# libea 0.2 - A brief introduction

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## 1 Introduction

LIBEA is a template based library written in C++14. The purpose of this software is to provide an extensible and reliable framework for writing evolutionary algorithms.

## 2 Building libea

To build LIBEA scons<sup>1</sup> is required. The library can be compiled and installed with the following commands:

After the installation you may want to build the test suite. Please note that this step requires the CppUnit<sup>2</sup> framework:

```
\# scons test-suite
```

If you want to generate the source code documentation please ensure Doxygen<sup>3</sup> is installed on your system and type in the following command:

```
\# scons doc
```

This documentation can be build with

```
# scons pdf
```

If you want to uninstall libea type in

```
\# sudo scons install -c
```

#### 3 The framework

## 3.1 A quick overview

In LIBEA individuals are represented in *sequences*. Theoretically any data type large enough to store the genotype of an individual is a valid sequence type. A group of sequences is called *population*.

To modify a sequence or evaluate its fitness a corresponding  $genome\ base\ class$  inherited from ea::AGenomeBase is required. Figure 1 illustrates this concept.

LIBEA offers the genome base class templates ea::PGenomeBase and ea::CPGenomeBase with the corresponding sequence types ea::Sequence and ea::CSequence. Any data type with a valid copy constructor is a supported gene type. There are already type aliases for many primitive data types available in LIBEA (see table 1).

The two offered base class templates are optimized for different use-cases. As shown in listing 1 the *ea::Sequence* structure has only two fields which are used to store the length of a sequence and its genes. Hence this kind of sequence is preferable if the application's memory footprint is critical.

<sup>1</sup>http://www.scons.org/

<sup>2</sup>http://freedesktop.org/wiki/Software/cppunit/

<sup>3</sup>http://www.stack.nl/~dimitri/doxygen/

Figure 1: sequences and corresponding genome base class

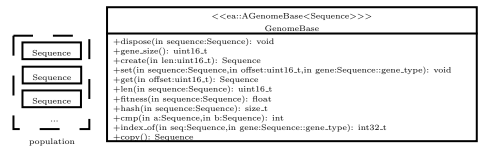


Table 1: genome base classes and corresponding sequence types offered by LIBEA

Table 1. Scholie base classes and corresponding sequence types offered by Eibbit		
GENOME BASE CLASS	CORRESPONDING SEQUENCE TYPE	GENE TYPE
ea::PGenomeBase <t></t>	ea::Sequence <t></t>	T
ea::Int32PenomeBase	ea::Sequence $<$ int $32_{-}t>$	$int32_t$
ea::UInt32PGenomeBase	ea::Sequence $<$ uint $32_{-}$ t $>$	$uint32_t$
ea::BinaryPGenomeBase	ea::Sequence <bool></bool>	bool
ea::DoublePGenomeBase	ea::Sequence <double></double>	double
ea::StringPGenomeBase	ea::Sequence <std::string></std::string>	std::string
ea::CPGenomeBase <t></t>	ea::CSequence <t></t>	Τ
ea::Int32CPenomeBase	ea::CSequence $<$ int $32_t>$	$int32_t$
ea::UInt32CPGenomeBase	ea::CSequence $<$ uint $32_t>$	$uint32_t$
ea::BinaryCPGenomeBase	ea::CSequence <bool></bool>	bool
ea::DoubleCPGenomeBase	ea::CSequence <double></double>	double
ea::StringCPGenomeBase	ea::CSequence <std::string></std::string>	std::string

In comparison the ea::CSequence structure has three additional fields. They are used to cache the evaluated fitness value and the hash of an individual (see listing 2). In general this sequence type is a better choice because the performance benefit might out weight the drawback of memory consumption.

The fields of both structures should **never be accessed directly**. LIBEA provides the functions  $ea::sequence\_len()$  and  $ea::sequence\_get()$  for reading genes from a sequence.

Listing 1: Sequence template declaration

```
template<typename TGene>
2
   struct Sequence
3
   {
4
     /*! Datatype of stored genes. */
5
     typedef TGene gene_type;
6
     /*! Number of stored genes. */
7
8
     uint16_t len;
      /*! Dynamic array holding genes. */
10
     TGene* genes;
11
   };
```

Listing 2: CSequence template declaration

```
{\bf template}{<}{\bf typename} \ \ {\rm TGene}{>}
1
2
    struct CSequence : Sequence<TGene>
3
      /*! Flags. */
4
5
      uint8_t flags;
6
      /*! Cached fitness value. */
7
      double fitness;
      /*! Cached hash value. */
8
9
      size_t hash;
10 };
```

#### 3.2 Core features

## 3.3 Working with sequences

Listing 3 demonstrates some basic functions of LIBEA. An individual represented in a sequence of integers is created (ll. 44-50) and copied (l. 62). The sequences are compared before and after modifying the second individual (ll. 67-75).

To access the genes and evaluate the fitness a genome base class is declared (l. 22). The template parameters specify the gene type and fitness function (ll. 5-19).

Listing 3: core functions

```
#include <iostream>
   #include <libea.hpp>
    // fitness function:
4
    class Fitness
5
6
    {
7
      public:
8
        double operator()(const ea::Sequence<int32_t>* const &seq)
             const
9
10
          double avg = 0;
11
12
           for(ea::sequence\_len\_t i = 0; i < ea::sequence\_len(seq);
13
             avg \; +\!\!=\; ea::sequence\_get \, (\, seq \; , \quad i \; ) \; ;
14
15
16
17
          return avg / ea::sequence_len(seq);
        }
18
19
    };
20
    // base class for modifying/accessing sequences:
21
    static ea::PGenomeBase<int32_t , Fitness> base;
22
23
24
    // print a sequence:
25 static void print_genome(ea::Sequence<int32_t>* seq)
26 {
```

```
27
      // print length, hash & fitness:
      \mathrm{std}::\mathrm{cout} \;<<\;"\,\mathrm{len}: \_" \;<<\;\mathrm{base}\,.\,\mathrm{len}\,(\,\mathrm{seq}\,) \;<<\;"\,, \_\mathrm{hash}: \_" \;<<\;\mathrm{base}\,.
28
          hash(seq) << ", \_fitness: \_" << base.fitness(seq) << std::
          endl;
29
30
      // print sequence:
31
      std::cout << "sequence:";
32
33
      for (ea::sequence_len_t i = 0; i < 10; ++i)
34
35
        std::cout << base.get(seq, i) << "";
36
37
38
      std::cout << std::endl;
39
40
    int main(int argc, char *argv[])
41
42
43
      // create new sequence:
44
      ea::Sequence<int32_t>* a = base.create(10);
45
      // set genes:
46
47
      for (ea::sequence\_len\_t i = 0; i < 10; ++i)
48
        base.set(a, i, i + 100);
49
50
51
      std::cout << "genome_a:" << std::endl;
52
53
      print_genome(a);
54
55
      // show index of genes:
56
      for (uint32_t g = 100; g \le 110; ++g)
57
58
        std::cout << "index\_of\_" << g << ":\_" << base.index\_of(a\,,
            g) << std::endl;
59
      }
60
      // copy sequence:
61
      ea::Sequence<int32_t>*b = base.copy(a);
62
      std::cout << "genome_b:" << std::endl;
63
64
      print_genome(b);
65
66
      // compare sequences:
      std::cout << "compare:" << base.cmp(a, b) << std::endl;
67
68
      // change single gene:
69
70
      base.set(b, 5, 42);
      std::cout << "genome_b:" << std::endl;</pre>
71
72
      print_genome(b);
73
74
      // compare sequences:
      std::cout << "compare:" << base.cmp(a, b) << std::endl;
75
76
77
      // free memory:
```

```
78 ea::dispose(base, { a, b });
79
80 return 0;
81 }
```

#### 3.4 Factories & output adapters

LIBEA offers the *ea::AFactory* base class as an option to create populations. Listing 4 shows how to create sequences of ten random integers using a custom factory (ll. 30-51). The values are generated with a random number generator (ll. 35-46).

The individuals are written to an ea::OutputAdapter (ll. 74-77). At the current stage LIBEA only provides an adapter for STL containers like std::vector, but new adapters can be added easily by implementing the ea::IOutputAdapter interface. To create an adapter use the ea::make\_output\_adapter() function.

Listing 4: custom factory

```
1 #include <stdint.h>
   #include <iostream>
3 #include <vector>
4
   #include <libea.hpp>
5
   // store genomes in sequences of int32\_t, this typedef makes
6
       the\ code\ more\ readable:
7
   typedef ea::Sequence<int32_t> Sequence;
9
   // fitness function:
10
   class Fitness
11
12
13
        double operator()(const Sequence* const &seq) const
14
          double avg = 0;
15
16
17
          for (ea::sequence\_len\_t i = 0; i < ea::sequence\_len(seq);
               ++i)
18
19
            avg += ea::sequence_get(seq, i);
20
21
22
          return avg / ea::sequence_len(seq);
        }
23
24
    };
25
26
    // base class for modifying/accessing sequences:
27
   static ea::PGenomeBase<int32_t , Fitness> base;
28
29
   // factory class:
30
   class Factory : public ea::AFactory<Sequence*>
31
32
      public:
33
        Sequence * create_sequence()
```

```
34
            int32_t genes[10];
35
36
37
            g.get_int32\_seq(1, 100, genes, 10);
38
            Sequence* seq = base.create(10);
39
40
            for(ea::sequence\_len\_t i = 0; i < 10; ++i)
41
42
43
              base.set(seq, i, genes[i]);
44
45
46
            return seq;
47
         }
48
49
       private:
50
         static ea::AnsiRandomNumberGenerator _g;
51
    };
52
53
    ea::AnsiRandomNumberGenerator Factory::_g;
54
55
    // debug function:
56
    static void print_genome(Sequence* seq)
57
       std::cout << "hash:" << base.hash(seq) << ", _fitness:" <<
58
           base.fitness(seq) << ", _sequence: _";
59
       for (ea::sequence_len_t i = 0; i < 10; ++i)
60
61
         \mathrm{std} :: \mathrm{cout} \; << \; \mathrm{base.get} \left( \; \mathrm{seq} \; , \; \; \mathrm{i} \; \right) \; << \; " \; \lrcorner " \; ;
62
63
64
       \mathtt{std} :: \mathtt{cout} \, <\!< \, \mathtt{std} :: \mathtt{endl} \, ;
65
66
67
68
    int main(int argc, char *argv[])
69
70
       Factory f;
71
       std::vector<Sequence*> population;
72
73
       // create adapter:
       {\bf auto}\ {\tt adapter}\ =\ {\tt ea::make\_output\_adapter}\ (\ {\tt population}\ )\ ;
74
75
76
       // create 100 genomes:
77
       f.create-population(100, adapter);
78
79
       // print genomes & free memory:
80
       std::for_each(begin(population), end(population), [](
           Sequence* seq)
81
82
         print_genome(seq);
83
       });
84
85
       ea::dispose(base, begin(population), end(population));
```

```
86
87 return 0;
88 }
```

## 3.5 Selection operators

Selection operators have to implement the ea::IIndexSelection interface (see listing 5). Instead of copying the selected genomes to an ea::OutputAdapter only the indexes are written. Access to the parent individuals is given through an adapter implementing the ea::IInputAdapter interface. Input adapters can be created for all STL containers providing random access iterators with the ea::make\_input\_adapter() function.

Listing 5: IIndexSelection

```
1
   template<typename TGenomeBase>
2
   class IIndexSelection
3
4
5
       typedef typename TGenomeBase::sequence_type sequence_type;
6
7
       ~IIndexSelection() {}
8
9
       virtual void select(IInputAdapter<sequence_type>& input,
           const uint32_t count, IOutputAdapter<uint32_t>& output
           ) = 0;
10
   };
```

Listing 6 creates a population with ten random sequences (ll. 43-58). Then five individuals are selected using the tournament selection operator (ll. 72-82).

Listing 6: selection operators

```
1 #include <stdint.h>
   #include <iostream>
3
   #include <vector>
4 #include <libea.hpp>
   // store genomes in sequences of int32_t, this typedef makes
6
       the code more readable:
7
   typedef ea::Sequence<int32_t> Sequence;
8
   // fitness function:
9
   class Fitness
10
11
12
     public:
        double operator()(const Sequence* const &seq) const
13
14
          double avg = 0;
15
16
17
          for (ea::sequence\_len\_t i = 0; i < ea::sequence\_len(seq);
18
19
            avg += ea::sequence_get(seq, i);
20
```

```
21
22
          return avg / ea::sequence_len(seq);
23
        }
24
   };
25
   static ea::Int32PGenomeBase<Fitness> base;
26
27
   static void print_genome(Sequence* seq)
28
29
30
      for(ea::sequence\_len\_t i = 0; i < ea::sequence\_len(seq); ++i
31
32
        std::cout << base.get(seq, i) << "";
33
34
35
     std::cout << std::endl;
36
37
   int main(int argc, char *argv[])
38
39
40
     ea::AnsiRandomNumberGenerator g;
41
42
     // create parent individuals:
43
     std::vector<Sequence*> population;
44
     for (ea::sequence_len_t i = 0; i < 10; ++i)
45
46
47
        auto seq = base.create(10);
48
49
        int32_t genes[10];
50
        g.get\_unique\_int32\_seq(0, 9, genes, 10);
51
52
        for(ea::sequence\_len\_t j = 0; j < 10; ++j)
53
54
          base.set(seq, j, genes[j]);
55
56
57
        population.push_back(seq);
58
59
60
     // create input adapter:
61
     auto input = ea::make_input_adapter(population);
62
63
     // print parent individuals:
64
     std::for_each(begin(population), end(population), [](
         Sequence* seq)
65
66
        print_genome(seq);
67
     });
68
69
     std::cout << std::endl;
70
71
     // select & print individuals:
72
      std::vector<std::size_t> children;
```

```
73
     auto output = ea::make_output_adapter(children);
74
75
     ea::TournamentSelection<ea::Int32PGenomeBase<Fitness>> sel;
     sel.select(input, 5, output);
76
77
     std::for_each(begin(children), end(children), [&population](
78
         uint32_t index)
79
       std::cout << "selected_child:_" << index << "=>_";
80
81
       print_genome(population[index]);
82
     });
83
     // cleanup:
84
85
     ea::dispose(base, begin(population), end(population));
86
87
     return 0;
   }
88
```

#### 3.6 Crossover operators

Crossover operators are inherited from ea::ACrossover and have to override the crossover() method (see listing 7).

Listing 7: ACrossover

```
template<typename TGenomeBase>
2
   class ACrossover
3
   {
     public:
4
5
       typedef typename TGenomeBase::sequence_type sequence_type;
6
7
       ~ACrossover() {}
8
       virtual uint32_t crossover(const sequence_type& a, const
9
           sequence_type& b, IOutputAdapter<sequence_type>&
           output) = 0;
10 };
```

Crossover operators create at least one child sequence from two parent individuals. The *crossover()* method returns the number of child sequences written to the given output adapter. Listing 8 shows an example.

In the example two parent individuals are created (ll. 41-42). Then two children are generated using a cycle crossover operator (ll. 51-55).

Listing 8: crossover operators

```
1 #include <stdint.h>
2 #include <iostream>
3 #include <vector>
4 #include <libea.hpp>
5
6 // store genomes in sequences of int32_t, this typedef makes the code more readable:
7 typedef ea::Sequence<int32_t> Sequence;
```

```
// fitness function:
9
10
  class Fitness
11
12
        double operator()(const Sequence* const &seq) const
13
14
          double avg = 0;
15
16
17
          for(ea::sequence\_len\_t i = 0; i < ea::sequence\_len(seq);
18
19
            avg += ea::sequence_get(seq, i);
20
21
22
          return avg / ea::sequence_len(seq);
23
24
    };
25
26
    static ea::Int32PGenomeBase<Fitness> base;
27
28
   static void print_genome(Sequence* seq)
29
30
      for(ea::sequence\_len\_t i = 0; i < ea::sequence\_len(seq); ++i
31
        std::cout << base.get(seq, i) << "_";
32
33
34
35
      std::cout << std::endl;
36
   }
37
38
   int main(int argc, char *argv[])
39
      // create parent individuals:
40
41
      auto a = base.create(10);
42
      auto b = base.create(10);
43
      for (ea::sequence_len_t i = 0; i < 10; ++i)
44
45
46
        base.set(a, i, i);
        base.set\left(b\,,\ i\;,\ 9\;-\;i\,\right);
47
48
49
      // create children:
50
51
      std::vector<Sequence*> children;
52
      auto adapter = ea::make_output_adapter(children);
53
      ea::CycleCrossover<ea::Int32PGenomeBase<Fitness>> crossover;
54
55
      std::size_t n = crossover.crossover(a, b, adapter);
56
      std::cout << "number\_of\_created\_children: \_" << n << std::
57
          endl;
58
```

```
std::for_each(begin(children), end(children), [](Sequence*
59
         seq)
60
        print_genome(seq);
61
62
      });
63
64
     // cleanup:
     ea::dispose(base, { a, b });
65
66
     ea::dispose(base, begin(children), end(children));
67
68
     return 0;
69
   }
```

## 3.7 Mutation operators

Mutation operators are inherited from ea::AMutation and have to override the create\_child() method (see listing 9).

Listing 9: AMutation

```
template<class TGenomeBase>
2
   class AMutation
3
   {
4
      public:
5
        typedef typename TGenomeBase::sequence_type sequence_type;
6
7
        virtual ~AMutation() {};
8
9
        virtual void mutate(sequence_type& sequence) = 0;
10
11
        sequence_type create_child(const sequence_type& sequence)
12
        {
          static TGenomeBase base;
13
14
          sequence_type m = base.copy(sequence);
15
16
          mutate(m);
17
18
          return m;
        }
19
20 };
```

In listing 10 a parent individual is created (ll. 43-48) and mutated in-place (l. 55). Afterwards an inverted child sequence is generated (ll. 59-61).

Listing 10: mutation operators

```
9
      public:
10
         double operator()(const Sequence* const &seq) const
11
12
13
           double count = 0;
14
           for (ea::sequence\_len\_t i = 0; i < ea::sequence\_len(seq);
15
16
              if(ea::sequence\_get(seq, i))
17
18
19
                ++count;
20
21
           }
22
23
           return count;
24
25
    };
26
27
    {\bf typedef} \ {\it ea}:: Binary PG enome Base {<} Fitness {>} \ Base;
28
    static Base base;
29
    static void print_genome(Sequence* seq)
30
31
32
      for(ea::sequence\_len\_t i = 0; i < ea::sequence\_len(seq); ++i
33
         std::cout << (base.get(seq, i) ? 1 : 0);
34
35
36
37
      \operatorname{std} :: \operatorname{cout} \,<< \,\operatorname{std} :: \operatorname{endl}\,;
38
    }
39
40
    int main(int argc, char *argv[])
41
      // create parent individual:
42
43
      auto a = base.create(10);
44
      for (ea::sequence\_len\_t i = 0; i < 10; ++i)
45
46
         base.set(a, i, i % 3);
47
48
49
50
      print_genome(a);
51
52
      // inplace mutation:
53
      ea::BitStringMutation < Base > muta;
54
55
      muta. mutate(a);
56
      print_genome(a);
57
      // create child:
58
      ea::InverseBitStringMutation <Base> mutb;
59
60
```

## 3.8 Processing operator pipelines

The *ea::Pipeline* namespace provides a framework to connect and process multiple genetic operators.

LIBEA processes operators by connecting elements to a pipeline. Pipeline elements read sequences from a source and write the processed data to a sink. Each element has to implement the *ea::Pipeline::IElelement* interface. LIBEA offers a corresponding pipeline element for all operator types.

A functor implementing the *ea::Pipeline::ITerminator* is executed after the last pipeline element has been processed to decide if the process should be repeated (see figure 2).

#### SelectionElement

The ea::Pipeline::SelectionElement pipeline element reads sequences from a source population and writes selected elements to the sink. Elements are selected by an underlying ea::IIndexSelection operator.

Creating a ea::Pipeline::SelectionElemenet instance two template parameters have to be defined: the desired selection operator and a functor inherited from ea::Pipeline::ASelectionSize. The functor is required to determine the number of sequences to select. LIBEA already offers two functors:

- ea::libea::SourceDivisor: divides the size of the source population by template parameter N
- $\bullet$  ea::libea::FixedSelectionSize: returns the template parameter N

#### CrossoverElement

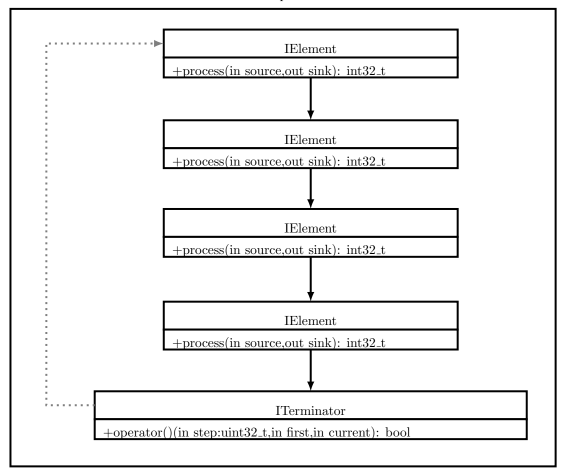
The ea::Pipeline::CrossoverElement pipeline element combines all sequences from a source population and produces children using the underlying ea::ACrossover operator. Generated children are written to the sink.

#### MutationElement

The ea::Pipeline::MutationElement reads sequences from a source population and writes them to the sink. When writing the sequences are mutated with a definable probability P using the underlying ea::AMutation operator.

Figure 2: connecting pipeline elements

## Pipeline



#### **ITerminator**

Terminators have to implement the *ea::Pipeline::ITerminator* interface (see listing 11). The () operator returns *true* if the process should be terminated and has the following parameters:

- ullet step: a counter incremented at the beginning of each iteration
- first: an input adapter providing access to the initial population
- current: an input adapter providing access to the current population

LIBEA offers the ea::Pipeline::ForLoopInidicator to repeat the pipeline process using step as loop counter.

#### Listing 11: ITerminator

```
1
        template<typename TGenomeBase>
2
        class ITerminator
3
4
          public:
            virtual ~ITerminator() {}
5
6
7
            virtual bool operator()(const uint32_t step,
                        IInputAdapter<typename TGenomeBase::
8
                            sequence_type>& first,
9
                        IInputAdapter<typename TGenomeBase::
                            sequence_type>& current) = 0;
10
        };
```

#### Travelling saleman problem

Listing 12 connects four genetic operators to find solutions for the popular travelling salesman problem.

A city is identified by an *uint8*<sub>-t</sub> (l. 11). A route is a sequence of cities (l. 12). Each city has a location which can be accessed by a *std::map* (ll. 14-31). The fitness function returns the sum of the distances between the cities of a route (ll. 34-51).

The initial population consists of 100 random routes created with a factory class (ll. 57-82). The mean and median fitness values are displayed (ll. 94-102) before processing the pipeline.

The operators are connected by their corresponding pipeline elements. At the beginning of each iteration sequences are selected by tournament selection. Then the selected sequences are used to create new children using edge recombination. After selecting 50 child sequences by double tournament selection the result is mutated using single swap mutation. The process is repeated 100 times (ll. 107-126).

After processing the pipeline mean and median fitness values of the generated child population are displayed (ll. 128-131).

Listing 12: pipeline operators

```
1 #include <libea.hpp>
   #include <vector>
3
   #include <map>
   #include <iostream>
6
   using namespace std;
7
   using namespace ea;
8
   using namespace ea::Pipeline;
9
10
   // type declarations:
   typedef uint8_t City;
11
12
   typedef CSequence<City>* Route;
13
14
   typedef struct
15
16
     uint16_t x;
```

```
17
      uint16_t y;
18
   } Point;
19
20
   // city map:
21
    static map<City , Point> Cities = {
22
      \{0, \{31, 9\}\},
23
      \{1, \{17, 82\}\},\
24
      \{2, \{56, 47\}\},
25
      \{3, \{70, 3\}\},
26
                  7 } },
       4, \{ 11, 
27
       5, { 93, 83 } },
      \{6, \{96, 22\}
28
29
        7, \{ 26, 97 \}
                     } },
30
       8, \{ 52, 58 \} \},
      { 9, { 68, 6 } };
31
32
33
    // fitness:
    class Fitness
34
35
36
      public:
37
        double operator()(const Route &seq) const
38
          double f = 0;
39
40
          for (ea::sequence\_len\_t i = 0; i < sequence\_len(seq) - 1;
41
42
43
            auto a = sequence_get(seq, i);
44
            auto b = sequence_get(seq, i + 1);
45
            f \leftarrow sqrt(pow(Cities[b].x - Cities[a].x, 2) + pow(
46
                 Cities [b].y - Cities [a].y, 2));
47
48
49
          return f;
        }
50
51
    };
52
    // base class type declaration:
53
    typedef CPGenomeBase<City , Fitness > Base;
54
55
56
    // factory:
   class RouteFactory : public ea::AFactory<Route>
57
58
      public:
59
        Route create_sequence()
60
61
62
          int32_t genes[10];
63
64
          _{g.get\_unique\_int32\_seq(0, 9, genes, 10)};
65
66
          auto seq = \_base.create(10);
67
68
          for(ea::sequence\_len\_t i = 0; i < 10; ++i)
```

```
69
              _base.set(seq, i, genes[i]);
70
71
72
73
           return seq;
         }
74
75
       private:
76
77
         static AnsiRandomNumberGenerator _g;
78
         static Base _base;
79
    };
80
    AnsiRandomNumberGenerator RouteFactory::_g;
81
82
     Base RouteFactory::_base;
83
84
    // process pipeline:
85
    int
    main(int argc, char *argv[])
86
87
       Base base;
88
89
       RouteFactory f;
90
       vector < Route > population;
91
       auto rnd = make_shared<ea::AnsiRandomNumberGenerator>();
92
       // create 100 routes:
93
94
       auto a = make_output_adapter(population);
95
96
       f.create_population(100, a);
97
98
       // create and process pipeline:
99
       auto source = make_input_adapter(population);
100
101
       cout << "mean_fitness_(parent):_" << mean<Base>(source) <<
           endl;
102
       cout << "median_fitness_(parent):" << median<Base>(source)
           << endl;
103
104
       vector < Route > children;
105
       auto cadapter = make_output_adapter(children);
106
107
       auto selection_a =
108
         SelectionElement < Base, SourceDivisor < Base >>
            (\mathbf{new}\ \mathsf{TournamentSelection} {<} \mathsf{Base}\,,\ \mathsf{std}:: \mathsf{less} {<} \mathbf{double} {>>} ())\,;
109
110
111
       auto selection_b =
         SelectionElement < Base, FixedSelectionSize < Base, 50>>(
112
113
           new DoubleTournamentSelection < Base, std::less < double >>()
                );
114
115
       auto crossover =
116
         CrossoverElement < Base >
            (new EdgeRecombinationCrossover<Base>());
117
118
119
       auto mutation =
```

```
120
         MutationElement<Base>
121
           (new SingleSwapMutation<Base>(), rnd);
122
123
      auto terminator = ForLoopTerminator < Base > (100);
124
      ITerminator < Base > & terminator ref = terminator;
125
       pipeline_process <Base > (source, cadapter, { & selection_a, &
126
          crossover, &selection_b, &mutation }, terminator_ref);
127
      source = make_input_adapter(children);
128
129
      cout << "mean_fitness_(parent):_" << mean<Base>(source) <<</pre>
130
      endl;
cout << "median_fitness_(parent):_" << median<Base>(source)
131
          << endl;
132
      // cleanup:
133
       dispose(base, begin(population), end(population));
134
      dispose(base, begin(children), end(children));
135
136
137
      return 0;
138
    }
```

# 4 Summary

As you have seen in this introduction LIBEA provides a basic framework to write evolutionary algorithms in a reliable way. There are some standard operators available which can be used with different kind of genomes.

The library has been written just for fun and I hope you might find it useful. To have a more detailed look at LIBEA you should generate the code documentation as described before in this document.