementation in JMetal due to a discrepancy between the author's code and the paper. The paper returns the ϵ -dominance archive while the code returns the leaders. This discrepancy causes a small difference in performance.

Parameter	Description	Default Value
populationSize	The size of the population	100
archiveSize	The size of the archive	100
maxEvaluations	The maximum number of evaluations for	25000
	adapting non-uniform mutation	
mutationProbability	The mutation probability for uniform and	1/N
	non-uniform mutation	
perturbation Index	Controls the shape of the distribution for	0.5
	uniform and non-uniform mutation	
epsilon	The ϵ values used by the ϵ -dominance	Problem dependent
	archive	

A.1.12 PAES

PAES is a multiobjective version of evolution strategy (Knowles and Corne, 1999). PAES tends to underperform when compared to other MOEAs, but it is often used as a baseline algorithm for comparisons. Like PESA-II, PAES uses the adaptive grid archive to maintain a fixed-size archive of solutions.

Parameter	Description	Default
		Value
archiveSize	The size of the archive	100
bisections	The number of bisections in the adaptive grid	8
	archive	
pm.rate	The mutation rate for polynomial mutation	1/N
pm. distribution Index	The distribution index for polynomial mutation	20.0

A.1.13 PESA-II

PESA-II is another multiobjective evolutionary algorithm that tends to underperform other MOEAs but is often used as a baseline algorithm in comparative studies (Corne et al., 2001). It is the successor to PESA (Corne and Knowles, 2000). Like PAES, PESA-II uses the adaptive grid archive to maintain a fixed-size archive of solutions.

Parameter	Description	Default Value
populationSize	The size of the population	10
archiveSize	The size of the archive	100
bisections	The number of bisections in the adaptive grid	8
	archive	

A.1.14 Random

The random search algorithm simply randomly generates new solutions uniformly throughout the search space. It is not intended as an "optimization algorithm" per se, but as a way to compare the performance of other MOEAs against random search. If an optimization algorithm can not beat random search, then continued use of that optimization algorithm should be questioned.

Parameter	Description	Default Value
populationSize	This parameter only has a use when parallelizing	100
	evaluations; it controls the number of solutions	
	that are generated and evaluated in parallel	
epsilon	The ϵ values used by the ϵ -dominance archive,	Unset
	which can either be a single value or a comma-	
	separated array (this parameter is optional)	

A.1.15 RSO

The repeated single objectives (RSO) algorithm solves multiobjective problems by running several single-objective optimizers independently with varying weights (Hughes, 2005). Any of the single-objective optimizers supported by the MOEA Framework can be utilized, and any properties supported by that optimizer can be defined. RSO is a useful tool for comparing single and multiobjective optimizers. The maximum number of evaluations is spread evenly across each single-objective instance.

Parameter	Description	Default Value
algorithm	The single-objective optimizer	"GA"
method	The scalarizing method	"min-max"
instances	The number of single-objective optimizers	100

A.1.16 RVEA

The reference vector guided evolutionary algorithm (RVEA) has many similarities with NSGA-III, but avoids use of Pareto dominance and uses an angle-penalized distance function for survival selection (Cheng et al., 2016). RVEA only works on problems with at least two objectives and can only use genetic operators requiring two parents. Like NSGA-III, the number of reference vectors is controlled by the number of objectives and the divisions parameter. For an M-objective problem, the number of reference vectors is:

$$H = \begin{pmatrix} M + divisions - 1 \\ divisions \end{pmatrix} \tag{A.3}$$

To use the two-layer approach, replace the divisions parameter with divisionsOuter and divisionsInner.

Parameter	Description	Default Value
populationSize	The size of the population. If unset, the pop-	Unset
	ulation size is equal to the number of refer-	
	ence vectors	
divisions	The number of divisions	Problem dependent
alpha	Controls the rate of change in the angle-	2
	penalized distance function	
adaptFrequency	The frequency (in generations) in which the	maxEvaluations /
	weights are adapted / scaled	(populationSize *
		10)

A.1.17 SMPSO

SMPSO is a multiobjective particle swarm optimization algorithm (Nebro et al., 2009).

Parameter	Description	Default
		Value
populationSize	The size of the population	100
archiveSize	The size of the archive	100
pm.rate	The mutation rate for polynomial mutation	1/N
pm. distribution In dex	The distribution index for polynomial mutation	20.0

A.1.18 SMS-EMOA

SMS-EMOA is an indicator-based MOEA that uses the volume of the dominated hypervolume to rank individuals (Beume et al., 2007).

Parameter	Description	Default Value
populationSize	The size of the population	100
offset	The reference point offset for computing hypervol-	100
	ume	

A.1.19 SPEA2

SPEA2 is an older but popular benchmark MOEA that uses the so-called "strength-based" method for ranking solutions (Zitzler et al., 2002a). The general idea is that the strength or quality of a solution is related to the strength of solutions it dominates.

Parameter	Description	Default Value
populationSize	The size of the population	100
offspringSize	The number of offspring generated every iteration	100
k	Crowding is based on the distance to the k -th near-	1
	est neighbor	

A.1.20 VEGA

VEGA is considered the earliest documented MOEA. While we provide support for VEGA, other MOEAs should be preferred as they exhibit better performance. VEGA is provided for its historical significance (Schaffer, 1985).

Parameter	Description	Default Value
populationSize	The size of the population	100

A.2 Single-Objective Optimizers

In addition to the multiobjective optimizers listed above, the MOEA Framework supports several single-objective optimizers. These single-objective optimizers can be used to solve both single and multiobjective problems. For multiobjective problems, additional weighting properties are provided.

All single objective algorithms support the "weights" and "method" properties. Both are optional. If not weights are given, the default is equal weights. "method" can either be "linear" or "min-max".

A.2.1 GA

GA is the standard genetic algorithm with elitism (Holland, 1975). A single elite individual is guaranteed to survive between generations.

Parameter	Description	Default Value
populationSize	The size of the population	100
method	The scalarization method	"linear"
weights	The scalarization weights	Equal weights

A.2.2 ES

ES is the standard (1 + 1) evolution strategies algorithm (Rechenberg, 1971). ES only supports real-valued variables. This means the population is size 1 and only 1 offspring is generated each iteration. The fittest solution survives to the next iteration. Additionally, ES uses a self-adaptive variation operator.

Parameter	Description	Default Value
populationSize	The size of the population	100
method	The scalarization method	"linear"
weights	The scalarization weights	Equal weights

A.2.3 DE

DE is the standard differential evolution algorithm (Storn and Price, 1997). DE only supports real-valued variables using the differential evolution operator. DE works by calculating the difference between two randomly-selected points and applying that difference to a third point.

Parameter	Description	Defa	ult Value
populationSize	The size of the population	100	
method	The scalarization method	"lir	near"
weights	The scalarization weights	Equa	al weights
de.crossoverRate	The DE crossover rate	See	documenta-
		tion	
de.stepSize	The DE step size	See	documenta-
		tion	

Appendix B

Additional Algorithms

In addition to the algorithms implemented natively within the MOEA Framework, as listed in Appendix A, the MOEA Framework can also run third-party optimization algorithms via a plugin mechanism. The current installation of MOEA Framework supports JMetal and the PISA Framework. JMetal algorithms are included by default. PISA Framework algorithms must be downloaded and compiled separately.

B.1 JMetal Algorithms

JMetal supports only the real-valued, binary, and permutation encodings. Each of the descriptions below will indicate which of these encodings, if any, the algorithm supports. For each encoding, a different set of parameters are available. For real-valued encodings, the additional parameters are:

Parameters	Description	Default
		Value
sbx.rate	The crossover rate for simulated binary crossover	1.0
${\bf sbx. distribution Index}$	The distribution index for simulated binary	15.0
	crossover	
pm.rate	The mutation rate for polynomial mutation	1.0/L
pm.distributionIndex	The distribution index for polynomial mutation	20.0

For binary encodings, the additional parameters are:

Parameters	Description	Default Value
1x.rate	The crossover rate for single-point crossover	0.9
bf.rate	The mutation rate for bit flip mutation	1.0/L

For permutation encodings, the additional parameters are:

Table B.1: List of available third-party optimization algorithms.

Algorithm	Туре	Real	Binary	Permutation	Grammar	Program	Constraints	Provider
AbYSS	Scatter Search	Yes	$N_{\rm O}$	No	N_{O}	N_{0}		JMetal
CellDE	Differential Evolution	Yes	No	No	$N_{\rm O}$	N_{0}		JMetal
DENSEA	Genetic Algorithm	Yes	Yes	Y_{es}	N_{0}	N_{0}		JMetal
ECEA	Genetic Algorithm	Yes	Yes	Yes	Yes	Yes		PISA
${ m FastPGA}$	Genetic Algorithm	Yes	Yes	Yes	No	N_{0}		JMetal
FEMO	Genetic Algorithm	Yes	Yes	Yes	Yes	Yes		PISA
$_{ m HypE}$	Indicator-Based	Yes	Yes	Yes	Yes	Yes		PISA
MOCell	Cellular	Yes	Yes	Yes	No	No		JMetal
MOCHC	CHC Algorithm	No	Yes	No	No	$N_{\rm O}$	Yes	JMetal
SEMO2	Genetic Algorithm	Yes	Yes	Yes	Yes	Yes		PISA
$_{ m SHV}$	Indicator-Based	Yes	Yes	Yes	Yes	Yes		PISA
SIBEA	Indicator-Based	Yes	Yes	Yes	Yes	Yes		PISA
SPAM	Indicator-Based	Yes	Yes	Yes	Yes	Yes		PISA

Parameters	Description	Default Value
pmx.rate	The crossover rate for PMX crossover	1.0
swap.rate	The mutation rate for the swap operator	0.35

Each of the supported JMetal algorithms and their parameters are detailed below.

AbYSS

AbYSS is a hybrid scatter search algorithm that uses genetic algorithm operators (Nebro et al., 2008). Use the string "AbYSS" when creating instances of this algorithm with the Executor. Only real-valued decision variables are supported. The following parameters are available:

Parameters	Description	Default Value
populationSize	The size of the population	20
archiveSize	The size of the archive	100
refSet1Size	The size of the first reference set	10
refSet2Size	The size of the second reference set	10
improvement Rounds	The number of iterations that the local search op-	1
	erator is applied	

B.1.1 CellDE

CellDE is a hybrid cellular genetic algorithm (meaning mating only occurs among neighbors) combined with differential evolution (Durillo et al., 2008). Use the string "CellDE" when creating instances of this algorithm with the Executor. CellDE defines its own parameters for its real-valued operators as listed below:

Parameters	Description	Default Value
populationSize	The size of the population	100
archiveSize	The size of the archive	100
feedBack	Controls the number of solutions from the archive	20
	that are fed back into the population	
${\rm de. crossoverRate}$	The crossover rate for differential evolution	0.1
de.stepSize	Control the size of each step taken by differential	0.5
	evolution	

B.1.2 DENSEA

DENSEA is the duplicate elimination non-domination sorting evolutionary algorithm (Greiner et al., 2006). Use the string "DENSEA" when creating instances of this algorithm with the Executor. DENSEA supports real-valued, binary, and permutation encodings. The following parameters are available:

Parameters	Description	Default Value
populationSize	The size of the population	100

B.1.3 FastPGA

FastPGA is a genetic algorithm that uses adaptive population sizing to solve time-consuming problems more efficiently (Eskandari et al., 2007). Use the string "FastPGA" when creating instances of this algorithm with the Executor. FastPGA supports real-valued, binary, and permutation encodings. The following parameters are available:

Parameters	Description	Default Value
maxPopSize	The maximum size of the population	100
initialPopulationSize	The initial size of the population	100
a	Constant controlling the population size	20.0
b	Multiplier controlling the population size	1.0
c	Constant controlling the number of offspring	20.0
d	Multiplier controlling the number of offspring	0.0
termination	If 0, the algorithm terminates early if all solutions	1
	like on the Pareto optimal front	

B.1.4 MOCell

MOCell is the multiobjective version of a cellular genetic algorithm (Nebro et al., 2007b). Use the string "MOCell" when creating instances of this algorithm with the Executor. MOCell supports real-valued, binary, and permutation encodings. The following parameters are available:

Parameters	Description	Default Value
populationSize	The size of the population	100
archiveSize	The size of the archive	100
feedBack	Controls the number of solutions from the archive	20
	that are fed back into the population	

B.1.5 MOCHC

MOCHC is a genetic algorithm that combines a conservative selection strategy with highly disruptive recombination, which unlike traditional MOEAs aims to produce offspring that are maximally different from both parents (Nebro et al., 2007a). Use the string "MOCHC" when creating instances of this algorithm with the Executor. MOCHC defines its own parameters for its search operators as listed below:

Parameters	Description	Default
		Value
initialConvergenceCount	The threshold (as a percent of the number of bits in	0.25
	the encoding) used to determine similarity between	
	solutions	
preserved Population	The percentage of the population that does not	0.05
	undergo cataclysmic mutation	
${\it convergenceValue}$	The convergence threshold that determines when	3
	cataclysmic mutation is applied	
populationSize	The size of the population	100
hux.rate	The crossover rate for the highly disruptive recom-	1.0
	bination operator	
bf.rate	The mutation rate for bit-flip mutation	0.35

B.2 PISA Algorithms

The MOEA Framework has been extensively tested with the PISA algorithms listed in Table B.1. PISA algorithms are provided by a third-party and are not distributed by the MOEA Framework, but the MOEA Framework can be configured to run such optimization algorithms. This section describes how to connect the MOEA Framework to a PISA algorithm.

The Platform and Programming Language Independent Interface for Search Algorithms (PISA), available at http://www.tik.ee.ethz.ch/pisa/, is a language-neutral programming interface for creating search and optimization algorithms (Bleuler et al., 2003). PISA specifies three components required for search algorithms:

- 1. selectors, which define the optimization algorithms;
- 2. variators, which define the optimization problems; and
- 3. a communication scheme using plaintext files.

This design offers several benefits. First, it clearly demarcates the responsibilities of algorithm experts from those of application engineers. The algorithm experts focus on designing and improving the behavior of optimization algorithms (i.e., selectors), whereas application engineers are responsible for encoding the details of their problem (i.e., variators). Second, the file-based communication scheme employed by PISA permits selectors and variators to be written in nearly any programming language, which may be paired with a selector/variator written in a completely different language. Third, the standardized communication scheme enables plug-and-play integration, allowing one module to be swapped out for another with little to no effort. For instance, one selector may be replaced by another simply by changing which executable is run.

The fundamental drawback of PISA is a result of its reliance on a file-based communication scheme. File access on modern computers remains a (relatively) slow operation, which

is further exacerbated by the need to constantly poll the communication files for changes. Nonetheless, PISA opens the door to a number of optimization algorithms, including:

- 1. Set Preference Algorithm for Multiobjective Optimization (SPAM)
- 2. Sampling-Based Hypervolume-Oriented Algorithm (SHV)
- 3. Simple Indicator-Based Evolutionary Algorithm (SIBEA)
- 4. Hypervolume Estimation Algorithm for Multiobjective Optimization (HypE)
- 5. Simple Evolutionary Multiobjective Optimizer (SEMO2)
- 6. Fair Evolutionary Multiobjective Optimizer (FEMO)
- 7. Epsilon-Constraint Evolutionary Algorithm (ECEA)

For this reason, the MOEA Framework provides the support necessary to integrate with the PISA library. The PISAAlgorithm class acts as a variator, which knows how to communicate with a PISA selector using the file-based communication protocol.

B.2.1 Adding a PISA Selector

A standardized method for adding PISA selectors to the MOEA Framework is provided. The steps required to add a new PISA selector are:

- 1. Download and extract a PISA selector
- 2. Edit global.properties
 - (a) Add the name of the selector, NAME, to the comma-separated list of available PISA selectors in org.moeaframework.algorithm.pisa.algorithms
 - (b) Add the property org.moeaframework.algorithm.pisa.NAME.command, which points to the program executable which starts the PISA selector
 - (c) Provide a list of parameters, in the order required by the PISA selector, with the property org.moeaframework.algorithm.pisa.NAME.parameters
 - (d) For each of the listed parameters, PARAM, add the property org. moeaframework.algorithm.pisa.NAME.PARAM to set the default value for the parameter. It is not necessary to list the seed parameter

For example, if we install the HypE selector, we would first download the HypE binaries from http://www.tik.ee.ethz.ch/pisa/. These binaries are typically packaged as a compressed file (.zip or .tar.gz). Next, extract the contents of this compressed file into the MOEA Framework installation folder. In this example, we extracted the contents to the folder pisa/hype_win. Finally, add the following lines to the global.properties file:

```
org.moeaframework.algorithm.pisa.algorithms=HypE
org.moeaframework.algorithm.pisa.HypE.command = ./pisa/hype_win/hype.exe
org.moeaframework.algorithm.pisa.HypE.parameters = seed, tournament, mating,
    bound, nrOfSamples
org.moeaframework.algorithm.pisa.HypE.parameter.tournament = 5
org.moeaframework.algorithm.pisa.HypE.parameter.mating = 1
org.moeaframework.algorithm.pisa.HypE.parameter.bound = 2000
org.moeaframework.algorithm.pisa.HypE.parameter.nrOfSamples = -1
```

Once completed, you should be able to run the diagnostic tool and confirm that HypE appears in the list of available algorithms. Additionally, HypE may be referenced throughout the MOEA Framework wherever the algorithm is specified as a string, such as:

```
new Executor()
    .withProblem("Kursawe")
    .withAlgorithm("HypE")
    .withMaxEvaluations(10000)
    .withProperty("bound", 1000)
    .withProperty("tournament", 2)
    .run();
```

B.2.2 Troubleshooting

I'm attempting to run the PISA algorithm, but nothing is happening. The task manager shows the PISA executable is running, but shows 0% CPU usage.

The MOEA Framework uses your system's default temporary directory as the location of the files used to communicate with the PISA selector. Some PISA selectors do not support paths containing spaces, and the path to the default temporary directory on older versions of Windows contains spaces. This causes a miscommunication between the MOEA Framework and PISA, which generally causes the MOEA Framework and PISA executables to stall.

The easiest workaround is to override the temporary directory location so that the space is removed. This can be accomplished by editing the <code>mglobal.properties</code> file and adding the line:

```
java.io.tmpdir = C:/temp/
```

PISA runs fine for a while, but eventually crashes with the message "Assertion failed: fp != null".

Some antivirus software is known to interfere with the file-based communication protocol used by PISA. Antivirus programs which actively monitor files for changes may lock the file during a scan, potentially blocking access by the PISA selector. Most PISA selectors will crash with the obscure message "Assertion failed: fp != null".

To verify this as the cause, you may temporarily disable your antivirus software and re-run the program. Once verified, a permanent solution involves adding an exception to the antivirus software to prevent scanning the PISA communication files. To implement this solution, first define the location of temporary files by adding the following line to the global properties file:

```
java.io.tmpdir = C:/temp/
```

Then add an exception to your antivirus software to disable scanning files located in this directory.

(Note: Disabling an antivirus program from scanning a folder will leave its contents unprotected. Follow these steps at your own risk.)

Appendix C

List of Variation Operators

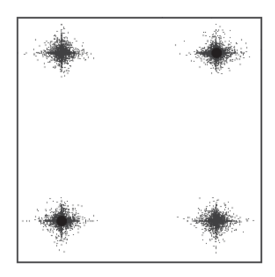
The following crossover and mutation operators are supported by the MOEA Framework.

Representation	Type	Abbr.
Real / Integer	Simulated Binary Crossover	sbx
Real / Integer	Polynomial Mutation	pm
Real / Integer	Differential Evolution	de
Real / Integer	Parent-Centric Crossover	pcx
Real / Integer	Simplex Crossover	spx
Real / Integer	Unimodal Normal Distribution Crossover	undx
Real / Integer	Uniform Mutation	um
Real / Integer	Adaptive Metropolis	am
Binary	Half-Uniform Crossover	hux
Binary	Bit Flip Mutation	bf
Permutation	Partially-Mapped Crossover	pmx
Permutation	Element Insertion	insertion
Permutation	Element Swap	swap
Subset	Subset Crossover	SSX
Subset	Subset Replacement	replace
Grammar	Single-Point Crossover for Grammars	gx
Grammar	Uniform Mutation for Grammars	gm
Program	Branch (Subtree) Crossover	bx
Program	Point Mutation	ptm
Any	Single-Point Crossover	1x
Any	Two-Point Crossover	2x
Any	Uniform Crossover	ux

C.1 Real-Valued Operators

Simulated Binary Crossover (SBX)

SBX attempts to simulate the offspring distribution of binary-encoded single-point crossover on real-valued decision variables (Deb and Agrawal, 1994). It accepts two parents and produces two offspring. An example of this distribution, which favors offspring nearer to the two parents, is shown below.



The distribution index controls the shape of the offspring distribution. Larger values for the distribution index generates offspring closer to the parents.

Parameters	Description	Default
		Value
sbx.rate	The probability that the SBX operator is applied	1.0
	to a decision variable	
sbx. distribution Index	The shape of the offspring distribution	15.0

Polynomial Mutation (PM)

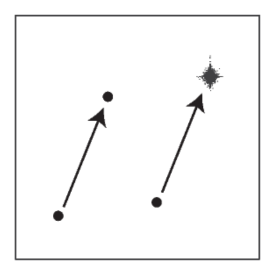
PM attempts to simulate the offspring distribution of binary-encoded bit-flip mutation on real-valued decision variables (Deb and Goyal, 1996). Similar to SBX, PM favors offspring nearer to the parent. It is recommended each decision variable is mutated with a probability of 1/N, where N is the number of decision variables. This results in one mutation per offspring on average.

The distribution index controls the shape of the offspring distribution. Larger values for the distribution index generates offspring closer to the parents.

Parameters	Description	Default
		Value
pm.rate	The probability that the PM operator is applied	1/N
	to a decision variable	
pm. distribution Index	The shape of the offspring distribution (larger val-	20.0
	ues produce offspring closer to the parent)	

Differential Evolution (DE)

Differential evolution works by randomly selecting three distinct individuals from a population. A difference vector is calculated between the first two individuals (shown as the left-most arrow in the figure below), which is subsequently applied to the third individual (shown as the right-most arrow in the figure below).

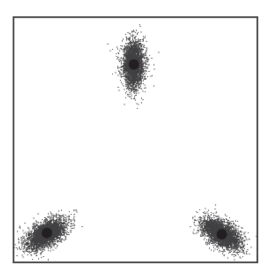


The scaling factor parameter adjusts the magnitude of the difference vector, allowing the user to decrease or increase the magnitude in relation to the actual difference between the individuals (Storn and Price, 1997). The crossover rate parameter controls the fraction of decision variables which are modified by the DE operator.

Parameters	Description	Default Value
de.crossoverRate	The fraction of decision variables modified by the	0.1
	DE operator	
de.stepSize	The scaling factor or step size used to adjust the	0.5
	length of each step taken by the DE operator	

Parent Centric Crossover (PCX)

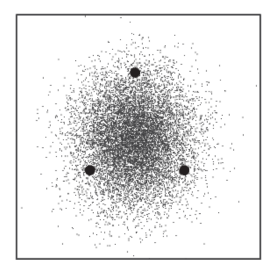
PCX is a multiparent operator, allowing a user-defined number of parents and offspring (Deb et al., 2002). Offspring are clustered around the parents, as depicted in the figure below.



Parameters	Description	Default Value
pcx.parents	The number of parents	10
pcx.offspring	The number of offspring generated by PCX	2
pcx.eta	The standard deviation of the normal distribution	0.1
	controlling the spread of solutions in the direction	
	of the selected parent	
pcx.zeta	The standard deviation of the normal distribution	0.1
	controlling the spread of solutions in the directions	
	defined by the remaining parents	

Unimodal Distribution Crossover (UNDX)

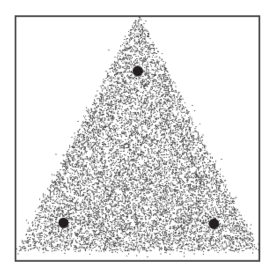
UNDX is a multiparent operator, allowing a user-defined number of parents and offspring (Kita et al., 1999; Deb et al., 2002). Offspring are centered around the centroid, forming a normal distribution whose shape is controlled by the positions of the parents, as depicted in the figure below.



Parameters	Description	Default Value
undx.parents	The number of parents	10
undx.offspring	The number of offspring generated by UNDX	2
undx.zeta	The standard deviation of the normal distribution	0.5
	controlling the spread of solutions in the orthogo-	
	nal directions defined by the parents	
undx.eta	The standard deviation of the normal distribution	0.35
	controlling the spread of solutions in the remaining	
	orthogonal directions not defined by the parents.	
	This value is divided by \sqrt{N} prior to use, where	
	N is the number of decision variables.	

Simplex Crossover (SPX)

SPX is a multiparent operator, allowing a user-defined number of parents and offspring (Tsutsui et al., 1999; Higuchi et al., 2000). The parents form a convex hull, called a simplex. Offspring are generated uniformly at random from within the simplex. The expansion rate parameter can be used to expand the size of the simplex beyond the bounds of the parents. For example, the figure below shows three parent points and the offspring distribution, clearly filling an expanded triangular simplex.



Parameters	Description	Default Value
spx.parents	The number of parents	10
spx.offspring	The number of offspring generated by UNDX	2
spx.epsilon	The expansion rate	3

Uniform Mutation (UM)

Each decision variable is mutated by selecting a new value within its bounds uniformly at random. The figure below depicts the offspring distribution. It is recommended each decision

variable is mutated with a probability of 1/N, where N is the number of decision variables. This results in one mutation per offspring on average.

Parameters	Description	Default Value
um.rate	The probability that the UM operator is applied	1/N
	to a decision variable	

Adaptive Metropolis (AM)

AM is a multiparent operator, allowing a user-defined number of parents and offspring (Vrugt and Robinson, 2007; Vrugt et al., 2009). AM produces normally-distributed clusters around each parent, where the shape of the distribution is controlled by the covariance of the parents.

Internally, the Cholesky decomposition is used to update the resulting offspring distribution. Cholesky decomposition requires that its input be positive definite. In order to guarantee this condition is satisfied, all parents must be unique. In the event that the positive definite condition is not satisfied, no offspring are produced and an empty array is returned by

Parameters	Description	Default Value
am.parents	The number of parents	10
am.offspring	The number of offspring generated by AM	2
am.coefficient	The jump rate coefficient, controlling the standard	2.4
	deviation of the covariance matrix. The actual	
	jump rate is calculated as $(am.coefficient/\sqrt{n})^2$	

C.2 Binary / Bit String Operators

Half Uniform Crossover (HUX)

Half-uniform crossover (HUX) operator. Half of the non-matching bits are swapped between the two parents.

Parameters	Description	Default Value
hux.rate	The probability that the UM operator is applied	1.0
	to a binary decision variable	

Bit Flip Mutation (BF)

Each bit is flipped (switched from a 0 to a 1, or vice versa) using the specified probability.

Parameters	Description	Default Value
bf.rate	The probability that a bit is flipped	0.01

C.3 Permutations

Partially Mapped Crossover (PMX)

PMX is similar to two-point crossover, but includes a repair operator to ensure the offspring are valid permutations (Goldberg and Jr., 1985).

Parameters	Description	Default Value
pmx.rate	The probability that the PMX operator is applied	1.0
	to a permutation decision variable	

Insertion Mutation

Randomly selects an entry in the permutation and inserts it at some other position in the permutation.

Parameters	Description	Default Value
insertion.rate	The probability that the insertion operator is ap-	0.3
	plied to a permutation decision variable	

Swap Mutation

Randomly selects two entries in the permutation and swaps their position.

Parameters	Description	Default Value
swap.rate	The probability that the swap operator is applied	0.3
	to a permutation decision variable	

C.4 Subsets

Subset Crossover (SSX)

SSX is similar to HUX crossover for binary strings, where half of the non-matching members are swapped between the two subsets.

Parameters	Description	Default Value
ssx.rate	The probability that the SSX operator is applied	0.9
	to a subset decision variable	

Replace Mutation

Randomly replaces one of the members in the subset with a non-member.

Parameters	Description	Default Value
replace.rate	The probability that the replace operator is ap-	0.3
	plied to a subset decision variable	

C.5 Grammars

Grammar Crossover (GX)

Single-point crossover for grammars. A crossover point is selected in both parents with the tail portions swapped.

Parameters	Description	Default Value
gx.rate	The probability that the GX operator is applied	1.0
	to a grammar decision variable	

Grammar Mutation (GM)

Uniform mutation for grammars. Each integer codon in the grammar representation is uniformly mutated with a specified probability.

Parameters	Description	Default Value
gm.rate	The probability that the GM operator is applied	1.0
	to a grammar decision variable	

C.6 Program Tree

Subtree Crossover (BX)

Exchanges a randomly-selected subtree from one program with a compatible, randomly-selected subtree from another program.

Parameters	Description	Default Value
gm.rate	The probability that the BX operator is applied to	1.0
	a program tree decision variable	

Point Mutation (PTM)

Mutates a program by randomly selecting nodes in the expression tree and replacing the node with a new, compatible, randomly-selected node.

Parameters	Description	Default Value
gm.rate	The probability that the PTM operator is applied	1.0
	to a program tree decision variable	

C.7 Generic Operators

Generic operators can be applied to any type. They work by simply swapping the value of the decision variable between the parents.

One-Point Crossover (1X)

A crossover point is selected and all decision variables to the left/right are swapped between the two parents. The two children resulting from this swapping are returned.

Parameters	Description	Default Value
1x.rate	The probability that one-point crossover is applied	1.0
	to produce offspring	

Two-Point Crossover (2X)

Two crossover points are selected and all decision variables between the two points are swapped between the two parents. The two children resulting from this swapping are returned.

Parameters	Description	Default Value
2x.rate	The probability that two-point crossover is applied	1.0
	to produce offspring	

Uniform Crossover (UX)

Crossover operator where each index is swapped with a specified probability.

Parameters	Description	Default Value
ux.rate	The probability that uniform crossover is applied	1.0
	to produce offspring	

Appendix D

List of Problems

The following test problems are packaged with the MOEA Framework. Problems marked with † have maximized objectives. The MOEA Framework negates the values of maximized objectives. Note that while many problems have similar Pareto fronts, their underlying problem definitions and properties can differ greatly.

Problem	# of Vars	# of Objs	# of Constrs	Type	Pareto Front
Belegundu	2	2	2	Real	
Binh	2	2	0	Real	8 S S S S S S S S S S S S S S S S S S S
Binh2	2	2	2	Real	\$ 3 8 9 0 20 40 00 80 100 100

Problem	# of Vars	# of Objs	# of Constrs	Type	Pareto Front
Binh3	2	3	0	Real	\$ 10 to 10 t
Binh4	2	3	2	Real	5 2 4 4 5 4 5 4 5 4 5 4 5 4 5 4 5 4 5 4
CF1	10	2	1	Real	3 3 3 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5
CF2	10	2	1	Real	
CF3	10	2	1	Real	2
CF4	10	2	1	Real	S 3 3 3 3 3 3 4 5 5 7 10

Problem	# of Vars	# of Objs	# of Constrs	Type	Pareto Front
${ m CF5}$	10	2	1	Real	3 3 3 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5
CF6	10	2	2	Real	3 3 3 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0
CF7	10	2	2	Real	3 3 3 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0
CF8	10	3	1	Real	\$ 3 2 40 52 64 64 66 10 60 10 60 10 10 10 10 10 10 10 10 10 10 10 10 10
CF9	10	3	1	Real	
CF10	10	3	1	Real	5 3 3 41 50 41 41 50 41 50 41 50 50 50 50 50 50 50 50 50 50 50 50 50

Problem	# of Vars	# of Objs	# of Constrs	Type	Pareto Front
$\mathrm{DTLZ1}_{-}\mathrm{N}^{1}$	4 + N	N	0	Real	3 3 3 3 3 0 4 0 5 100 100 100 100 100 100 100 100 100
DTLZ2_N	9+N	N	0	Real	3 3 3 3 3 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5
DTLZ3_N	9+N	N	0	Real	2
$\mathrm{DTLZ4}_{-}\mathrm{N}$	9+N	N	0	Real	3 - 3 - 3 - 3 - 3 - 3 - 3 - 3 - 3 - 3 -
DTLZ7_N	19 + N	N	0	Real	2 2 3 3 2 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0
Fonseca	2	2	0	Real	3 - 3 - 3 - 3 - 3 - 3 - 3 - 3 - 3 - 3 -

 $^{^{1}\}mathrm{DTLZ}$ problems are scalable to any number of objectives. Replace N with the number of objectives.

Problem	# of Vars	# of Objs	# of Constrs	Type	Pareto Front
Fonseca2	3	2	0	Real	3 3 5 5 5 6 6 8 13
Jimenez †	2	2	4	Real	87 97 98 98 98 98 98 98 98 98 98 98 98 98 98
Kita†	2	2	3	Real	\$7 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0
Kursawe	3	2	0	Real	9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9
Laumanns	2	2	0	Real	8 2 2 25 32 35 560
Lis	2	2	0	Real	3 - 3 - 3 - 3 - 3 - 3 - 3 - 3 - 3 - 3 -

Problem	# of Vars	# of Objs	# of Constrs	Type	Pareto Front
LZ1	30	2	0	Real	2
LZ2	30	2	0	Real	2
LZ3	30	2	0	Real	3 3 3 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5
LZ4	30	2	0	Real	3 3 3 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5
LZ5	30	2	0	Real	2
LZ6	10	3	0	Real	

Problem	# of Vars	# of Objs	# of Constrs	Type	Pareto Front
LZ7	10	2	0	Real	3 3 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5
LZ8	10	2	0	Real	3 3 3 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5
LZ9	30	2	0	Real	3 - 3 - 3 - 3 - 3 - 3 - 3 - 3 - 3 - 3 -
Murata	2	2	0	Real	2 2 3 3 3 2 2 2 2 2 5 5 5 6 6 6 6 6 7 6 7 6 7 6 7 6 7 6 7 6
Obayashi †	2	2	1	Real	50 mg - 10 ds ds ds ds d2
OKA1	2	2	0	Real	3 2 3 3 3 4 5 5 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6

Problem	# of Vars	# of Objs	# of Constrs	Type	Pareto Front
OKA2	3	2	0	Real	
Osyczka	2	2	2	Real	27 2 27 2 28 2 25 2 26 2 26 2 26 2 26 2 26 2 26
Osyczka2	6	2	6	Real	R 3 3 3 4 2 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5
Poloni †	2	2	0	Real	2 2 4 6 8 10 12 14
Quagliarella	16	2	0	Real	S 2 3 4
Rendon	2	2	0	Real	R

Problem	# of Vars	# of Objs	# of Constrs	Type	Pareto Front
Rendon2	2	2	0	Real	3 45 45 45 35 35 25 28
Schaffer	1	2	0	Real	3 2 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3
Schaffer2	1	2	0	Real	2
Srinivas	2	2	2	Real	0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0
Tamaki †	3	3	1	Real	
Tanaka	2	2	2	Real	2

Problem	# of Vars	# of Objs	# of Constrs	Type	Pareto Front
UF1	30	2	0	Real	
UF2	30	2	0	Real	3 - 3 - 3 - 3 - 3 - 3 - 3 - 3 - 3 - 3 -
UF3	30	2	0	Real	3 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0
UF4	30	2	0	Real	
UF5	30	2	0	Real	3 - 3 - 3 - 3 - 3 - 3 - 3 - 3 - 3 - 3 -
UF6	30	2	0	Real	3 - 3 - 3 - 3 - 3 - 3 - 3 - 3 - 3 - 3 -

Problem	# of Vars	# of Objs	# of Constrs	Type	Pareto Front
UF7	30	2	0	Real	3 - 3 - 3 - 3 - 3 - 3 - 3 - 3 - 3 - 3 -
UF8	30	3	0	Real	
UF9	30	3	0	Real	
UF10	30	3	0	Real	2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2
UF11	30	5	0	Real	
UF12 UF13	30 30	5 5	$0 \\ 0$	Real Real	
Viennet	2	3	0	Real	\$ 1 10 10 10 10 10 10 10 10 10 10 10 10 1
Viennet2	2	3	0	Real	10 10 10 10 10 10 10 10 10 10 10 10 10 1

Problem	# of Vars	# of Objs	# of Constrs	Type	Pareto Front
Viennet3	2	3	0	Real	
Viennet4	2	3	3	Real	5 6 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9
$WFG1_{-}N^{2}$	9+N	N	0	Real	S 0 0 05 15 15 20
WFG2_N	9+N	N	0	Real	3 n o o o o o o o o o o o o o o o o o o
WFG3_N	9+N	N	0	Real	S - 0 0 0 1 1 1 1 2 2 2 3 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0
$ m WFG4_N$	9+N	N	0	Real	S n 0 10 15 20 100 100 100 100 100 100 100 100 100

 $^{^{2}}$ WFG problems are scalable to any number of objectives. Replace N with the number of objectives.

Problem	# of Vars	# of Objs	# of Constrs	Type	Pareto Front
${ m WFG5_N}$	9+N	N	0	Real	S =
${ m WFG6_N}$	9+N	N	0	Real	S 0 - 0 12 15 20 000 000 000 000 000 000 000 000 000
$ m WFG7_N$	9+N	N	0	Real	S 0 0 0 10 15 20 100 100 100 100 100 100 100 100 100
${ m WFG8_N}$	9+N	N	0	Real	S a a a a a a a a a a a a a a a a a a a
${ m WFG9}_{-}{ m N}$	9+N	N	0	Real	S 4 -
ZDT1	30	2	0	Real	3 3 3 3 4 5 6 6 6 1 1 5

Problem	# of Vars	$\# ext{ of } $	# of Constrs	Type	Pareto Front
ZDT2	30	2	0	Real	3 - 3 - 3 - 3 - 3 - 3 - 3 - 3 - 3 - 3 -
ZDT3	30	2	0	Real	
ZDT4	10	2	0	Real	3 3 3 3 3 3 3 4 5 5 5 5 5 5 6 7 7 7 8 7 8 7 8 7 8 7 8 7 8 7 8 7 8 7
ZDT5	80	2	0	Binary	
ZDT6	10	2	0	Real	3 3 3 3 3 4 23 56 67 23 58 10

Appendix E

Errors and Warning Messages

This chapter provides a comprehensive list of errors and warning messages that may be encountered when using the MOEA Framework. The error or warning message is shown in *italic text*, followed by details and possible fixes for the issue.

E.1 Errors

Errors halt the execution of the program and produce an error message to the standard error stream (i.e., the console). Most errors can be corrected by the user.

Exception in thread "main" java.lang.NoClassDefFoundError: <class>

Thrown when Java is starting but is unable to find the specified class. Ensure the specified class is located on the Java classpath. If the class is located in a JAR file, use

```
java -classpath "$CLASSPATH:/path/to/library.jar" ...
```

If the class is an individual class file in a folder, use

```
java -classpath "$CLASSPATH:/path/to/folder/"
```

Also ensure you are using the correct classpath separator. Linux users will use the colon (:) as the above examples demonstrate. Windows and Cygwin users should use the semi-colon (;).

Error occurred during initialization of VM or Too small initial heap for new size specified

This Java error occurs when the initial heap size (allocated memory) is too small to instantiate the Java virtual machine (VM). This error is likely caused by the -Xmx command line option requesting less memory than is necessary to start the VM. Increasing the -Xmx value may resolve this issue. Also ensure the -Xmx argument is properly formatted. For instance, use -Xmx128m and NOT -Xmx128.

Error occurred during initialization of VM or Could not reserve enough space for object heap or Could not create the Java virtual machine

This Java error occurs when there is insufficient heap size (allocated memory) to instantiate the Java virtual machine (VM). This error is likely caused by the -Xmx command line option requesting more memory than is available on the host system. This error may also occur if other running processes consume large quantities of memory. Lowering the -Xmx value may resolve this issue.

Exception in thread "main" java.lang.OutOfMemoryError: GC overhead limit exceeded

Java relies on a garbage collector to detect and free memory which is no longer in use. This process is usually fast. However, if Java determines it is spending too much time performing garbage collection (98% of the time) and is only recovering a small amount of memory (2% of the heap), this error is thrown. This is likely caused when the inuse memory approaches the maximum heap size, leaving little unallocated memory for temporary objects. Try increasing the maximum heap size with the -Xmx command line argument.

Assertion failed: fp != NULL, file <filename>, line linenumber>

PISA modules communicate using the file system. Some anti-virus software scans the contents of files before read and after write operations. This may cause one of the PISA communication files to become inaccessible and cause this error. To test if this is the cause, try disabling your anti-virus and re-run the program.

A more permanent and secure solution involves adding an exception to the anti-virus software to prevent active monitoring of PISA communication files. For example, first add the line

```
java.io.tmpdir=<folder>
```

to plobal properties and set <folder> to some temporary folder where the PISA communication files will be stored. Then configure your anti-virus software to ignore the contents of <folder>.

problem does not have an analytical solution

Attempted to use SetGenerator to produce a reference set for a problem which does not implement AnalyticalProblem. Only AnalyticalProblems, which provide a method for generating Pareto optimal solutions, can be used with SetGenerator.

input appears to be newer than output

Several of the command line utilities read entries in an input file and write the corresponding outputs to a separate output file. If the last modified date on the input file is newer than the date on the output file, this exception is thrown. This suggests that the input file has been modified unexpectedly, and attempting to resume with a partially evaluated output file may result in incorrect results. To resolve:

- 1. If the input file is unchanged, use the --force command line option to override this check.
- 2. If the input file is changed, delete the output file and restart evaluation from the beginning.

no reference set available

Several of the command line utilities require a reference set. The reference set either is provided by the problem (through the ProblemProvider), or supplied by the user via a command line argument. This exception occurs if neither approach provides a reference set.

unable to load reference set

Indicates that a reference set is specified, but it could not be loaded. The error message should contain additional information about the underlying cause for the load failure.

output has more entries than input

Thrown by the Evaluator or ResultFileEvaluator command line utilities when attempting to resume evaluation of a partially evaluated file, but the output file contains more entries than the input file. This implies the input file was either modified, or a different input file was supplied than originally used to produce the output file. Unless the original input file is found, do not attempt to recover from this exception. Delete the output file and restart evaluation from the beginning.

maxEvaluations not defined

Thrown by the Evaluator command line utility if the maxEvaluations property has not been defined. This property must either be defined in the parameter input file or through the -x maxEvaluations=<value> command line argument.

unsupported decision variable type

Thrown when the user attempts to use an algorithm that does not support the given problem's decision variable encoding. For instance, GDE3 only supports real-valued encodings, and will throw this exception if binary or permutation encoded problems are provided.

not enough bits or not enough dimensions

The Sobol sequence generator supports up to 21000 dimensions and can produce up to 2147483647 samples $(2^{31}-1)$. While unlikely, if either of these two limits are exceeded, these exceptions are thrown.

invalid number of parents

Attempting to use CompoundVariation in a manner inconsistent with its API specification will result in this exception. Refer to the API documentation and the restrictions on the number of parents for a variation operator.

binary variables not same length or permutations not same size

Thrown by variation operators which require binary variables or permutations of equal length, but the supplied variables differ in length.

invalid bit string

Thrown by ResultFileReader if either of the following two cases occurs:

- 1. The binary variable length differs from that specified in the problem definition.
- 2. The string encoding in the file contains invalid characters.

In either case, the binary variable stored in the result file could not be read.

invalid permutation

Thrown by ResultFileReader if either of the following two cases occurs: 1) the permutation length differs from that specified in the problem definition; and 2) the string encoding in the file does not represent a valid permutation. In either case, the permutation stored in the result file could not be read.

no provider for <name>

Thrown by the service provider interface (org.moeaframework.core.spi) codes when no provider for the requested service is available. Check the following:

- 1. If a nested exception is reported, the nested exception will identify the failure.
- 2. Ensure <name> is in fact provided by a built-in or third-party provider. Check spelling and case sensitivity.
- 3. If <name> is supplied by a third-party provider, ensure the provider is located on the Java classpath. If the provider is in a JAR file, use

```
java -classpath "$CLASSPATH:/path/to/provider.jar"
...
```

If the provider is supplied as class files in a folder, use

```
java -classpath "$CLASSPATH:/path/to/folder/"
```

Also ensure you are using the correct classpath separator. Linux users will use the colon (:) as the above examples demonstrate. Windows and Cygwin users should use the semi-colon (;).

error sending variables to external process or error receiving variables from external process

Thrown when communicating with an external problem, but an I/O error occurred that disrupted the communication. Numerous situations may cause this exception, such as the external process terminating unexpectedly.

end of stream reached when response expected

Thrown when communicating with an external process, but the connection to the external process closed. This is most likely the result of an error on the external process side which caused the external process to terminate unexpectedly. Error messages printed to the standard error stream should appear in the Java error stream.

response contained fewer tokens than expected

Thrown when communicating with an external problem, and the external process has returned an unexpected number of entries. This is most likely a configuration error where the defined number of objectives or constraints differs from what is actually returned by the external process.

unable to serialize variable

Attempted to serialize a decision variable to send to an external problem, but the decision variable is not one of the supported types. Only real variables are supported.

restart not supported

PISA supports the ability to reuse a selector after a run has completed. The MOEA Framework currently does not support this feature. This exception is thrown if the PISA selector attempts to reset.

expected END on last line or unexpected end of file or invalid selection length

These exceptions are thrown when communicating with PISA processes, and the files produced by the PISA process appear to be incomplete or malformed. Check the implementation of the PISA codes to ensure they follow the correct protocol and syntax.

invalid variation length

This exception is caused by an incorrect configuration of PISA. The following equality must hold

children*(mu/parents) = lambda,

where mu is the number of parents selected by the PISA process, parents is the number of parent solutions required by the variation operator, children is the number of offspring produced by a single invocation of the variation operator, and lambda is the total number of offspring produced during a generation.

no digest file

Thrown when attempting to validate a data file using a digest file, but no such digest file exists. Processing of the data file should cease immediately for sensitive applications where data integrity is essential. If the digest file simply hasn't yet been produced but the file contents are verified, the FileProtection command line utility can optionally generate digest files.

invalid digest file

Thrown when attempting to validate a date file using a digest file, but the digest file is corrupted or does not contain a valid digest. Processing of the data file should cease immediately for sensitive applications where data integrity is essential.

digest does not match

Thrown when attempting to validate a data file using a digest file, but the actual digest of the data file does not match the expected digest contained in the digest file. This indicates that the data file or the digest file are corrupted. Processing of the data file should cease immediately for sensitive applications where data integrity is essential.

unexpected rule separator or rule must contain at least one production or invalid symbol or rule must start with non-terminal or rule must contain at least one production or codon array is empty

Each of these exceptions originates in the grammatical evolution code, and indicate specific errors when loading or processing a context free grammar. The specific error message details the cause.

unable to mkdir idirectory;

For an unknown reason, the underlying operating system was unable to create a directory. Check to ensure the location of the directory is writable. One may also manually create the directory.

no scripting engine for extension jext;

Attempted to use the Java Scripting APIs, but no engine for the specified file extension could be found. To resolve:

- 1. Check that the extension is valid. If not, supply the file extension for the scripting language required.
- 2. Ensure the scripting language engine is listed on the classpath. The engine, if packaged in a JAR, can be specified with

```
java -classpath "$CLASSPATH:/path/to/engine.jar"
```

Also ensure you are using the correct classpath separator. Linux users will use the colon (:) as the above example demonstrates. Windows and Cygwin users should use the semi-colon (;).

no scripting engine for jname;

Attempted to use the Java Scripting APIs, but no engine with the specified name was found.

- 1. Check that the name is valid. If not, supply the correct name for the scripting language engine as required.
- 2. Ensure the scripting language engine is listed on the classpath. The engine, if packaged in a JAR, can be specified with

```
java -classpath "$CLASSPATH:/path/to/engine.jar"
```

Also ensure you are using the correct classpath separator. Linux users will use the colon (:) as the above example demonstrates. Windows and Cygwin users should use the semi-colon (;).

file has no extension

Attempted to use a script file with ScriptedProblem, but the filename does not contain a valid extension. Either supply the file extension for the scripting language required, or use the constructor which accepts the engine name as an argument.

scripting engine not invocable

Thrown when using a scripting language engine which does not implement the Invocable interface. The scripting language does not support methods or functions, and thus can not be used as intended.

requires two or more groups

Attempted to use one of the n-ary statistical tests which require at least two groups. Either add a second group to compare against, or remove the statistical test.

could not locate resource inameż

Thrown when attempting to access a resource packages within the MOEA Framework, but the resource could not be located. This is an error with the distribution. Please contact the distributor to correct this issue.

insufficient number of entries in row

Attempted to read a data file, but the row was missing one or more entries. The exact meaning depends on the specific data file, but generally this error means the file is incomplete, improperly formatted or corrupted. See the documentation on the various file types to determine if this error can be corrected.

invalid entry in row

Attempted to read a data file, but an entry was not formatted correctly. See the documentation on the various file types to determine if this error can be corrected.

invoke calculate prior to getting indicator values

Attempted to retrieve one of the indicator values prior to invoking the calculate method. When using QualityIndicator, the calculate method must be invoked prior to retrieving any of the indicator values.

```
not a real variable or
not a binary variable or
not a boolean variable or
not a permutation
```

The EncodingUtils class handles all the type checking internally. If any of the arguments are not of the expected type, one of these exceptions is thrown. Ensure the argument is of the expected type. For example, ensure variable is a BinaryVariable when calling EncodingUtils.asBinary(variable).

```
invalid number of values or invalid number of bits
```

Attempted to set the decision variable values using an array, but the number of elements in the array does not match the required number of elements. For EncodingUtils. setReal and EncodingUtils.setInt, ensure the number of real-valued/integer-valued decision variables being set matches the array length. For EncodingUtils. setBinary, ensure the number of bits expressed in the binary variable matches the array length.

lambda function is not valid

In genetic programming, a lambda function was created with an invalid body. The body of a lambda function must be fully defined and strongly typed. If not, this exception is thrown. Check the definition of the lambda function and ensure all arguments are non-null and are of the correct type. Check the error output to see if any warning messages were printed that detail the cause of this exception.

index does not reference node in tree

Attempted to use one of the node.getXXXAt() methods, but the index referred to a node not within the tree. This is similar to an out-of-bounds exception, as the index pointed to a node outside the tree. Ensure the index is valid.

malformed property argument

The Evaluator and Solve command line utilities support setting algorithm parameters on the command line with the -x option. The parameters should be of the form:

-x name=value

or if multiple parameters are set:

-x name1=value1; name2=value2; name3=value3

This error is thrown if the command line argument is not in either of these two forms. Check the command line argument to ensure it is formatted correctly.

key not defined in accumulator: <key>

Thrown when attempting to access a key in an Accumulator object that is not contained within the Accumulator. Use accumulator.keySet() to see what keys are available and ensure the requested key exists within the accumulator.

an unclean version of the file exists from a previous run, requires manual intervention

Thrown when ResultFileWriter or MetricFileWriter attempt to recover data from an interrupted run, but it appears there already exists an "unclean" file from a previous recovery attempt. If the user believes the unclean file contains valid data, she can copy the unclean file to its original location. Or, she can delete the unclean file to start fresh. The org.moeaframework.analysis.sensitivity.cleanup property in plobal.properties controls the default behavior in this scenario.

requires at least two solutions or objective with empty range

These two exceptions are thrown when using the Normalizer with a degenerate population. A degenerate population either has fewer than two solutions or the range of any objective is below computer precision. In this scenario, the population can not be normalized.

lower bound and upper bounds not the same length

When specifying the --lowerBounds and --upperBounds arguments to the Solve utility, the number of values in the comma-separated list must match.

invalid variable specification įvalue $\dot{\xi}$, not properly formatted invalid real specification įvalue $\dot{\xi}$, expected $R(jlb\dot{\xi},jub\dot{\xi})$ invalid binary specification įvalue $\dot{\xi}$, expected $B(jlength\dot{\xi})$ invalid permutation specification įvalue $\dot{\xi}$, expected $P(jlength\dot{\xi})$ invalid variable specification įvalue $\dot{\xi}$, unknown type

The --variables argument to the Solve utility allows specifying the types and ranges of the decision variables. These error messages indicate that one or more of the variable specifications is invalid. The message will identify the problem. An example variable specification is provided below:

```
--variables "R(0;1),B(5),P(10),R(-1;1)"
```

Also, always surround the argument with quotes as shown in this example.

must specify either the problem, the variables, or the lower and upper bounds arguments

The Solve command line utility operates on both problems defined within the MOEA Framework (by name) or problems external to the MOEA Framework, such as an executable. For problems identified by name, the --problem argument must be specified. For external problems, (1) if the problem is real-valued, you can use the --lowerBounds and --upperBounds arguments; or (2) use the --variables argument to specify the decision variables and their types.

E.2 Warnings

Warnings are messages printed to the standard error stream (i.e., the console) that indicate an abnormal or unsafe condition. While warnings do not indicate an error occurred, they do indicate caution is required by the user.

no digest file exists to validate <FILE>

Attempted to validate the file but no digest file exists. This indicates that the framework could not verify the authenticity of the file.

saving result file without variables, may become unstable

Occurs when writing a result file with the output of decision variables suppressed. The suppression of decision variable output is a user-specified option. The warning "may become unstable" indicates that further use of the result file may result in unexpected errors if the decision variables are required.

unsupported decision variable type, may become unstable

Occurs when reading or writing result files which use unsupported decision variable types. When this occurs, the program is unable to read or write the decision variable, and its value is therefore lost. The warning "may become unstable" indicates that further use of the result file may result in unexpected errors if the decision variables are required.

duplicate solution found

Issued by ReferenceSetMerger if any of the algorithms contribute identical solutions. If this warning is emitted, the contribution of each algorithm to the reference set is invalid. Use SetContribution instead to compute the contribution of overlapping sets to a reference set.

can not initialize unknown type

Emitted by RandomInitialization if the problem uses unsupported decision variable types. The algorithm will continue to run, but the unsupported decision variables will remain initialized to their default values.

an error occurred while writing the state file or an error occurred while reading the state file

Occurs when checkpoints are enabled, but the algorithm does not support checkpoints or an error occurred while reading or writing the checkpoint. The execution of the algorithm will continue normally, but no checkpoints will be produced.

multiple constraints not supported, aggregating into first constraint

Occurs when an algorithm implementation does not support multiple constraints. This occurs primarily with the JMetal library, which only uses a single aggregate constraint violation value. When translating between JMetal and the MOEA Framework, the first objective in the MOEA Framework is assigned the aggregate constraint violation value; the remaining objectives become 0.

increasing MOEA/D population size

The population size of MOEA/D must be at least the number of objectives of the problem. If not, the population size is automatically increased.

checkpoints not supported when running multiple seeds

Emitted by the Executor when the withCheckpointFile(...) and accumulateAcrossSeeds(...) options are both used. Checkpoints are only supported for single-seed evaluation. The Executor will continue without checkpoints.

checkpoints not supported by algorithm

Emitted by the Executor if the algorithm is not Resumable (i.e., does not support checkpoints). The Executor will continue without checkpoints.

 $Provider\ org. moeaframework. algorithm. jmetal. JMetal Algorithms\ could\ not\ be\ instantiated: java.lang. No Class Def Found Error: <class>$

This warning occurs when attempting to instantiate the JMetal algorithm provider, but the JMetal library could not be found on the classpath. This is treated as a warning and not an exception since a secondary provider may exist for the specified algorithm. If no secondary provider exists, a ProviderNotFoundException will be thrown. To correct, obtain the latest JMetal library from http://jmetal.sourceforge.net/ and list it on the classpath as follows:

java -classpath "\$CLASSPATH:/path/to/JMetal.jar"

Also ensure you are using the correct classpath separator. Linux users will use the colon (:) as the above example demonstrates. Windows and Cygwin users should use the semi-colon (;).

unable to negate values in <file>, incorrect number of values in a row

Emitted by the Negater command line utility when one of the files it is processing contains an invalid number of values in a row. The file is expected to contain the same number of values in a row as values passed to the -d, --direction command line argument. The file will not be modified if this issue is detected.

unable to negate values in <file>, unable to parse number

Emitted by the Negater command line utility when one of the files it is processing encounters a value it is unable to parse. The columns being negated must be numeric values. The file will not be modified if this issue is detected.

```
argument is null or
<class> not assignable from <class>
```

When validating an expression tree using the node.isValid() method, details identifying why the tree is invalid are printed. The warning "argument is null" indicates the tree is incomplete and contains a missing argument. Check to ensure all arguments of all nodes within the tree are non-null. The warning "<class> not assignable from <class>" indicates the required type of an argument did not match the return type of the argument. If this warning appears when using Sequence, For or While nodes, ensure you specify the return type of these nodes using the appropriate constructor.

unable to parse solution, ignoring remaining entries in the file or insufficient number of entries in row, ignoring remaining rows in the file

Occurs when MetricFileReader or ResultFileReader encounter invalid data in an input file. They automatically discard any remaining entries in the file, assuming they are corrupt. This is primarily intended to allow the software to automatically recover from a previous, interrupted execution. These warnings are provided to inform the user that invalid entries are being discarded.

Unable to find the file ifile?

This warning is shown when running an example that must load a data file but the data file could not be found. Ensure that the examples directory is located on your classpath:

```
java -classpath "$CLASSPATH:examples" ...
```

Also ensure you are using the correct classpath separator. Linux users will use the colon (:) as the above example demonstrates. Windows and Cygwin users should use the semi-colon (;).

incorrect number of names, using defaults

Occurs when using the --names argument provided by ARFFConverter and AerovisConverter to provide custom names for the decision variables and/or objectives, but the number of names provided is not correct. When providing names for only the objectives, the number of names must match the number of objectives. When providing names for both variables and objectives, the number of names must match the number of variables and objectives in the data file. Otherwise, this warning is displayed and the program uses default names.

population is empty, can not generate ARFF file

The ARFFConverter outputs an ARFF file using the last entry in a result file. If the last entry is empty, then no ARFF file is generated.

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