Elements of Programming

Transformations and their orbits

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Transformations and Their Orbits

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This chapter defines a transformation as a unary regular function from a type to itself. Successive applications of a transformation starting from an initial value determine an orbit of this value. Depending only on the regularity of the transformation and the finiteness of the orbit, we implement an algorithm for determining orbit structures that can be used in different domains. For example, it could be used to detect a cycle in a linked list or to analyze a pseudorandom number generator. We derive an interface to the algorithm as a set of related procedures and definitions for their arguments and results. This analysis of an orbit-structure algorithm allows us to introduce our approach to programming in the simplest possible setting.
Homogeneous predicates and operations

While there are functions from any sequence of types to any type, particular classes of signatures commonly occur. In this book we frequently use two such classes: **homogeneous predicates** and **operations**. Homogeneous predicates are of the form $T \times \ldots \times T \rightarrow \text{bool}$; operations are functions of the form $T \times \ldots \times T \rightarrow T$. While there are $n$-ary predicates and $n$-ary operations, we encounter mostly unary and binary homogeneous predicates and unary and binary operations.
Concepts *Predicate* and *UnaryPredicate*

A *predicate* is a functional procedure returning a truth value:

\[
\text{Predicate}(P) \triangleq \\
\text{FunctionalProcedure}(P) \\
\land \text{Codomain}(P) = \text{bool}
\]

A homogeneous predicate is one that is also a homogeneous function:

\[
\text{HomogeneousPredicate}(P) \triangleq \\
\text{Predicate}(P) \\
\land \text{HomogeneousFunction}(P)
\]

A *unary predicate* is a predicate taking one parameter:

\[
\text{UnaryPredicate}(P) \triangleq \\
\text{Predicate}(P) \\
\land \text{UnaryFunction}(P)
\]
An operation is a homogeneous function whose codomain is equal to its domain:

\[
Operation(Op) \triangleq \text{HomogeneousFunction}(Op)
\wedge \text{Codomain}(Op) = \text{Domain}(Op)
\]
Examples of operations

Examples of operations:

```c
int abs(int x) {
    if (x < 0) return -x; else return x;
} // unary operation

double euclidean_norm(double x, double y) {
    return sqrt(x * x + y * y);
} // binary operation

double euclidean_norm(double x, double y, double z) {
    return sqrt(x * x + y * y + z * z);
} // ternary operation
```

**Lemma**

euclidean_norm(x, y, z) = euclidean_norm(euclidean_norm(x, y), z)

This lemma shows that the ternary version can be obtained from the binary version. For reasons of efficiency, expressiveness, and, possibly, accuracy, the ternary version is part of the computational basis for programs dealing with three-dimensional space.
A procedure is *partial* if its definition space is a subset of the direct product of the types of its inputs; it is *total* if its definition space is equal to the direct product. We follow standard mathematical usage, where partial function includes total function. We call partial procedures that are not total *nontotal*. Implementations of some total functions are nontotal on the computer because of the finiteness of the representation. For example, addition on signed 32-bit integers is nontotal.
Dealing with nontotal functions

A nontotal procedure is accompanied by a precondition specifying its definition space. To verify the correctness of a call of that procedure, we must determine that the arguments satisfy the precondition. Sometimes, a partial procedure is passed as a parameter to an algorithm that needs to determine at runtime the definition space of the procedural parameter. To deal with such cases, we define a definition-space predicate with the same inputs as the procedure; the predicate returns true if and only if the inputs are within the definition space of the procedure. Before a nontotal procedure is called, either its precondition must be satisfied, or the call must be guarded by a call of its definition-space-space predicate.

Exercise

Implement a definition-space predicate for addition on 32-bit signed integers.
This chapter deals with unary operations, which we call *transformations*:

\[
\text{Transformation}(F) \triangleq \\
\text{Operation}(F) \\
\wedge \text{UnaryFunction}(F) \\
\wedge \text{DistanceType} : \text{Transformation} \rightarrow \text{Integer}
\]

We discuss \text{DistanceType} in the next section.
Transformations are self-composable: $f(x), f(f(x)), f(f(f(x)))$, and so on. The definition space of $f(f(x))$ is the intersection of the definition space and result space of $f$. This ability to self-compose, together with the ability to test for equality, allows us to define interesting algorithms.

When $f$ is a transformation, we define its powers as follows:

$$f^n(x) = \begin{cases} 
  x & \text{if } n = 0, \\
  f^{n-1}(f(x)) & \text{if } n > 0 
\end{cases}$$
Concept *Integer*

To implement an algorithm to compute $f^n(x)$, we need to specify the requirement for an integer type. We study various concepts describing integers in Chapter ???. For now we rely on the intuitive understanding of integers. Their models include signed and unsigned integral types, as well as arbitrary-precision integers, with these operations and literals:

<table>
<thead>
<tr>
<th>Specifications</th>
<th>C++</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sum</td>
<td>+</td>
</tr>
<tr>
<td>Difference</td>
<td>−</td>
</tr>
<tr>
<td>Product</td>
<td>·</td>
</tr>
<tr>
<td>Quotient</td>
<td>/</td>
</tr>
<tr>
<td>Remainder</td>
<td>mod</td>
</tr>
<tr>
<td>Zero</td>
<td>0</td>
</tr>
<tr>
<td>One</td>
<td>1</td>
</tr>
<tr>
<td>Two</td>
<td>2</td>
</tr>
</tbody>
</table>

where I is an integer type.
That leads to the following algorithm:

```cpp
template<typename F, typename N>
    requires(Transformation(F) && Integer(N))
    Domain(F) power_unary(Domain(F) x, N n, F f)
{
    // Precondition: n ⩾ 0 ∧ (∀i ∈ N) 0 < i ≤ n ⇒ f^i(x) is defined
    while (n != N(0)) {
        n = n - N(1);
        x = f(x);
    }
    return x;
}
```
Reachability, cyclic and terminal elements, and orbits

To understand the global behavior of a transformation, we examine the structure of its *orbits*: elements reachable from a starting element by repeated applications of the transformation. \( y \) is *reachable* from \( x \) under a transformation \( f \) if for some \( n \geq 0 \), \( y = f^n(x) \). \( x \) is *cyclic* under \( f \) if for some \( n \geq 1 \), \( x = f^n(x) \). \( x \) is *terminal* under \( f \) if and only if \( x \) is not in the definition space of \( f \). The *orbit* of \( x \) under a transformation \( f \) is the set of all elements reachable from \( x \) under \( f \).

**Lemma**

An orbit does not contain both a cyclic and a terminal element.

**Lemma**

An orbit contains at most one terminal element.
Distance types

If \( y \) is reachable from \( x \) under \( f \), the *distance* from \( x \) to \( y \) is the least number of transformation steps from \( x \) to \( y \). Obviously, distance is not always defined.

Given a transformation type \( F \), \( \text{DistanceType}(F) \) is an integer type large enough to encode the maximum number of steps by any transformation \( f \in F \) from one element of \( T = \text{Domain}(F) \) to another. If type \( T \) occupies \( k \) bits, there can be as many as \( 2^k \) values but only \( 2^k - 1 \) steps between distinct values. Thus if \( T \) is a fixed-size type, an integral type of the same size is a valid distance type for any transformation on \( T \). (Instead of using the distance type, we allow the use of any integer type in \texttt{power_unary}, since the extra generality does not appear to hurt there.)

It is often the case that all transformation types over a domain have the same distance type. In this case the type function \( \text{DistanceType} \) is defined for the domain type and defines the corresponding type function for the transformation types.
The existence of \texttt{DistanceType} leads to the following procedure:

\begin{verbatim}
  template<typename F>
    requires(Transformation(F))
  DistanceType(F) distance(Domain(F) x, Domain(F) y, F f)
  {
    // Precondition: \textit{y is reachable from} x \textit{under} f
    typedef DistanceType(F) N;
    N n(0);
    while (x != y) {
      x = f(x);
      n = n + N(1);
    }
    return n;
  }
\end{verbatim}
Orbits have different shapes. An orbit of $x$ under a transformation is

- **infinite** if it has no cyclic or terminal elements
- **terminating** if it has a terminal element
- **circular** if $x$ is cyclic
- **$\rho$-shaped** if $x$ is not cyclic, but its orbit contains a cyclic element

An orbit of $x$ is **finite** if it is not infinite. Figure 1 illustrates the various cases.
Orbit shapes

- Infinite
- Terminating
- Circular
- \( \rho \)-shaped

Figure: Orbit Shapes
Structure of orbits

The *orbit cycle* is the set of cyclic elements in the orbit and is empty for infinite and terminating orbits. The *orbit handle*, the complement of the orbit cycle with respect to the orbit, is empty for a circular orbit. The *connection point* is the first cyclic element, and is the first element of a circular orbit and the first element after the handle for a ρ-shaped orbit. The *orbit size* $o$ of an orbit is the number of distinct elements in it. The *handle size* $h$ of an orbit is the number of elements in the orbit handle. The *cycle size* $c$ of an orbit is the number of elements in the orbit cycle.

**Lemma**

\[ o = h + c \]
Distances

**Lemma**
The distance from any point in an orbit to a point in a cycle of that orbit is always defined.

**Lemma**
If $x$ and $y$ are distinct points in a cycle of size $c$,

$$c = \text{distance}(x, y, f) + \text{distance}(y, x, f)$$

**Lemma**
If $x$ and $y$ are points in a cycle of size $c$, the distance from $x$ to $y$ satisfies

$$0 \leq \text{distance}(x, y, f) < c$$
Finite orbit assumption

If we observe the behavior of a transformation, without access to its definition, we cannot determine whether a particular orbit is infinite: It might terminate or cycle back at any point. If we know that an orbit is finite, we can use an algorithm to determine the shape of the orbit. Therefore there is an implicit precondition of orbit finiteness for all the algorithms in this chapter.

There is, of course, a naive algorithm that stores every element visited and checks at every step whether the new element has been previously encountered. Even if we could use hashing to speed up the search, such an algorithm still would require linear storage and would not be practical in many applications. However, there is an algorithm that requires only a constant amount of storage.
The following analogy helps to understand the algorithm. If a fast car and a slow one start along a path, the fast one will catch up with the slow one if and only if there is a cycle. If there is no cycle, the fast one will reach the end of the path before the slow one. If there is a cycle, by the time the slow one enters the cycle, the fast one will already be there and will catch up eventually. Carrying our intuition from the continuous domain to the discrete domain requires care to avoid the fast one skipping past the slow one.¹

¹?, page 7 attributes this algorithm to Robert W. Floyd.
The discrete version of the algorithm is based on looking for a point where fast meets slow. The *collision point* of a transformation \( f \) and a starting point \( x \) is the unique \( y \) such that

\[
y = f^n(x) = f^{2n+1}(x)
\]

and \( n \geq 0 \) is the smallest integer satisfying this condition. This definition leads to an algorithm for determining the orbit structure that needs one comparison of fast and slow per iteration. To handle partial transformations, we pass a definition-space predicate to the algorithm:
```cpp
template<typename F, typename P>
    requires(Transformation(F) && UnaryPredicate(P) &&
             Domain(F) == Domain(P))
Domain(F) collision_point(const Domain(F)& x, F f, P p)
{
    // Precondition: p(x) ⇔ f(x) is defined
    if (!p(x)) return x;
    Domain(F) slow = x;  // slow = f^0(x)
    Domain(F) fast = f(x);  // fast = f^1(x)
    // n ← 0 (completed iterations)
    while (fast != slow) {
        slow = f(slow);  // slow = f^{n+1}(x) ∧ fast = f^{2n+1}(x)
        if (!p(fast)) return fast;
        fast = f(fast);  // slow = f^{n+1}(x) ∧ fast = f^{2n+2}(x)
        if (!p(fast)) return fast;
        fast = f(fast);  // slow = f^{n+1}(x) ∧ fast = f^{2n+3}(x)
        // n ← n + 1
    }
    return fast;  // slow = f^{n}(x) ∧ fast = f^{2n+1}(x)
    // Postcondition: return value is terminal point or collision point
}
```
Partial correctness of collision_point

We establish the correctness of collision_point in three stages: (1) verifying that it never applies f to an argument outside the definition space; (2) verifying that if it terminates, the postcondition is satisfied; and (3) verifying that it always terminates.

While f is a partial function, its use by the procedure is well defined, since the movement of fast is guarded by a call of p. The movement of slow is unguarded, because by the regularity of f, slow traverses the same orbit as fast, so f is always defined when applied to slow.

The annotations show that if, after \( n \geq 0 \) iterations, fast becomes equal to slow, then fast = \( f^{2n+1}(x) \) and slow = \( f^n(x) \). Moreover, n is the smallest such integer, since we checked the condition for every \( i < n \).
Termination of collision_point

If there is no cycle, \( p \) will eventually return false because of finiteness. If there is a cycle, \( \text{slow} \) will eventually reach the connection point (the first element in the cycle). Consider the distance \( d \) from \( \text{fast} \) to \( \text{slow} \) at the top of the loop when \( \text{slow} \) first enters the cycle: \( 0 \leq d < c \). If \( d = 0 \), the procedure terminates. Otherwise the distance from \( \text{fast} \) to \( \text{slow} \) decreases by 1 on each iteration. Therefore the procedure always terminates; when it terminates, \( \text{slow} \) has moved a total of \( h + d \) steps.
The following procedure determines whether an orbit is terminating:

```cpp
template<typename F, typename P>
  requires(Transformation(F) && UnaryPredicate(P) &&
           Domain(F) == Domain(P))
bool terminating(const Domain(F)& x, F f, P p)
{
    // Precondition: \( p(x) \iff f(x) \) is defined
    return !p(collision_point(x, f, p));
}
```
Sometimes, we know either that the transformation is total or that the orbit is nonterminating for a particular starting element. For these situations it is useful to have a specialized version of `collision_point`:

```cpp
template<typename F>
  requires(Transformation(F))
Domain(F)
collision_point_nonterminating_orbit(const Domain(F)& x, F f)
{
  Domain(F) slow = x; // slow = f^0(x)
  Domain(F) fast = f(x); // fast = f^1(x)
  // n ← 0 (completed iterations)
  while (fast != slow) {
    slow = f(slow); // slow = f^{n+1}(x) ∧ fast = f^{2n+1}(x)
    fast = f(fast); // slow = f^{n+1}(x) ∧ fast = f^{2n+2}(x)
    fast = f(fast); // slow = f^{n+1}(x) ∧ fast = f^{2n+3}(x)
    // n ← n + 1
  }
  return fast; // slow = f^n(x) ∧ fast = f^{2n+1}(x)
  // Postcondition: return value is collision point
}
```
Position of \textit{collision point}

In order to determine the cycle structure—handle size, connection point, and cycle size—we need to analyze the position of the collision point.

When the procedure returns the collision point

\[ f^n(x) = f^{2n+1}(x) \]

\( n \) is the number of steps taken by \textit{slow}, and \( 2n + 1 \) is the number of steps taken by \textit{fast}.

\[ n = h + d \]

where \( h \) is the handle size and \( 0 \leq d < c \) is the number of steps taken by \textit{slow} inside the cycle. The number of steps taken by \textit{fast} is

\[ 2n + 1 = h + d + qc \]

where \( q > 0 \) is the number of full cycles completed by \textit{fast} when \textit{slow} enters the cycle. Since \( n = h + d \),

\[ 2(h + d) + 1 = h + d + qc \]
Position of collision\_point, continued

Simplifying gives

\[ qc = h + d + 1 \]

Let us represent \( h \) modulo \( c \):

\[ h = mc + r \]

with \( 0 \leq r < c \). Substitution gives

\[ qc = mc + r + d + 1 \]

or

\[ d = (q - m)c - r - 1 \]

\( 0 \leq d < c \) implies

\[ q - m = 1 \]

so

\[ d = c - r - 1 \]

and \( r + 1 \) steps are needed to complete the cycle.
Therefore the distance from the collision point to the connection point is

\[ e = r + 1 \]
Distinguishing circular from \( \rho \)-shaped case

In the case of a circular orbit \( h = 0, r = 0 \), and the distance from the collision point to the beginning of the orbit is

\[ e = 1 \]

Circularity, therefore, can be checked with the following procedures:

```cpp
template<typename F>
    requires(Transformation(F))
bool circular_nonterminating_orbit(const Domain(F)& x, F f)
{
    return x == f(collision_point_nonterminating_orbit(x, f));
}

template<typename F, typename P>
    requires(Transformation(F) && UnaryPredicate(P) &&
                 Domain(F) == Domain(P))
bool circular(const Domain(F)& x, F f, P p)
{
    // Precondition: \( p(x) \Leftrightarrow f(x) \) is defined
    Domain(F) y = collision_point(x, f, p);
    return p(y) && x == f(y);
}
```
Finding connection point

We still don’t know the handle size $h$ and the cycle size $c$. Determining the latter is simple once the collision point is known: Traverse the cycle and count the steps.

To see how to determine $h$, let us look at the position of the collision point:

$$f^{h+d}(x) = f^{h+c-r-1}(x) = f^{mc+r+c-r-1}(x) = f^{(m+1)c-1}(x)$$

Taking $h+1$ steps from the collision point gets us to the point $f^{(m+1)c+h}(x)$, which equals $f^h(x)$, since $(m+1)c$ corresponds to going around the cycle $m+1$ times. If we simultaneously take $h$ steps from $x$ and $h+1$ steps from the collision point, we meet at the connection point. In other words, the orbits of $x$ and 1 step past the collision point converge in exactly $h$ steps, which leads to the following sequence of algorithms:
template<typename F>
    requires(Transformation(F))
Domain(F) convergent_point(Domain(F) x0, Domain(F) x1, F f)
{
    // Precondition: (\exists n \in \text{DistanceType}(F)) n \geq 0 \land f^n(x0) = f^n(x1)
    while (x0 != x1) {
        x0 = f(x0);
        x1 = f(x1);
    }
    return x0;
}
template<typename F>
    requires(Transformation(F))
Domain(F)
connection_point_nonterminating_orbit(const Domain(F)& x, F f)
{
    return convergent_point(
        x,
        f(collision_point_nonterminating_orbit(x, f)),
        f);
}
connection_point

template<typename F, typename P>
    requires(Transformation(F) && UnaryPredicate(P) &&
             Domain(F) == Domain(P))
Domain(F) connection_point(const Domain(F)& x, F f, P p)
{
    // Precondition: p(x) ⇔ f(x) is defined
    Domain(F) y = collision_point(x, f, p);
    if (!p(y)) return y;
    return convergent_point(x, f(y), f);
}
Determining whether orbits intersect

**Lemma**
If the orbits of two elements intersect, they have the same cyclic elements.

**Exercise**
Design an algorithm that determines, given a transformation and its definition-space predicate, whether the orbits of two elements intersect.

**Exercise**
The precondition of `convergent_point` ensures termination. Implement an algorithm `convergent_point_guarded` for use when that precondition is not known to hold, but there is an element in common to the orbits of both $x_0$ and $x_1$. 
Representing orbit sizes

The natural type to use for the sizes $o$, $h$, and $c$ of an orbit on type $T$ would be an integer count type large enough to count all the distinct values of type $T$. If a type $T$ occupies $k$ bits, there can be as many as $2^k$ values, so a count type occupying $k$ bits could not represent all the counts from 0 to $2^k$. There is a way to represent these sizes by using distance type.
Representing sizes

An orbit could potentially contain all values of a type, in which case \( o \) might not fit in the distance type. Depending on the shape of such an orbit, \( h \) and \( c \) would not fit either. However, for a \( \rho \)-shaped orbit, both \( h \) and \( c \) fit. In all cases each of these fits: \( o - 1 \) (the maximum distance in the orbit), \( h - 1 \) (the maximum distance in the handle), and \( c - 1 \) (the maximum distance in the cycle). That allows us to implement procedures returning a triple representing the complete structure of an orbit, where the members of the triple are as follows:

<table>
<thead>
<tr>
<th>Case</th>
<th>( m_0 )</th>
<th>( m_1 )</th>
<th>( m_2 )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Terminating</td>
<td>( h - 1 )</td>
<td>0</td>
<td>terminal element</td>
</tr>
<tr>
<td>Circular</td>
<td>0</td>
<td>( c - 1 )</td>
<td>( x )</td>
</tr>
<tr>
<td>( \rho )-shaped</td>
<td>( h )</td>
<td>( c - 1 )</td>
<td>connection point</td>
</tr>
</tbody>
</table>
template<typename F> 
    requires(Transformation(F)) 
triple<DistanceType(F), DistanceType(F), Domain(F)> 
orbit_structure_nonterminating_orbit(const Domain(F)& x, F f) 
{
    typedef DistanceType(F) N;
    Domain(F) y = connection_point_nonterminating_orbit(x, f);
    return triple<N, N, Domain(F)>(distance(x, y, f), 
                                    distance(f(y), y, f), 
                                    y);
}
template<typename F, typename P>
    requires(Transformation(F) &&
             UnaryPredicate(P) && Domain(F) == Domain(P))
triple<DistanceType(F), DistanceType(F), Domain(F)>
orbit_structure(const Domain(F)& x, F f, P p)
{
    // Precondition: p(x) ⇔ f(x) is defined
    typedef DistanceType(F) N;
    Domain(F) y = connection_point(x, f, p);
    N m = distance(x, y, f);
    N n(0);
    if (p(y)) n = distance(f(y), y, f);
    // Terminating: m = h − 1 ∧ n = 0
    // Otherwise: m = h ∧ n = c − 1
    return triple<N, N, Domain(F>>(m, n, y);
}
Exercises

Exercise
Derive formulas for the count of different operations ($f$, $p$, equality) for the algorithms in this chapter.

Exercise
Use `orbit_structure_nonterminating_orbit` to determine the average handle size and cycle size of the pseudorandom number generators on your platform for various seeds.
Actions

Algorithms often use a transformation $f$ in a statement like

$$x = f(x);$$

Changing the state of an object by applying a transformation to it defines an *action* on the object. There is a duality between transformations and the corresponding actions: An action is definable in terms of a transformation, and vice versa:

```cpp
void a(T& x) { x = f(x); } // action from transformation
```

and

```cpp
T f(T x) { a(x); return x; } // transformation from action
```

Despite this duality, independent implementations are sometimes more efficient, in which case both action and transformation need to be provided. For example, if a transformation is defined on a large object and modifies only part of its overall state, the action could be considerably faster.
Exercise

Rewrite all the algorithms in this chapter in terms of actions.
Another way to detect a cycle is to repeatedly test a single advancing element for equality with a stored element, while replacing the stored element at ever increasing intervals. This and other ideas are described in ?, ?, and ?. Implement other algorithms for orbit analysis, compare their performance for different applications, and develop a set of recommendations for selecting the appropriate algorithm.
Abstraction allowed us to define abstract procedures that can be used in different domains. Regularity of types and functions is essential to make the algorithms work: fast and slow follow the same orbit because of regularity. Developing nomenclature is essential (e.g., orbit kinds and sizes). Affiliated types, such as distance type, need to be precisely defined.
End of chapter