Optimizing smallpt
Daves Scies, Sideharth Bhu
Hudel Hobers
November 4th, 2020

# Optimizing smallpt

Davean Scies, Siddharth Bhat

Haskell Exchange

November 4th, 2020

Optimizing smallpt

What is smallpt anyway?

1. S

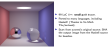
- 2. 99 LoC C++: small path tracer.
- 3. Ported to many languages, including Haskell!
- 4. Haskell port was by Vo Minh Thu. Thanks a ton!
- 5. Start from noteed's original source; SHA the output image from the Haskell source for baseline.
- 6. Perfect for an optimization case study.
- 7. Plan: Quick walk through Haskell code, end up at C++ (clang++) performance.



- ▶ 99 LoC C++: small path tracer.
- Ported to many languages, including Haskell! (Thanks to Vo Minh Thu/noteed).
- Start from noteed's original source; SHA the output image from the Haskell source for baseline.



-What is smallpt anyway?



- 1. S
- 2. 99 LoC C++: small path tracer.
- 3. Ported to many languages, including Haskell!
- 4. Haskell port was by Vo Minh Thu. Thanks a ton!
- Start from noteed's original source; SHA the output image from the Haskell source for baseline.
- 6. Perfect for an optimization case study.
- 7. Plan: Quick walk through Haskell code, end up at C++ (clang++) performance.

```
struct Vec {
 double x, y, z; // position, also color (r,q,b)
  ... methods...
struct Ray { Vec o, d; Ray(Vec o_, Vec d_) : o(o_), d(d_) {} };
enum Refl_t { DIFF, SPEC, REFR }; // material types, used in radiance()
struct Sphere {
 double rad; // radius
 Vec p, e, c; // position, emission, color
 Refl_t refl: // reflection type (DIFFuse, SPECular, REFRactive)
  ... methods ...
 double intersect(const Ray &r) const // returns distance, 0 if nohit
Sphere spheres[] = {//Scene: radius, position, emission, color, material
 Sphere(1e5, Vec( 1e5+1,40.8,81.6), Vec(), Vec(.75,.25,.25), DIFF), //Left
  ... initialization ...
};
```

#### Optimizing smallpt

-11-05

4 D > 4 A > 4 B > 4 B > B 9 Q (>

What is smallpt anyway?

where the  $\xi$   $= \xi_1$   $= \xi_2$   $= \xi_3$   $= \xi_3$ 

- 1. S
- 2. Has geometric primitives: vectors, spheres, materials
- 3. Entirely number-based, no real data structure

```
Vec radiance(const Ray &r, int depth, unsigned short *Xi){
```

```
4□ > 4률 > 4를 > 4를 > 를 ∽Q
```

Optimizing smallpt

What is smallpt anyway?

- 1. **S**
- 2. Most of the compute cost is spent in the function that traces rays.
- 3. is called radiance

radiance

radiance

```
Vec radiance(const Ray &r, int depth, unsigned short *Xi){
```

```
radiance
radiance
```

radiance

radiance

```
Optimizing smallpt

What is smallpt anyway?

What is smallpt anyway?
```

- 1. **S**
- 2. Recursively calls itself a bunch of times

```
Vec radiance(const Ray &r, int depth, unsigned short *Xi){
```

```
if (
           ) if (
                                          else
if (
                  ){
                      radiance
} else if (
                      radiance
if (
                      radiance
 radiance
                              radiance
 radiance
                              radiance
```

## Optimizing smallpt

What is smallpt anyway?

- 1. S
- 2. Recursion is guarded by a lot of control flow

```
Vec radiance(const Ray &r, int depth, unsigned short *Xi){
```

```
Vec x=r.o+r.d*t, n=(x-obj.p).norm(), nl=n.dot(r.d)<0?n:n*-1, f=obj.c;
            ) if (
if (
                                           else
if (
                  ){
                       radiance
} else if (
                       radiance
if ((cos2t=1-nnt*nnt*(1-ddn*ddn))<0)
                       radiance
  radiance
                               radiance
  radiance
                               radiance
```

#### Optimizing smallpt

-11-02



- 1. S
- 2. The control flow and computation is very numeric in nature

```
Vec radiance(const Ray &r, int depth, unsigned short *Xi){
```

```
Vec x=r.o+r.d*t, n=(x-obj.p).norm(), nl=n.dot(r.d)<0?n:n*-1, f=obj.c;
            ) if (erand48(Xi) )
if (
                                           else
if (
                   ){
                  erand48(Xi)
                                  erand48(Xi)
                       radiance
} else if (
                       radiance
if ((cos2t=1-nnt*nnt*(1-ddn*ddn))<0)
                       radiance
                                erand48(Xi)
  radiance
                               radiance
  radiance
                               radiance
```

#### Optimizing smallpt

60-TT-

What is smallpt anyway?



- 1. **S**
- 2. We use erand48 for randomness

```
Vec radiance(const Ray &r. int depth, unsigned short *Xi){
 double t;
                                         // distance to intersection
 int id=0:
                                         // id of intersected object
 if (!intersect(r, t, id)) return Vec(); // if miss, return black
  const Sphere &obj = spheres[id];
                                         // the hit object
 Vec x=r.o+r.d*t, n=(x-obj.p).norm(), nl=n.dot(r.d)<0?n:n*-1, f=obj.c;
 double p = f.x > f.y && f.x > f.z ? f.x : f.y > f.z ? f.y : f.z; // max refl
 if (++depth>5) if (erand48(Xi)<p) f=f*(1/p); else return obj.e; //R.R.
 if (obj.refl == DIFF){
                                         // Ideal DIFFUSE reflection
   double r1=2*M PI*erand48(Xi), r2=erand48(Xi), r2s=sgrt(r2);
   Vec w=n1, u=((fabs(w.x)>.1?Vec(0,1):Vec(1))\%w).norm(), v=w\%u;
   Vec d = (u*cos(r1)*r2s + v*sin(r1)*r2s + w*sart(1-r2)).norm():
   return obj.e + f.mult(radiance(Ray(x,d),depth,Xi));
 } else if (obi.refl == SPEC)
                                         // Ideal SPECULAR reflection
   return obj.e + f.mult(radiance(Ray(x,r.d-n*2*n.dot(r.d)),depth,Xi));
  Ray reflRay(x, r.d-n*2*n.dot(r.d)); // Ideal dielectric REFRACTION
  bool into = n.dot(n1)>0;
                                         // Ray from outside going in?
 double nc=1, nt=1.5, nnt=into?nc/nt:nt/nc, ddn=r.d.dot(nl), cos2t;
  if ((cos2t=1-nnt*nnt*(1-ddn*ddn))<0) // Total internal reflection
   return obj.e + f.mult(radiance(reflRay,depth,Xi));
  Vec tdir = (r.d*nnt - n*((into?1:-1)*(ddn*nnt+sqrt(cos2t)))).norm();
 double a=nt-nc, b=nt+nc, R0=a*a/(b*b), c = 1-(into?-ddn:tdir.dot(n));
 double Re=RO+(1-RO)*c*c*c*c*c.Tr=1-Re.P=.25+.5*Re.RP=Re/P.TP=Tr/(1-P);
  return obj.e + f.mult(depth>2 ? (erand48(Xi) < P ? // Russian roulette
   radiance(reflRav.depth.Xi)*RP:radiance(Rav(x.tdir).depth.Xi)*TP) :
   radiance(reflRay,depth,Xi)*Re+radiance(Ray(x,tdir),depth,Xi)*Tr);
```



#### Optimizing smallpt

0-11-05

—What is smallpt anyway?

#### What is smallpt anyway?

We consider the property of t

- 1. S
- 2. The full code continues to be more of the same

# Initial Haskell Code: radiance $(1\times)$

```
radiance :: Ray -> CInt -> Ptr CUShort -> IO Vec
radiance ray@(Ray o d) depth xi = case intersects ray of
(Nothing,__) -> return zerov
(Just t,Sphere _r p e c refl) -> do
```

```
continue f = case refl of - BRANCHING
DIFF -> do
    r1 <- ((2*pi)*) `fmap` erand48 xi -- RMG</pre>
```

#### radiance

```
SPEC -> do
  rad <- radiance -- RECURSION
REFR -> do
```

```
if
then do
rad <- radiance
```

## Optimizing smallpt

2020-11-05

Initial Haskell Code: radiance  $(1\times)$ 



- 1. S
- 2. the original source has the same computation in haskell

## Initial Haskell Code: Data structures

```
cross :: Vec -> Vec -> Vec
(.*) :: Vec -> Double -> Vec
infixl 7 .*
len :: Vec -> Double
norm :: Vec -> Vec
norm v = v .* recip (len v)
dot :: Vec -> Vec -> Double
maxv :: Vec -> Double

data Ray = Ray Vec Vec -- origin, direction
data Refl = DIFF | SPEC | REFR -- material types, used in radiance
-- / radius, position, emission, color, reflection
data Sphere = Sphere Double Vec Vec Refl
```

data Vec = Vec {-# UNPACK #-} !Double {-# UNPACK #-} !Double {-# UNPACK #-} !Double

#### Optimizing smallpt

11-05

Initial Haskell Code: Data structures

name of the of the of the (1) of the (1) of the of the the o

Initial Haskell Code: Data structures

- 1. S
- 2. We implement the same data structures in Haskell
- 3. Note that Ray, Sphere not having unpack

## Initial Haskell Code: scene data

```
spheres :: [Sphere]
spheres =
 [ Sphere 1e5 (Vec (1e5+1) 40.8 81.6) 0 (Vec 0.75 0.25 0.25) DIFF --Left
 , Sphere 1e5 (Vec (99-1e5) 40.8 81.6) 0 (Vec 0.25 0.25 0.75) DIFF --Rght
 , Sphere 1e5 (Vec 50 40.8 1e5)
                                      0 0.75 DIFF --Back
 , Sphere 1e5 (Vec 50 40.8 (170-1e5)) 0 0
                                             DIFF --Frnt
 , Sphere 1e5 (Vec 50 1e5 81.6)
                                      0 0.75 DIFF --Botm
 , Sphere 1e5 (Vec 50 (81.6-1e5) 81.6) 0 0.75 DIFF -- Top
 , Sphere 16.5 (Vec 27 16.5 47)
                                0 0.999 SPEC --Mirr
 , Sphere 16.5 (Vec 73 16.5 78)
                                0 0.999 REFR --Glas
 , Sphere 600 (Vec 50 681.33 81.6)
                                             DIFF] -- Lite
                                     12 0
```





- 1. S
- 2. this list will be walked many times, as it contains our scene information.

## Initial Haskell code: Sphere intersection

```
intersect :: Ray -> Sphere -> Maybe Double
intersect (Ray o d) (Sphere r p _e _c _refl) =
 if det<0 then Nothing else f (b-sdet) (b+sdet)
 where op = p - o -- Numeric
        eps = 1e-4
        b = dot op d
        det = b*b - dot op op + r*r -- Numeric
        sdet = sqrt det
       f a s = if a>eps then Just a else if s>eps then Just s else Nothing
intersects :: Ray -> (Maybe Double, Sphere)
intersects ray = (k, s)
 where (k,s) = foldl' f (Nothing, undefined) spheres -- Spheres iterated over
        f(k',sp) s' = case(k',intersect ray s') of
                  (Nothing, Just x) -> (Just x,s')
                  (Just y, Just x) \mid x < y \rightarrow (Just x,s')
                  _ -> (k',sp)
```



## Optimizing smallpt

20-11-05

Initial Haskell code: Sphere intersection

-> Sphere -> Maybe Double 4) (Sphere F p \_e \_c \_pef1) -

interest. (May a d) (Sphere r p .e. c. \_nexi) —
If detCO them Subling also r (G-adet) (B-adet)
Union op - p - a - - Fanoria
op - it - d

act - bet - d

adet - bet - dut op op - r v - Fanoria
adet - agt det

and - age one per the last a close if a sep then last a close little per the last a close if a sep then last a close little per - (height boddle, piners) interested any - ( $x_1, x_2$ ) interested any - ( $x_1, x_2$ ) interested per - ( $x_1, x_2$ )

Initial Haskell code: Sphere intersection

- 1. S
- 2. Responsible for figuring out what the ray hits.
- 3. We iterate over the list of spheres.
- 4. Once again, numeric heavy.
- 5. Use a Maybe to indicate whether we've found an answer or not.

# Initial Haskell Code: radiance $(1\times)$

```
radiance :: Ray -> CInt -> Ptr CUShort -> IO Vec
radiance ray@(Ray o d) depth xi = case intersects ray of
 (Nothing,_) -> return zerov
 (Just t,Sphere _r p e c refl) -> do
   let x = o 'addy' (d 'mulvs' t)
       n = norm $ x 'subv' p
       nl = if n 'dot' d < 0 then n else n 'mulvs' (-1)
       pr = maxv c
       depth' = depth + 1
       continue f = case refl of
         DIFF -> do
           r1 <- ((2*pi)*) 'fmap' erand48 xi
           r2 <- erand48 xi
           let r2s = sqrt r2
               w@(Vec wx _ _) = nl
               u = norm $ (if abs wx > 0.1 then (Vec 0 1 0) else (Vec 1 0 0)) `cross` w
               v = w `cross` u
               d' = norm $ (u`mulvs`(cos r1*r2s)) `addv` (v`mulvs`(sin r1*r2s)) `addv` (w`mulvs`sqrt (1-r2))
           rad <- radiance (Ray x d') depth' xi
           return $ e 'addv' (f 'mulv' rad)
         SPEC -> do
           let d' = d `subv` (n `mulvs` (2 * (n'dot'd)))
           rad <- radiance (Ray x d') depth' xi
           return $ e 'addy' (f 'mulv' rad)
         REFR -> do
           let reflRay = Ray x (d `subv` (n `mulvs` (2* n`dot`d)))
               into = n'dot'n1 > 0
               nc = 1
               nnt = if into then nc/nt else nt/nc
               ddn= d'dot'nl
               cos2t = 1-nnt*nnt*(1-ddn*ddn)
           if cos2t<0
             then do
               rad <- radiance reflRay depth' xi
```



#### Optimizing smallpt

2020-11-05

Initial Haskell Code: radiance (1×)



- 1. S
- 2. Branch heavy
- 3. Recursive
- 4. Uses an RNG

# Initial Haskell Code: Entry point $(1 \times)$

```
smallpt :: Int -> Int -> Int -> IO ()
smallpt w h nsamps = do
 c <- VM.replicate (w * h) 0
 allocaArray 3 \xi -> -- Create mutable memory
    flip mapM_ [0..h-1] $ \y -> do -- Loop
     writeXi xi v
     for_[0..w-1] \ x \rightarrow do -- Loop
        let i = (h-y-1) * w + x
        for_ [0..1] \sy -> do -- Loop
          for_ [0..1] \sx -> do -- Loop
            r <- newIORef 0 -- Create mutable memory
            for_ [0..samps-1] \_s -> do -- Loops, Loops
              r1 <- (2*) <$> erand48 xi
              rad <- radiance (Ray (org+d.*140) (norm d)) 0 xi -- Crunch
              modifyIORef r (+ rad .* recip (fromIntegral samps)) -- Write
            ci <- VM.unsafeRead c i
            Vec rr rg rb <- readIORef r</pre>
            VM.unsafeWrite c i $
                ci + Vec (clamp rr) (clamp rg) (clamp rb) .* 0.25 -- Write
            . . .
```

#### Optimizing smallpt

0-11-05

4 D > 4 A > 4 B > 4 B > B 9 Q (>

Initial Haskell Code: Entry point  $(1\times)$ 

ci + Yec (class sw) (class sw) (class sh) .\* 0.25 -- Svite

Initial Haskell Code: Entry point (1x)

- 1. S
- 2. Uses mutability
- 3. Performs number crunchy loops
- 4. Finally, writes results out

# Initial Haskell Code: File I/O $(1\times)$

```
withFile "image.ppm" WriteMode $ \hdl -> do
    hPrintf hdl "P3\n%d %d\n%d\n" w h (255::Int)
    flip mapM_ [0..w*h-1] \i -> do
        Vec r g b <- VM.unsafeRead c i
        hPrintf hdl "%d %d %d " (toInt r) (toInt g) (toInt b)</pre>
```

```
Initial Haskell Code: RNG (1\times)
```

```
foreign import ccall unsafe "erand48"
  erand48 :: Ptr CUShort -> IO Double
```





- 1. S
- 2. We use the RNG to decide randomly in which direction to send rays
- 3. Point out the use of foreign CCall.

Restrict export list to 'main'  $(1.13\times)$ 

-module Main where

+module Main (main) where



- 1. S
- 2. The very first thing to do is to let the compiler actually optimize.
- 3. Compiler can't know how exported functions are used.
- 4. Export lists not just about encapsulation.

# Restrict export list to 'main' $(1.13\times)$

```
-module Main where
```

+module Main (main) where

- Exported functions could be used by something unknown.
- Original versions must be available.





Restrict export list to 'main' (1.13×)

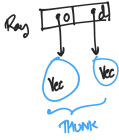
- - 1. S 2. The very first thing to do is to let the compiler actually optimize.
- 3. Compiler can't know how exported functions are used.

Restrict export list to 'main'  $(1.13\times)$ 

4. Export lists not just about encapsulation.

# Mark entries of Ray and Sphere as UNPACK and Strict $(1.07 \times)$

data Ray = Ray Vec Vec



- ▶ By default, all fields are *thunks* to rest of computation
- ► Pure, allow equational reasoning.

Optimizing smallpt

 $^-$ Mark entries of Ray and Sphere as UNPACK and Strict (1.07×)

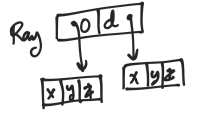
PLANT

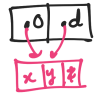
Mark entries of Ray and Sphere as UNPACK and Strict (1.07×)

By default, all fields are thunks to rest of computation
 Pure, allow equational reasoning.

# Mark entries of Ray and Sphere as UNPACK and Strict $(1.07 \times)$

data Ray = Ray !Vec !Vec





- ▶ When strict, elements are *pointers* to known structures
- pointers enable sharing!

Optimizing smallpt

Mark entries of Ray and Sphere as UNPACK and Strict  $(1.07\times)$ 

Mark entries of Ray and Sphere as UNPACK and Strict (1.07 $\times$ )



data Ray - Ray (Sec (Sec



► When strict, elements are pointers to known structures
► pointers enable sharing!



- ▶ When unpacked, elements are *members* of the parent.
- Larger, but eliminate pointer chasing.



# Mark entries of Ray and Sphere as UNPACK and Strict $(1.07 \times)$

```
data Vec = Vec {-# UNPACK #-} !Double
              {-# UNPACK #-} !Double
              {-# UNPACK #-} !Double
-data Ray = Ray Vec Vec -- origin, direction
+data Ray = Ray !Vec !Vec -- origin, direction
data Refl = DIFF | SPEC | REFR -- material types, used in radiance
-- radius, position, emission, color, reflection
-data Sphere = Sphere Double Vec Vec !Refl
+data Sphere = Sphere {-# UNPACK #-} !Double
                     {-# UNPACK #-} !Vec
                     {-# UNPACK #-} !Vec
                     {-# UNPACK #-} !Vec !Refl
struct Vec { double x, y, z; }
struct Ray { std::function<Vec()> v; std::function<Vec()> w; };
struct RayUnpack { double xv, yv, int zv;
                  double xw, yw, zw; };
```



#### Optimizing smallpt

 $^-$ Mark entries of Ray and Sphere as UNPACK and Strict (1.07imes)

American Section 2 of Section 2

Mark entries of Ray and Sphere as UNPACK and Strict (1.07×)

- 1. D
- Strictness in the arguments means that they're evaluated when instantiated, not when demanded.
- 3. Unpacking removes indirection from doing a memory lookup for components.
- 4. Means we have to copy everything into the data structure that it is unpacked into.
- 5. We don't unpack ray (Lots of calculations on its components, want those to fuse)
- 6. Unpack Sphere its static

## Use a pattern synonym to unpack Refl in Sphere $(1.07\times)$



## Optimizing smallpt

20-11-0

Use a pattern synonym to unpack Refl in Sphere (1.07 imes)



Use a pattern synonym to unpack Refl in Sphere (1.07×)

- D
- 2. UnboxedSums are recent
- 3. UnboxedSums are very unpleasant
- 4. We're using an older trick to fake the unboxing here instead.
- 5. In this case it isn't much of a win, but it illustrates the technique.

# Change from maximum on a list to max $(1.08\times)$

```
-maxv (Vec a b c) = maximum [a,b,c]
+\max v (Vec a b c) = \max a (\max b c)
    let x = o `addv` (d `mulvs` t)
         n = norm $ x `subv` p
         nl = if n `dot` d < 0 then n else n `mulvs` (-1)</pre>
         pr = maxv c
         depth' = depth + 1
         continue f = case refl of
           DIFF -> do
    if depth'>5
       then do
         er <- erand48 xi
        let !pr = maxv c
```

## Optimizing smallpt

0-11-05

↓□ → ↓□ → ↓ = → ↓ = → ∫ へ ○

Change from maximum on a list to max  $(1.08 \times)$ 

```
are (New a h c) = maximum (p,h,c)
are (New a h c) = max (num h c)
are = "simble" (num h c)
are = num f c (num h c)
are = num f
```

Change from maximum on a list to max (1.08×)

- 1. D
- 2. finicky optimization.
- 3. Minimal usage of this function.
- 4. GHC does not evaluate at compile time, only has RULES

## Convert erand48 to pure Haskell $(1.09\times)$

```
-foreign import ccall unsafe "erand48"
- erand48 :: Ptr CUShort -> IO Double
+erand48 :: IORef Word64 -> IO Double
+erand48 !t = do -- | Some number crunchy thing.
+ r <- readIORef t
+ 1et. x' = 0x5deece66d * r + 0xb
       d_word = 0x3ff000000000000 . | . ((x' .&. 0xfffffffffff) `unsafeShiftL` 4)
       d = castWord64ToDouble d word - 1.0
+ writeTORef t x'
+ pure d
-radiance :: Ray -> CInt -> Ptr CUShort -> IO Vec
+radiance :: Ray -> Int -> IORef Word64 -> IO Vec -- IORef with state
 radiance ray@(Ray o d) depth xi = case intersects ray of
   c <- VM.replicate (w * h) zerov
- allocaArrav 3 $ \xi -> -- Old RNG state
       flip mapM_ [0..h-1] $ \y -> do
+ xi <- newIORef O -- New RNG state
+ flip mapM_ [0..h-1] $ \y -> do
       writeXi xi y
```



# Optimizing smallpt Convert erand48 to pure Haskell (1.09×) Convert erand48 to pure Haskell (1.09×)

- 1. S
- 2. The entire premise of this talk is that Haskell can be as fast as C.
- 3. We're opening the black box of what erand48 does to GHC
- Further any impedance mismatch, such as FFI almost universally has to have, carries some bookkeeping overhead.
- 5. If our Haskell code was as fast as the C code moving the code into Haskell would be a win, if it was slightly slower it could still be a win.
- 6. Often considering your Haskell code's performance is a better option and easier than reimplementing something in C.
- 7. As is the way with optimizations, this is not universally true.

## Remove mutability: Erand48 Monad

```
-erand48 :: IORef Word64 -> IO Double
-erand48 !t = do
- r <- readIORef t
+data ET a = ET !Word64 !a deriving Functor
+newtype Erand48 a = Erand48 { runErand48' :: Word64 -> ET a } deriving Functor
+instance Applicative Erand48 where
+instance Monad Erand48 where
+runWithErand48 :: Int -> Erand48 a -> a
+erand48 :: Erand48 Double
-radiance :: Ray -> Int -> IORef Word64 -> IO Vec
-radiance ray@(Ray o d) depth xi = case intersects ray of
+radiance :: Rav -> Int -> Erand48 Vec
+radiance ray@(Ray o d) depth = case intersects ray of
            r1 <- (2*pi*) <$> erand48 xi
            r2 <- erand48 xi
             r1 <- (2*pi*) <$> erand48
            r2 <- erand48
                              then (.* rp) <$> radiance reflRay depth' xi
                              else (.* tp) <$> radiance (Ray x tdir) depth' xi
                              then (.* rp) <$> radiance reflRav depth'
                               else (.* tp) <$> radiance (Ray x tdir) depth'
```



## Optimizing smallpt

0-11-05

Remove mutability: Erand48 Monad



- 1. D
- 2. All these mutability locations throw in extra RTS code, extra sequencing that blocks the compiler's optimization, and dependency chains.
- 3. Sometimes we need mutability for performance.
- 4. SSA is normal to compilers though.
- 5. We almost start at SSA as a functional language.
- 6. don't break it when you don't have a good reason.

## Removing mutation: eliminate IORef and Data. Vector. Mutable

```
- c <- VM.replicate (w * h) 0
- xi <- newIORef 0
- flip mapM_ [0..h-1] $ \y -> do
       writeXi xi y
       for_{0..w-1} \ x \rightarrow do
         let i = (h-v-1) * w + x
         for_{0..1} \ v \rightarrow do
          for_ [0..1] \sx -> do
             r <- newIORef 0
             for [0..samps-1] \setminus s \rightarrow do
               r1 <- (2*) <$> erand48 xi
       img = (`concatMap` [(h-1), (h-2)..0]) $\y -> runWithErand48 y do
         for [0..w-1] \ x \rightarrow do
           (\protect\) foldlM pf 0 [(sy, sx) | sy <- [0,1], sx <- [0,1]]) \ci (sy, sx) -> do
             Vec rr rg rb <- (\f -> foldlM f 0 [0..samps-1]) \ !r _s -> do
               r1 <- (2*) <$> erand48
               modifyIORef r (+ rad .* recip (fromIntegral samps))
             ci <- VM.unsafeRead c i
             Vec rr rg rb <- readIORef r
             VM.unsafeWrite c i $ ci + Vec (clamp rr) (clamp rg) (clamp rb) .* 0.25
               pure (r + rad .* recip (fromIntegral samps))
             pure (ci + Vec (clamp rr) (clamp rg) (clamp rb) .* 0.25)
```

```
Removing mutation: eliminate IORef and Data . Vector . Mutable
Optimizing smallpt
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        \begin{aligned} & \text{for}_{i} = \{ x_{i}, x_{i} \in \mathcal{Y}_{i} \mid x_{i} = x \\ & \text{for}_{i} = \{ x_{i}, x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i}, x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i}, x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i}, x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i}, x_{i} \neq x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i}, x_{i} \neq x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i}, x_{i} \neq x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i}, x_{i} \neq x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i}, x_{i} \neq x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i}, x_{i} \neq x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i}, x_{i} \neq x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i}, x_{i} \neq x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i}, x_{i} \neq x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i}, x_{i} \neq x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i}, x_{i} \neq x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i}, x_{i} \neq x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i}, x_{i} \neq x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i}, x_{i} \neq x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i} \neq x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i} \neq x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i} \neq x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i} \neq x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i} \neq x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i} \neq x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i} \neq x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i} \neq x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i} \neq x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i} \neq x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i} \neq x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i} \neq x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i} \neq x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i} \neq x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i} \neq x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i} \neq x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i} \neq x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i} \neq x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i} \neq x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i} \neq x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i} \neq x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} = \{ x_{i} \neq x_{i} \} \mid x_{i} > x \\ & \text{for}_{i} 
                                                                                                                                                                                                                                                                                     Removing mutation: eliminate IORef and
                                                                                                                                                                                                                                                                                 Data Vector Mutable
```

1. D



4 D > 4 A > 4 B > 4 B > B 9 Q (>

# Set everything in smallpt to be strict $(1.17\times)$

```
smallpt :: Int -> Int -> Int -> IO ()
smallpt w h nsamps = do
- let samps = nsamps `div` 4
       org = Vec 50 52 295.6
      dir = norm \$ Vec 0 (-0.042612) (-1)
       cx = Vec (fromIntegral w * 0.5135 / fromIntegral h) 0 0
       cy = norm (cx `cross` dir) `mulvs` 0.5135
+ let !samps = nsamps `div` 4
       !org = Vec 50 52 295.6
       !dir = norm \$ Vec 0 (-0.042612) (-1)
       !cx = Vec (fromIntegral w * 0.5135 / fromIntegral h) 0 0
       !cv = norm (cx `cross` dir) `mulvs` 0.5135
- r1 <- (2*) `fmap` erand48 xi
- let dx = if r1<1 then sqrt r1-1 else 1-sqrt(2-r1)
- r2 <- (2*) `fmap` erand48 xi
- let dy = if r2<1 then sqrt r2-1 else 1-sqrt(2-r2)
      d = ...
- rad <- radiance (Ray (org`addv`(d`mulvs`140)) (norm d)) 0 xi
+ !r1 <- (2*) `fmap` erand48 xi
+ let !dx = if r1<1 then sqrt r1-1 else 1-sqrt(2-r1)
+ !r2 <- (2*) `fmap` erand48 xi
+ let !dy = if r2<1 then sqrt r2-1 else 1-sqrt(2-r2)
       !d = ...
              pure $! r + rad .* recip (fromIntegral samps)
             pure $! ci + Vec (clamp rr) (clamp rg) (clamp rb) .* 0.25
```

#### Optimizing smallpt

-11-05

◆ロ → ← 同 → ← 三 → へ ○ へ ○

Set everything in smallpt to be strict  $(1.17 \times)$ 



Don't senselessly bang everything in sight.

- 1.
- 2. This is not a recommendation, this is a warning.
- 3. We get a speedup here but it can also regress performance. Some of these bangs are regressions that are hidden.

## Why strictness may be bad

```
let foo = let x = error "ERR" in \y -> y let foo0pt = \y -> y
```





- 1. S
- 2. Consider the function foo and fooOpt. These are equivalent
- 3. The fact that x is not used allows us to eliminiate computing x
- 4. Consider the next version
- 5. Illegal, we need to have x, because it doesn't produce ERR
- 6. we can't equationally reason about the program anymore.
- 7. Makes it harder for GHC. GHC is conservative about bangs
- ${\it 8. \ } In hibits\ compiler\ from\ optimizing$

## Why strictness may be bad

```
let foo = let x = error "ERR" in \y -> y
let fooOpt = \y -> y
let foo' = let !x = error "ERR" in \y -> y
```





Why strictness may be bad



Why strictness may be bad

- 1. S
- 2. Consider the function foo and fooOpt. These are equivalent
- 3. The fact that x is not used allows us to eliminiate computing x
- 4. Consider the next version
- 5. Illegal, we need to have x, because it doesn't produce ERR
- 6. we can't equationally reason about the program anymore.
- 7. Makes it harder for GHC. GHC is conservative about bangs
- 8. Inhibits compiler from optimizing

## Why strictness may be bad

```
let foo = let x = error "ERR" in \y -> y
let fooOpt = \y -> y
let foo' = let !x = error "ERR" in \y -> y
let foo'Opt = \y -> y -- ERROR! forcing foo'=foo'Opt should give "ERR"
```





Why strictness may be bad

Why strictness may be bad

- 1. S
- 2. Consider the function foo and fooOpt. These are equivalent
- 3. The fact that x is not used allows us to eliminiate computing x
- 4. Consider the next version
- 5. Illegal, we need to have x, because it doesn't produce ERR
- 6. we can't equationally reason about the program anymore.
- 7. Makes it harder for GHC. GHC is conservative about bangs
- 8. Inhibits compiler from optimizing

# Reduce to only useful strictnesses in smallpt( $1.17 \times$ )

```
- let !samps = nsamps `div` 4
       !org = Vec 50 52 295.6
       !dir = norm \$ Vec 0 (-0.042612) (-1)
       !cx = Vec (fromIntegral w * 0.5135 / fromIntegral h) 0 0
       !cv = norm (cx `cross` dir) .* 0.5135
  let samps = nsamps `div` 4
       org = Vec 50 52 295.6
       dir = norm \$ Vec 0 (-0.042612) (-1)
       cx = Vec (fromIntegral w * 0.5135 / fromIntegral h) 0 0
       cv = norm (cx `cross` dir) .* 0.5135
               !r1 <- (2*) <$> erand48
               r1 <- (2*) <$> erand48
               !r2 <- (2*) < $> erand 48
               r2 <- (2*) <$> erand48
               !rad <- radiance (Ray (org+d.*140) (norm d)) 0
               rad <- radiance (Ray (org+d.*140) (norm d)) 0
               pure $! r + rad .* recip (fromIntegral samps)
             pure $! ci + Vec (clamp rr) (clamp rg) (clamp rb) .* 0.25
               pure (r + rad .* recip (fromIntegral samps))
             pure (ci + Vec (clamp rr) (clamp rg) (clamp rb) .* 0.25)
```





trad <- radiance (Ray (orged.\*560) (norm d)) (

pure \$1 x + rad .\* recip (from[ategral samps) pure \$1 ci + Vec (clamp rV) (clamp rg) (clamp rb) .\* 0.25

- 1. D
- 2. Bangs force evaluation.
- 3. The computation might diverge.
- 4. Thus the compiler can no longer move the computation around or simplify it.
- Useless work.
- 6. A little thinking about how the variables are used or looking at core allows us to select which ones we bang selectively.

## Use strictness strategically in entire project

```
if det<0 then Nothing else f (b-sdet) (b+sdet)

where op = p - o

eps = 1e-4

b = dot op d

det = b*b - dot op op + r*r

sdet = sqrt det

f a s = if a>eps then Just a else if s>eps then Just s else Nothing

if det<0

then Nothing

else

let !eps = 1e-4

!sdet = sqrt det

!a = b-sdet

!s = b+sdet

in if a>eps then Just a else if s>eps then Just s else Nothing
```



#### Optimizing smallpt

Use strictness strategically in entire project



Use strictness strategically in entire project

- 1. D
- Sometimes (point out 'intersect') we have to rearrange the code though when we use bangs.
- 3. Bangs tell the compiler to make more efficient code, but take away the compiler's options in how to do so.
- 4. Only take away the compiler's liberties when it's using them poorly.
- 5. Becomes intuitive.

## Remove Maybe from intersect(s) $(1.32\times)$

```
Old: Use Maybe Double to represent (was-hit?:bool, hit-distance: Double)
 New: use (1/0) to represent not (was-hit?)
-intersect :: Ray -> Sphere -> Maybe Double
+intersect :: Rav -> Sphere -> Double
intersect (Ray o d) (Sphere r p _e _c _refl) =
- if det<0 then Nothing else f (b-sdet) (b+sdet)
+ if det<0 then (1/0.0) else f (b-sdet) (b+sdet)
   where op = p `subv` o
         f a s = if a>eps then Just a else if s>eps then Just s else Nothing
         f a s = if a>eps then a else if s>eps then s else (1/0.0)
-intersects :: Ray -> (Maybe Double, Sphere)
+intersects :: Ray -> (Double, Sphere)
intersects rav = (k, s)
- where (k.s) = foldl' f (Nothing, undefined) spheres
         f (k',sp) s' = case (k',intersect ray s') of
                   (Nothing, Just x) -> (Just x,s')
                   (Just y, Just x) \mid x < y \rightarrow (Just x,s')
                   _ -> (k',sp)
+ where (k,s) = foldl' f (1/0.0,undefined) spheres
         f(k', sp) s' = let !x = intersect ray s' in if <math>x < k' then (x, s') else (k', sp)
radiance :: Ray -> Int -> STRefU s Word64 -> ST s Vec
radiance ray@(Ray o d) depth xi = case intersects ray of
- (Nothing, ) -> return zerov
- (Just t,Sphere _r p e c refl) -> do
+ (t,_) \mid t == (1/0.0) -> return zerov
+ (t,Sphere _r p e c refl) -> do
```

◄□▶ ◀圖▶ ◀불▶ ◀불▶ 불 ∽9<</p>

#### Optimizing smallpt

Remove Maybe from intersect(s)  $(1.32 \times)$ 

| The region of the content of the c

Remove Maybe from intersect(s) (1.32×)

- 1. S
- 2. This is a far more performance critical version of what we saw with 'maximum' vs. 'max'.
- 3. innermost functions are of critical importance. remove Maybe which significantly reduces the boxing
- 4. Since a Ray that fails to intersect something can be said to intersect at infinity, Double already actually covers the structure at play
- 5. This also reduces allocation.

## Hand unroll the fold in intersects $(1.35\times)$

```
intersects :: Ray -> (Double, Sphere)
-intersects ray = (k, s)
- where (k,s) = foldl' f (1/0.0, undefined) spheres
+intersects ray =
+ f (... (f (f (intersect ray sphLeft, sphLeft) sphRight) ...)
+ where
    f(k', sp) s' = let !x = intersect ray s' in if x < k' then (x, s') else (k', sp)
-spheres :: [Sphere]
-spheres = let s = Sphere ; z = zerov ; (.*) = mulvs ; v = Vec in
- [ s 1e5 (v (1e5+1) 40.8 81.6) z (v 0.75 0.25 0.25) DIFF --Left
- , s 1e5 (v (-1e5+99) 40.8 81.6) z (v 0.25 0.25 0.75) DIFF --Rght
. . .
+sphLeft, sphRight, ... :: Sphere
+sphLeft = Sphere 1e5 (Vec (1e5+1) 40.8 81.6)
                                                 zerov (Vec 0.75 0.25 0.25) DIFF
+sphRight = Sphere 1e5 (Vec (-1e5+99) 40.8 81.6) zerov (Vec 0.25 0.25 0.75) DIFF
```



#### Optimizing smallpt

Hand unroll the fold in intersects  $(1.35\times)$ 

Contraction of the Contraction o

Hand unroll the fold in intersects (1.35×)

- 1. D
- 2. 'intersects' is very hot
- 3. Loop unrolling
- 4. Many compilers do this for us, and there are special versions of it like Duff's Device.
- 5. Sadly GHC doesn't
- 6. Can do variants by hand.
- 7. RULE could handle each one specificly (only exactly that one)?

## Custom datatype for intersects parameter passing

```
Old: Tuple with possibly-uenevaluated Double and Sphere
New: Reference to a guaranteed-to-be-evaluated Double and Sphere
-intersects :: Ray -> (Double, Sphere)
+data T = T !Double !Sphere
+intersects :: Rav -> T
intersects rav =
    f ( ... f (intersect ray sphLeft, sphLeft) sphRight) ... sphLite
+ f ( ... f (T (intersect ray sphLeft) sphRight) ... sphLite
  where
    f(k', sp) s' =
        let !x = intersect ray s' in if x < k' then (x, s') else (k', sp)
    f !(T k' sp) !s' =
        let !x = intersect ray s' in if x < k' then T x s' else T k' sp
radiance :: Ray -> Int -> Erand48 Vec
radiance ray@(Ray o d) depth = case intersects ray of
- (!t,_) \mid t == 1/0.0 \rightarrow return 0
- (!t,!Sphere _r p e c refl) -> do
+ (T t_{-}) | t == 1/0.0 -> return 0
+ (T t (Sphere _r p e c refl)) -> do
    let !x = o + d .* t
         !n = norm \$ x - p
         !nl = if dot n d < 0 then n else negate n
```



#### Optimizing smallpt

20-11-05

Custom datatype for intersects parameter passing

> ta = norm \$ x - p tal = if dot n d < 0 then n else negate n

- 1. D
- 2. We can optimize data passing.
- 3. Want: Data strict, but not unpacked.
- 4. Compiler knows its evaluated but no copying
- 5. A normal tuple lacks strictness information.
- 6. An unboxed tuple forces copying
- 7. Strict Tuple.
- 8. This exists in libraries of course, but we wanted to illustrate it.

## Optimize file writing

```
build-depends:
      base >= 4.12 && < 4.15
    , bytestring >= 0.11
-toInt :: Double -> Int
-toInt x = floor $ clamp x ** recip 2.2 * 255 + 0.5 
+toInt :: Double -> BB.Builder -- O(1) concatenation
+toInt x = BB.intDec (floor (clamp x ** recip 2.2 * 255 + 0.5)) <> BB.char8 ' '
. . .
  withFile "image.ppm" WriteMode $ \hdl -> do
         hPrintf hdl "P3\nd" \d\n\d\n" w h (255::Int)
        for_ img \(Vec r g b) -> do
          hPrintf hdl "%d %d %d " (toInt r) (toInt g) (toInt b)
         BB.hPutBuilder hdl $
          BB.string8 "P3\n" <> -- efficient builders for ASCII
          BB.intDec w <> BB.char8 ' ' <> BB.intDec h <> BB.char8 '\n' <>
          BB.intDec 255 <> BB.char8 '\n' <>
           (mconcat $ fmap (\( (Vec r g b) -> toInt r <> toInt g <> toInt b) img)
```



#### Optimizing smallpt

Optimize file writing



Optimize file writing

build-depends: base >= 4.12 && < 4.15

- 1. D
- 2. Strings are inefficient
- 3. 'bytestring' has some efficient writing code, so we just convert to that for a modest gain.

# Use LLVM backend $(1.87\times)$

+package smallpt-opt

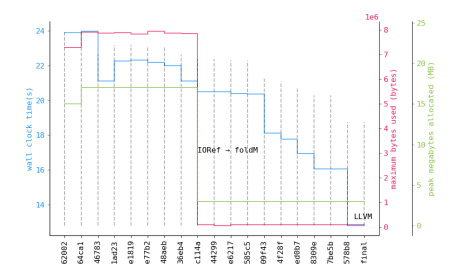
+ ghc-options: -fllvm





- 1. S
- 2. Finally, this particular code is quite numeric heavy.
- 3. There are optimizations for numeric heavy code we're missing in GHC.
- 4. LLVM has an extensive library of laws to optimize low level numeric ops.
- 5. LLVM is too low-level to understand haskell as haskell.
- 6. LLVM makes decisions with the tacit assumption that the assembly came from a C-like language, which is often to the detriment of a Haskell-like language.
- 7. In this case, as the code is "fortran-like", LLVM wins.

## The view from the mountaintop





1. D



## Avoid CPU ieee754 slow paths

```
intersect :: Ray -> Sphere -> Double
 intersect (Ray o d) (Sphere r p _e _c _refl) =
  if det<0
- then 1/0.0
+ then 1e20
   else
    . . .
    in if a>eps then a else if s>eps then s else 1/0.0
    in if a>eps then a else if s>eps then s else 1e20
radiance :: Ray -> Int -> Erand48 Vec
radiance ray@(Ray o d) depth = case intersects ray of
- (T t_) | t == 1/0.0 \rightarrow return 0
+ (T 1e20 _) -> return 0
```



#### Optimizing smallpt

-11-05

-Avoid CPU ieee754 slow paths

Avoid CPU ieee754 slow paths

intersect (i) by  $\rightarrow$  Sphere  $\rightarrow$  Double intersect (by  $\nu$  of) (Sphere  $\nu$   $\rho$   $_{\nu}$  =  $_{\nu}$  =  $_{\nu}$  = 1 than 1/0.0 • than 1/0.0 • than 1/0.0 • shar • If Subpet then a sales if support then a slar 1/0.0 • in if Subpet then a sale if given then a slar 1/0.0

vadiance :: Ray -> Int -> Eranddê Vec vadiance ray#CRay o d) depth - case intersects ray of

radiance rayb(kay o 4) depth = case intersects
- (T t \_) | t = 1/0.0 -> return 0
- (T 1600 \_) -> return 0
- ...

- 1. D
- 2. We used +Inf to match the Maybeness
- 3. C++ code set 1e20 s the horizon
- 4. Mechanical sympathy is important.
- 5. Know how the CPU (abstractly) executes slow path / fast path.

## Fix differences with C++ version

```
- if depth>2
+ if depth'>2 -- depth' = depth + 1
```





- 1. **S**
- 2. Since the sha1 of the output didn't match the C++ version we started investigating.
- 3. clang++, g++ actually produce different sha1s
- 4. unincremented depth was being used in one branch, causing us to do more work
- 5. now confident to say we're doing the same computation as C++

## **Takeaways**

- ► The unrolling in 'intersects' is ugly.
- (We feel) the maintainability of this code hasn't been significantly harmed.
- ▶ We're faster than clang++ and within 6% of g++
- ► Haven't exhausted the optimization opportunities.
- ▶ GHC could learn to do several of these optimizations for us.
- ▶ Others are just good Haskell style.
- Clean Haskell is often performant Haskell.



Takeaways

(We feel) the realistalisability of this code hasn't been significantly haveed.
 We're faster than clarg++ and within 6% of g++
 Haven't enhanced the optimization opportunities.

GHC could learn to do enveral of these optimizations for us
 Others are just good Haskell style.
 Clean Haskell is often performant Haskell.

Cital ratios is com percental ration

1. D

4□ > 4□ > 4□ > 4□ > 4□ > 4□ > 4□



# Experimental data

▶ All test were on an otherwise idle Equinix Metal c3.small.x86 (Intel Xeon E-2278G with 32GiB RAM, Ubuntu 20.04).

