15-150 Fall 2015 Homework 10

Out: Wednesday, 11 November 2015 Due: Tuesday, 17 November 2015 at 23:59 EST

1 Introduction

In this homework, you will see two simple implementations of a sequence library, use reduce to generalize find, and later implement the Barnes-Hut approximation algorithm for the n-body problem.

1.1 Getting The Homework Assignment

The starter files for the homework assignment have been distributed through our git repository, as usual.

1.2 Submitting The Homework Assignment

Submissions will be handled through Autolab, at https://autolab.cs.cmu.edu

In preparation for submission, your hw/10 directory should contain a file named exactly hw10.pdf containing your written solutions to the homework.

To submit your solutions, run make from the hw/10 directory (that contains a code folder and a file hw10.pdf). This should produce a file hw10.tar, containing the files that should be handed in for this homework assignment. Open the Autolab web site, find the page for this assignment, and submit your hw10.tar file via the "Handin your work" link.

The Autolab handin script does some basic checks on your submission: making sure that the file names are correct; making sure that no files are missing; making sure that your code compiles cleanly. Note that the handin script is *not* a grading script—a timely submission that passes the handin script will be graded, but will not necessarily receive full credit. You can view the results of the handin script by clicking the number corresponding to the "check" section of your latest handin on the "Handin History" page. If this number is 0.0, your submission failed the check script; if it is 1.0, it passed.

Remember that your written solutions must be submitted in PDF format—we do not accept MS Word files or other formats.

All the code that you want to have graded for this assignment should be contained in barnes-hut.sml, shrubseq.sml,sizeseq.sml, and tester.sml, and must compile cleanly. If you have a function that happens to be named the same as one of the required functions but does not have the required type, it will not be graded.

1.3 Due Date

This assignment is due on Tuesday, 17 November 2015 at 23:59 EST. Remember that you may use a maximum of one late day per assignment, and that you are allowed a total of three late days for the semester.

1.4 Methodology

You must use the five step methodology discussed in class for writing functions, for **every** function you write in this assignment. Recall the five step methodology:

- 1. In the first line of comments, specify the type of the function.
- 2. In the second line of comments, specify via a REQUIRES clause any assumptions about the arguments to be passed to the function.
- 3. In the third line of comments, specify via an ENSURES clause what the function computes (what it returns when applied to an argument that satisfies the assumptions in REQUIRES).
- 4. Implement the function.
- 5. Provide testcases, generally in the format val <return value> = <function> <argument value>.

For example, for the factorial function presented in lecture:

```
(* fact : int -> int
  * REQUIRES: n >= 0
  * ENSURES: fact(n) ==> n! *)
fun fact (0 : int) : int = 1
  | fact (n : int) : int = n * fact(n-1)

(* Tests: *)
val 1 = fact 0
val 6 = fact 3
val 720 = fact 6
```

1.4.1 Emphatic Warning

CM will not return cleanly if any of the files listed in the sources have no code in them. Because we want you to learn how to write modules from scratch, we have handed out a few files that are empty except for a few place holder comments. That means that there are a few files in the sources.cm we handed out that are commented out, so that when you first get your tarball CM.make "sources.cm" will work cleanly.

You must uncomment these lines as you progress through the assignment! If you forget, it will look like your code compiles cleanly even though it almost certainly doesn't.

2 Tree Sequences

In lecture we covered an abstract data structure called Sequences, which you made use of in the last assignment (and will continue to use for the rest of the course). So far, we have avoided discussing implementations of a Sequence library; thanks to representational independence¹, the implementation details are not important for most of the code you will write in this class. The next few problems will give you a brief introduction to implementing sequences using a familiar data structure: binary trees!

We provide the following (shortened) signature in the file tseq.sig:

```
signature TSEQ =
sig
    type 'a seq

exception Range

val length : 'a seq -> int
    val nth : int -> 'a seq -> 'a
    val tabulate : (int -> 'a) -> int -> 'a seq
end
```

These are the only functions on sequences you will use for this part of the assignment. You should not use the Seq library functions, nor implement any functions outside this signature (but keep them in mind for the second part of the assignment, Barnes-Hut!)

2.1 Sequential Shrubberies

We can implement the above signature using a variety of underlying data structures; with correct code, the only difference for a user should be the run-time of our functions. In the file **shrubseq.sml** we have provided the beginnings of a sequence implementation using the following datatype:

For a type t, a value of type t seq is a binary tree with data of type t stored at the leaves. Additionally, we have the following *balance invariant*: For any Node(L,R) the number of leaves in L and in R differ by at most 1. Your code must ensure that this invariant is always

¹Recall that representational independence is the idea that we can use multiple implementations of a data structure to represent the same signature, in this case ordered lists of elements (sequences).

maintained. Note that since we do not support an insert or delete operation on sequences, it is enough to guarantee that our trees are balanced when we first create them.

Task 2.1 (2 pts). We have provided an implementation of the length function - it should look very familiar from your previous work with binary trees. What is the work and span of length? Briefly explain why.

Task 2.2 (10 pts). Implement the **nth** function. Your implementation must use **length**, but NO other helper functions, and will have work in O(n) and span in $O((\log n)^2)$. Remember that sequences are indexed beginning at zero; your code should raise a Range exception if the argument index is invalid for the given sequence.

Note: There is a way to implement nth in O(n) work and $O(\log n)$ span using a helper function and certain assumptions about the underlying tree—you should **not** write this implementation. Your code for this task should not be very long or complex; in the next section, you will see an alternative optimization that makes this simple implementation more efficient.

Task 2.3 (10 pts). Implement the tabulate function. DO NOT use length, nth, or any other helper functions. Your implementation must have work in $O(n \log n)$ and span in $O((\log n)^2)$ (you may assume that the input function has O(1) work and span). Remember to ensure that your trees are balanced!

(There is also an implementation of tabulate with O(n) work and $O(\log n)$ span, but you do not not need to write that unless you want to.)

Do not forget to test your implementation; you can test your functions together by using tabulate to create a simple sequence, then checking for correct values with length and nth.

Clearly, the above implementation is not ideal; your answer to the first question above should give you a clue as to why. There are several ways to improve our implementation; in the next task, you will do exactly that, focusing on our implementation of length.

2.2 Sizable Shrubberies

We would prefer for our sequences to implement length with constant work and span. One simple way to achieve this is to modify our underlying datatype. In the file sizeseq.sml we provide the following datatype:

As before, for any type t a value of type t seq is a balanced binary tree with data stored at the leaves. This time, however we store an additional value of type int in each node. It is your responsibility to figure out what the additional value represents. Make sure, however, that your implementation can easily meet the Big-O bounds we have given you for each function.

Task 2.4 (2 pts). Implement length in sizeseq.sml; your implementation must have O(1) work and span.

Task 2.5 (1 pts). Implement **nth** so that the function has work and span in $O(\log n)$.

From the last two problems, you should see how an improvement in one part of the code can lead to simple yet fast code in other areas. However, this comes at a slight trade-off.

Task 2.6 (2 pts). Implement tabulate in sizeseq.sml; you should be able to re-use most of your code from the previous section.

Task 2.7 (3 pts). Do the work and span of tabulate change, and if so, what are the new Big-O bounds? Briefly explain why, and describe the cost trade-off that occurs between tabulate and length (hint: consider more than just time).

Remember to test your implementations of length, nth, and tabulate as before.

These two implementations show how we can achieve O(1) time for length and a reasonable run-time for some of our functions using balanced trees, but in other cases we are restricted by our choice of data structure. Using alternative implementations with arrays or vectors (like the Seq library you will use for Barnes-Hut) we can actually achieve O(1) work for nth and span for tabulate. In this class we will focus primarily on using sequences for functional programming, but if you are interested you can learn more about this type of problem through further courses in data structures and algorithms such as 15-210.

3 Just a Monoid in the Category of Endofunctors

Recall the function findOne from Homework 7. Recall that using findOne, we implemented findTwo. In this section, we will generalize it to find n unique elements using reduce.

First, consider the infixed function

with the following definition:

infix >=>

Notice that >=> is associative, that is, f >=> (g >=> h) = (f >=> g) >=> h. You can think of >=> as pipelining functions, and if any of them evaluate to NONE, then the entire thing evaluates to NONE.

Task 3.1 (2 pts).

What is the identity/neutral element of >=>, that is, find the function id such that

$$\forall (f: 'a \rightarrow 'a \text{ option}). f >=> id = id >=> f = f$$

You should put your solution in the pdf, not in the code. You do not need to prove correctness or give justification.

Notice that because we now have an identity element for >=>, we can use it with reduce.

Task 3.2 (3 pts).

To make sure you understand what is really going on with >=>, write a simple function equivalent to

$$(fn \ 0 \Rightarrow NONE \mid x \Rightarrow SOME(3 \mod x)) \Rightarrow (fn \ x \Rightarrow SOME(x+1))$$

When we say simple function, we mean a function that has no pattern matching on the result of a function application. For example, $(fn 5 \Rightarrow SOME(10000) \mid x \Rightarrow SOME(x+1))$ is a simple function. You should put your solution in the pdf, not in the code. You do not need to give a proof or justification.

Now, let us generalize find to n elements. Recall the function

with the behaviour

 $\mbox{findOne p T s } k = \begin{cases} \mbox{s } \mbox{v} & \mbox{where v is a value in T such that p v = true, if there is one.} \\ k \mbox{ () otherwise} \end{cases}$

You can find the definition of findOne and tree inside the structure FindOne

Task 3.3 (15 pts).

In the functor FindMany, use Seq.reduce, Seq.tabulate, >=>, and findOne to implement

such that findN p n T evaluates to SOME L if there exists n unique² elements that satisfy p in T, with L containing exactly n unique elements that satisfy p in T. Otherwise, it evaluates to NONE. We can always find a list of 0 elements in the empty tree³. We have provided an infixed helper function inside:''a * ''a list -> bool for your convenience. findN should not be recursive.

Hint: generate a sequence of functions and reduce with >=>.

²When we say unique, we mean elements distinct under =

³The trees we give you have no empty tree, but your code should work for an arbitrary tree with associated findOne function

4 Barnes-Hut

For the remainder of this assignment, you will be implementing the Barnes-Hut algorithm for efficient n-body simulation, an extension of the ideas you learned in lecture.

4.1 Preface

The following points explore the various libraries you will be using for this homework. Please read through each section to ensure you fully understand the code infrastructure for the coming tasks.

4.1.1 The plane problem

For n-body simulations, we need a representation of numbers (or scalars) to talk about distance between bodies in a 2D plane. We have two options for numerical representation:

1. **Reals.** Reals are fast to compute and work well for visualization, but alas they are only a floating point precision approximation to real numbers —not actual real numbers in the mathematical sense. In particular, addition and multiplication on values of type real are not always associative, and multiplication does not always distribute over addition. This means that you can do the same sequence of operations in two slightly different orders and get drastically different results:

```
- 10E30 + (~10E30 + 1.0);

val it = 0.0 : real

- (10E30 + ~10E30) + 1.0;

val it = 1.0 : real
```

Hence, reals make it difficult to test our code for correctness.

2. **Rationals.** Rationals (or all numbers which we can represent as a fraction $\frac{a}{b}$ where $a, b \in \mathbb{Z}$) are not built into SML but give us a precise representation for non-integer numeric values. Consequently, rationals make testing more accurate than if we used reals.

Computation on rationals, however, is slow because we have to do arbitrary precision integer arithmetic. Moreover, we cannot compute Euclidean distance (i.e., in a 2D plane) with rationals. If x and y are rational numbers, it is not necessarily the case that $\sqrt{x^2 + y^2}$ is a rational number. We can, however, compute the Manhattan distance between points⁴.

⁴http://en.wikipedia.org/wiki/Taxicab_geometry

4.1.2 The plane solution

Rather than choose one or the other, we will instead parameterize our Barnes-Hut algorithm by giving it an implementation of numbers inside a package called Plane. Plane contains a number of useful functions and types, including Plane.scalar, Plane.point, and Plane.vec for scalars, points, and vectors, respectively.

We have provided to you two implementations of Plane using reals and rationals which you can find in lib/realplaneargs.sml and lib/ratplaneargs.sml, respectively. The file lib/space.sig lists all of the functions common to both implementations of the plane. The lib/makeplane.sml and lib/makescalar.sml also have some common functions to the Plane signature which you may find useful.

4.1.3 Scalar library

The type Scalar.scalar is equipped with the following functions:

- Scalar.plus: Scalar.scalar * Scalar.scalar -> Scalar.scalar which computes the sum of two scalars.
- Scalar.minus : Scalar.scalar * Scalar.scalar -> Scalar.scalar which computes the difference of two scalars.
- Scalar.times: Scalar.scalar * Scalar.scalar -> Scalar.scalar which computes the product of two scalars.
- Scalar.divide: Scalar.scalar * Scalar.scalar -> Scalar.scalar which computes the quotient of two scalars.
- Scalar.compare: Scalar.scalar * Scalar.scalar -> order which computes the ordering between two scalars.
- Scalar.fromRatio : IntInf.int * IntInf.int -> Scalar.scalar Scalar.fromRatio (x,y) evaluates to the value of type Scalar.scalar which represents $\frac{x}{y}$.
- Scalar.toString: Scalar.scalar -> string which computes a string representation of a scalar.

There are other helper functions in the file implemented in terms of these; see lib/scalar.sig for a description. By using only the above operations, your code will work with either implementation of the plane.

4.1.4 Point and vector libraries

In order to write our implementation of the Barnes-Hut algorithm, we need several operations on vectors and points in space, many of which we discussed in lecture. The type Plane.point

is used to represent a point in space, and the type Plane.vec is used to represent vectors of velocity, acceleration, etc. Whatever the definition of Scalar.scalar is, we define the type of points and vectors as in lecture (along with a datatype constructor):

datatype Plane.point = Plane.Coord of Scalar.scalar * Scalar.scalar
datatype Plane.vec = Plane.Vec of Scalar.scalar * Scalar.scalar

This uses Scalar.scalar for numbers. Note you may see the type Plane.scalar while debugging - this is defined to be the same as Scalar.scalar. Here we give a brief description of some of the functions you may need on this assignment:

- Plane.distance: Plane.point -> Plane.point -> Scalar.scalar
 Plane.distance p1 p2 evaluates to the distance between the points p1 and p2.
- Plane.midpoint: Plane.point -> Plane.point -> Plane.point Plane.midpoint p1 p2 evaluates to the midpoint of the points p1 and p2.
- Plane.head: Plane.vec -> Plane.point
 Plane.head v evaluates to the point that corresponds to the displacement of v from the origin.
- Plane.sum : ('a -> Plane.vec) -> 'a Seq.seq -> Plane.vec Plane.sum f s evaluates to the vector that corresponds to the sum of the sequence of vectors that results from mapping f on s. This corresponds to the mathematical notion of a sum of vectors, sum f $\langle s_1, \ldots, s_n \rangle = \sum_{i=1}^n f(s_i)$.

4.1.5 Other libraries

There is a lot more starter code for this assignment than for any previous assignment. Only edit the files barnes_hut.sml and tester.sml. When writing your solution, you should use functions specified in:

- lib/bbox.sig
- lib/space.sig
- lib/scalar.sig
- lib/mechanics.sig
- ../../src/sequence/sequencecore.sig
- ../../src/sequence/sequence.sig

The signature in lib/bbox.sig is the same as the BOX signature that you implemented in Homework 9 with some additional functions. Do not use your own implementation for this homework. We will be releasing an implementation for you to use on Thursday November 12, 2015 due to the late day for the previous homework.

4.1.6 Testing

You have two methods for testing your code:

- If you want to check your code for correctness, you will use the rational implementation of the Plane (see Section 4.6.1).
- If you want to visualize your n-body simulation, you will use the real implementation of the Plane (see Section 4.6.2).

4.2 Naïve simulation

Recall that the pieces of information we need about a body in space are its mass, location, and velocity. This is represented by the type definition

```
type body = Plane.Scalar.scalar * Plane.point * Plane.vec
```

The type body is used to represent the different bodies in the *n*-body simulation. Specifically, in an expression (m, p, v) of type body, m is the mass of the body, p is its position, and v is the vector representing its velocity.

The naive implementation that we gave in lecture has been rewritten using planes and the following function:

```
accelerations : body Seq.seq -> Plane.vec Seq.seq
```

in naiveNBody.sml. This function transforms a sequence of bodies into a sequence in which the element at position i represents the acceleration for the i-th body.

However, this implementation is accurate on large inputs but unacceptably slow for an actual simulation. You will implement a more efficient approximation to find accelerations.

4.3 The Barnes-Hut algorithm

Barnes-Hut groups bodies into quadrants and uses a threshold value θ to determine whether each individual body is "far enough" away from a group of other bodies. If so, Barnes-Hut groups the other bodies into a big pseudobody and uses that for the acceleration calculation instead. This results in a loss of accuracy, but a dramatic speedup in terms of runtime—while the old algorithm had work in $O(n^2)$, this algorithm's work is in $O(n \log n)$ if the threshold value is well-chosen.

To calculate the effect of a pseudobody on another body, it is important to know the total mass of all the bodies represented by the pseudobody and also their center of mass or barycenter. Therefore, when we form a pseudobody, we will compute a tuple (m, c) such that m: Scalar.scalar is the total mass of the bodies and c: Plane.point is the barycenter. To compute the barycenter, we compute a weighted average of the vectors corresponding to the displacement of each body's position from the origin. For example, if the positions are given by the set $\{p_i \mid i \in I\}$ and the corresponding masses are given by the set $\{m_i \mid i \in I\}$ then we compute the following vector:

$$\mathbf{R} = \frac{\sum_{i \in I} m_i \mathbf{r}_i}{\sum_{i \in I} m_i}$$

where \mathbf{r}_i is the vector corresponding to the displacement of position p_i from the origin. The barycenter is then the point that results from displacing the origin point by \mathbf{R} .

Given the total mass and barycenter, we approximate the acceleration due to all the bodies in the group as the acceleration due to a single body located at the barycenter with mass equal to the total mass.

4.3.1 Computing the barycenter

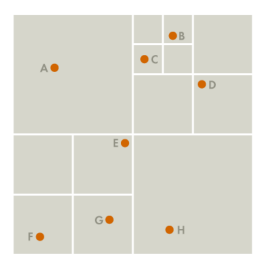
We will begin by writing the **barycenter** function to compute the total mass and barycenter of a sequence of pairs of masses and points.

Task 4.1 (10 pts). Write the function

that computes the pair (m, c) where m is the total mass of the bodies in the sequence (i.e., the sum of the first components of the pairs) and c is the barycenter. You should use the provided scale_point function to compute a weighted vector from a mass and position. You should also use the Plane.sum function.

4.3.2 Grouping bodies

To group bodies, we will partition them into quadrants. Starting at the center of the plane we are considering, we divide the field into quadrants, then recursively group the bodies in each quadrant, stopping when a region has either zero or one body in it. Here's a sample division of a set of bodies⁵:



Note: this diagram does not precisely represent what your BH tree looks like, because although we use quadrants for dividing the bodies, we group the bodies within a quadrant by their bounding box and then subdivide that box.

This process yields a tree-structured division of space with the following datatype:

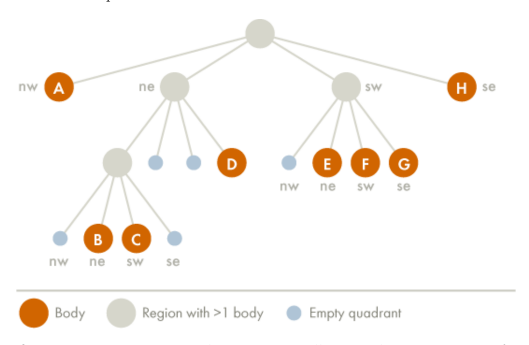
```
datatype bhtree =
    Null
    | Singleton of body
    | Box of (Scalar.scalar * Plane.point) * Scalar.scalar * (bhtree Seq.seq)
```

⁵Credit: http://arborjs.org/docs/barnes-hut

Null represents a region with no bodies in it. Singleton b represents a region with exactly the body b in it. For Box ((m, c), bd, sq), we have:

- m is the total mass of the bodies contained in the region.
- c is the barycenter of the bodies contained in the region.
- bd is the diameter of the bounding box of the region. A bounding box for a set of bodies is a rectangular region which contains all the bodies in the set. Some functions will reference the type BB.box which represents the bounding box, and you will find functions related to the bounding box in lib/bbox.sig. For our tree, however, you will only store the diameter of the bounding box, not the box itself.
- sq is the sequence of the four quadrants in the region. The invariant is that this sequence is always of length four and that the four child bhtree's are, in order, the top-left (also called nw), top-right (ne), bottom-left (sw), and bottom-right (se) quadrants of the region, respectively.

So the above example would have a tree that looks like:



As a first step in constructing this tree, we will write the quadrantize function to partition a sequence of bodies into four sub-sequences: one for each quadrant.

Task 4.2 (10 pts). Write the function

quadrantize : BB.box -> body Seq.seq -> body Seq.seq Seq.seq

that takes a bounding box, a sequence of bodies in that box, and returns a sequence with 4 inner sequences of bodies. The sequence at index i should containing all the bodies in quadrant i, where the quadrants are numbered starting at 0 in the following order: top-left (nw) has number 0, top-right (ne) has number 1, bottom-left (sw) has number 2, and bottom-right (se) has number 3. Note that you must use the function BB.quadrant to determine which quadrant each body belongs to. You may find taking a look at lib/bbox.sig helpful.

4.3.3 Growing the tree

We now have the tools we need to compute a bhtree from a sequence of bodies.

Task 4.3 (10 pts). Write the function

```
compute_tree : body Seq.seq -> bhtree
```

such that compute_tree s evaluates to T, where T is the tree decomposition of s. You may assume that no two bodies in s occupy the same position (*i.e.*, have equal position components). Your implementation does not have to handle arguments that violate this condition. You must use Seq.map to compute the recursive calls in parallel!

Note 1: you will need to compute the bounding box for the given bodies and store its diameter inside Boxes of your tree. There are certain helper functions which you may find useful for this task. First, you have

```
position : body -> Plane.point
```

which returns the position of a body. Second, you have

```
BB.hull : Plane.point Seq.seq -> BB.box
```

which given a sequence of points returns a bounding box containing all points in the sequence (BB.hull raises BB.Invalid if the input sequence is empty). Lastly, you have

```
BB.diameter : BB.box -> Scalar.scalar
```

which given a bounding box returns the diameter of the bounding box.

Note 2: If a large set of bodies is partitioned into several subsets and the barycenter of each subset is known, the barycenter of the whole set can be more efficiently computed as the barycenter of the barycenters of the subsets. In this instance, the barycenter of the bodies in a bounding box can be computed as the barycenter of the four quadrants' barycenters. You should use this observation to compute the barycenter in the recursive case of compute_tree by applying the barycenter function you wrote earlier to the sequence of the barycenters of the quadrants given by the results of the recursive calls. You may find the function center_of_mass: bhtree -> Scalar.scalar * Plane.point helpful to project the relevant data from the sequence of recursive results.

4.3.4 Computing acceleration

Now that we can calculate the tree determined by a group of bodies, we can use it to efficiently compute an approximation of the acceleration of all the bodies at this particular timestep. This brings us back to the threshold value θ mentioned above.

The reason Barnes-Hut is more efficient than the naïve approach to the n-body problem is that it does not compute the exact acceleration—instead, it uses θ to determine exactly how precise to be. Whenever your algorithm reaches a region with more than one body in it (that is, a Box in the tree), it checks to see if $\frac{d}{r} \leq \theta$ (where d is the diameter of the region's bounding box and r is the distance from the body being checked to the region's barycenter). If it is, then the region is treated as one large body located at its barycenter (which we have conveniently already calculated!). Otherwise, the region gets decomposed into quadrants and the respective accelerations from the bodies in each quadrant are computed recursively, combined, and returned.

Task 4.4 (10 pts). Write the function

groupable : Plane.point -> Plane.point -> Scalar.scalar -> Scalar.scalar -> bool

such that given a point p, a location c of a pseudobody, a diameter bd for the pseudobody's bounding box and a threshold t, groupable p c bd t evaluates to true if $\frac{d}{r} \leq \theta$ and false otherwise. Recall that in this equation, d refers to the diameter of the pseudobody's bounding box, r refers to the distance between the point and the pseudobody, and θ corresponds to the threshold argument t. You may find the function Plane.distance useful.

Task 4.5 (10 pts). Write the function

bh_acceleration : bhtree -> Scalar.scalar -> body -> vec

such that bh_acceleration T threshold b computes the acceleration on b from the tree T according to the algorithm described above. (Hint: The function accOnPoint in lib/mechanics.sig may be useful.) You should use Plane.sum to compute the recursive calls in parallel and add the resulting accelerations!

The provided function barnes_hut uses your compute_tree and bh_acceleration functions to form the Barnes-Hut tree for a sequence of bodies and then uses that to compute the acceleration on each body in the sequence.

4.4 Testing your code

Now that you have written all of the important code, you can actually visualize an n-body simulation with some of the infrastructure that we have given you!

Barnes-Hut is more difficult to test than the previous homeworks because the data structures are more complex and examples are harder to write out by hand. We encourage you to test your code using all of the following techniques. Although you should take advantage of the visualizer and transcripts, you must also write your own tests for Tasks 4.1-4.4 in tester.sml. You will also find some utilities to help you test in tester.sml.

- Run the visualizer! The existence of many bugs will be pretty easy (and funny) to spot.
- You can use the functions provided in tester.sml. This might be especially helpful for debugging particular functions. In it you will find functions to test for equality of sequences, boxes, and much more.
- For the final task 4.5, you should test your overall implementation by running on the rational plane and diffing the output against ours, as described below. You do not need to include hard-coded tests for this one task in your SML file. Running the visualization with the output from the floating point scalars should give you a rough idea of if your code is correct, but comparing with our output on the rational scalars will be more precise.

Note that the Barnes-Hut algorithm that you implement is not the same as the naïve quadratic algorithm! The former is an approximation of the latter. Therefore, you should expect the visualization of the output of your code to be roughly the same as the visualization of the output of the naïve code, but the outputs themselves will not agree.

4.5 Compiling your code

To load your code, at the REPL issue the command

- CM.make "sources.cm";

4.6 Generating transcripts

Once you have done the CM.make, you can use the functor Transcripts in transcripts.sml to generate transcipts. A transcript is a file in which each line represents the position of all the bodies at a time step. Using Transcripts produces many example transcripts, but if you want to make your own simulations, you can write your own functor to produce transcripts for your own universes. In particular, the visualizer is a lot more fun if you simulate the solar system for a year, but we left this out because generating the rational version of the transcript takes too long.

4.6.1 Rational transcripts

You can use a rational plane transcript to test the correctness of your code.

To create rational plane transcripts, you can type the following commands into the REPL:

```
- CM.make "sources.cm";
- RatTrans.go();
```

This will create a number of files that all end in auto.txt, and may take a few minutes. You can safely delete these files at any time.

We have provided you with a number of rational transcripts in the folder tests. The transcripts that you generate in this way should match the transcripts that we have provided exactly. To compare your transcripts with ours, use the UNIX utility diff. For example,

```
diff rat.onebody.1day.auto.txt tests/rat.onebody.1day.auto.txt
```

If diff doesn't make any output, as in

```
bovik@unix34 hw10 % diff rat.onebody.1day.auto.txt tests/rat.onebody.1day.auto.txt bovik@unix34 hw10 %
```

then the files are exactly the same and your code agrees with ours on that case. If diff does produce output, as in

```
bovik@unix34 hw10 % diff rat.onebody.1day.auto.txt tests/rat.onebody.1day.auto.txt 2c2
< (0, 0),(0, 57909100000),(0, ~108208930000),(0, ~149600000000),(0, 227939100000),(0, 778547200000),(0, ~1433449370000),(0, 2876679082000),(0, ~45)
---
> (0, 0),(0, 57909100000),(0, ~108208930000),(0, ~149600000000),(0, 227939100000),(0, 778547200000),(0, ~1433449370000),(0, 2876679082000),(0, ~4503443661000)
bovik@unix34 hw10 %
```

then your code does not agree with ours on that case.⁶

We've provided a script comparetranscripts.sh that will help you do this as well. If your code is correct, the output produced by running comparetranscripts.sh should look like this:

```
bovik@unix34 hw10 % comparetranscripts.sh checking rat.onebody.1day.auto.txt against tests/rat.onebody.1day.auto.txt checking rat.onebody.1yr.auto.txt against tests/rat.onebody.1yr.auto.txt
```

⁶diff is a robust and useful UNIX utility for comparing text; you can learn more about it starting at http://en.wikipedia.org/wiki/Diff.

```
checking rat.onebody.2weeks.auto.txt against tests/rat.onebody.2weeks.auto.txt checking rat.system.1day.auto.txt against tests/rat.system.1day.auto.txt checking rat.system.2day.auto.txt against tests/rat.system.2day.auto.txt checking rat.twobody.1day.auto.txt against tests/rat.twobody.1day.auto.txt checking rat.twobody.2day.auto.txt against tests/rat.twobody.2day.auto.txt checking rat.twobody.3day.auto.txt against tests/rat.twobody.3day.auto.txt checking rat.twobody.4day.auto.txt against tests/rat.twobody.4day.auto.txt checking rat.twobody.5day.auto.txt against tests/rat.twobody.5day.auto.txt checking rat.twobody.5day.auto.txt against tests/rat.twobody.5day.auto.txt checking rat.twobody.6day.auto.txt against tests/rat.twobody.6day.auto.txt bovik@unix34 hw10 %
```

4.6.2 Real transcripts

You can use a real plane transcript to run the visualizer.

To create real plane transcripts, you can type the following commands into the REPL:

```
- CM.make "sources.cm";
- RealTrans.go();
```

As stated above, calculations with reals are much faster, and the transcipts should be created almost instantly. All the transcripts end in auto.txt and you can safely delete any of these files.

Once you have produced a transcript file, you can visualize it by navigating to

```
http://www.cs.cmu.edu/~15150/visualizer/
```

You can then load a transcript file in one of two ways: either dragging and dropping the transcript file into the dashed box, or using the file browser to select the file manually. **The visualizer will only work with real transcripts!** Once you select a transcript, click "Go!" to run the visualizer.

In the visualization, objects are displayed as circles. Note that the scale of the display will be chosen automatically so that all bodies are visible on-screen at all times. If one body is very far away from the others at some step of the simulation, this can result in objects at different positions appearing very close or even being in distinguishable in the visualization.

Note: running RealTrans.go() will generate more transcripts of greater complexity than using the rational transcripts. Try putting real.system.1001day.auto.txt into the visualizer for interesting results.