Classical Zeno Effect Seen in the Coherence of Light Sources

In this notebook, we wish to repeat the results of [Soruce name blah blah]. In this paper, the authors perform an example of the Zeno effect in the classical limit, showing that use of slits to cause a distrubance on a beam of light can cause the measured intensity over a set distance to increase.

To do this, we have 4 equations we need to find:

1.
$$J(x_1,x_2)=\langle E(x_1)E^*(x_2)\rangle=rac{1}{2a}exp\Big[rac{(x_1-x_2)^2}{d^2}\Big]$$
2. $P=\int_{-a}^a J(x,x)dx$
3. $\mu_g=rac{1}{P}\cdot \left[\int\int_{-a}^a |J(x_1,x_2)|^2
ight]^{1/2}$
4. $J(x_1',x_2')=rac{k}{2\pi z}\cdot\int\int_{-a}^a J(x_1,x_2) imes K(x_1'-x_1)K^*(x_2'-x_2)dx_1dx_2$

Where:

$$K(x) = exp\Big[rac{ikx^2}{2z}\Big]$$

We solve this set of equation numerically.

Main.workspace3.Data

```
    Struct containg all variables and data arrays needed for this calculation.

# Arguments
- n::Int : Dimensionality of the system
- zmax::Float : Distance from source slit to detector

    - nr::Int : Number of slits the beam is to pass through, excluding souce and

 detector slit.

    - dd::Float - 'transverse coherence length is proportional to dd'

    mutable struct Data

     n; zmax; nr; dd;
      a; b; c; d; e;
      х;
      z; step; ss;
      function Data(n, zmax, nr, dd)
          a = complex(zeros(n,n))
          b = complex(zeros(n,n))
          c = complex(zeros(n,n))
          d = complex(zeros(n,n))
          e = complex(zeros(n,n))
```

```
x = zeros(n) * 0.0

z = zmax/nr
    #step seems to be our 'dx'
step = 2/(n-1)

#This is the only
ss = step * (1-1im)/(2*sqrt(pi * z))

return new(n,zmax,nr,dd,a,b,c,d,e,x,z,step,ss)
end
end
```

Main.workspace3.calculatePower

```
Impliments equation (2)
function calculatePower(dat::Data)
power = 0

for i in 1:dat.n-1
power += dat.a[i,i] * dat.step
end
#This currently can return a complex number. Is this correct?
return power
end
```

Main.workspace3.calculateCoherence

```
Impliments equation (3)
"""

function calculateCoherence(dat::Data, power)
    coh = 0;

for i in 1:dat.n
    for j in 1:dat.n

        coh += abs(dat.a[i,j])^2 * 2 * dat.step^2
    end
end

coh = sqrt(coh)/power
return coh;
end
```

getCentreIntensity (generic function with 1 method)

```
    function getCentreIntensity(dat::Data)
    return dat.a[convert(Int64, (dat.n+1)/2), convert(Int64, (dat.n+1)/2)]
    end
```

initExperimentValues (generic function with 1 method)

```
function initExperimentValues(dat::Data)
  for i in 1:dat.n
      dat.x[i] = -1 + 2 * (i-1)/(dat.n-1);
  end

for i in 1:dat.n
  for j in 1:dat.n

#dat.a = J(x1,x2) -> Equation 1 (a=1)
  dat.a[i,j] = exp(- (dat.x[i] - dat.x[j])^2 / dat.dd^2)/2

#Equation for K(x) and K*(x)
  #We have also included the numberical
```

iterateOverSlits (generic function with 2 methods)

```
function iterateOverSlits(dat::Data, print_all=false)
      Pretty sure this calculates over equation 4 ?
      for m in 1:dat.nr
          # println("Iteration $m our of $dat.nr")
          #print_all && println("\tStarting first Quadrant")
          #Fill the first octant of e
          for i in 1:convert(Int64, (dat.n+1)/2)
              for j in 1:i
                   dat.e[i,j] = 0
                   for k in 1:dat.n
                       dat.c[k,j] = 0
                       for l in 1:dat.n
                           dat.c[k,j] += dat.a[k,l] * dat.b[l,j]
                       dat.e[i,j] += dat.d[i,k] * dat.c[k,j]
                  end
              end
          end
          #print_all && println("\tStarting below Quandrant")
          #Fill the octant below
          for i in convert(Int64, (dat.n+3)/2):dat.n
              for j in 1:(dat.n-i+1)
                   dat.e[i,j] = 0
                   for k in 1:dat.n
                       dat.c[k,j] = 0
                       for l in 1:dat.n
                           dat.c[k,j] += dat.a[k,l] * dat.b[l,j]
                       end
                       dat.e[i,j] += dat.d[i,k]* dat.c[k,j]
                  end
              end
          end
          #print_all && println("\tStarting final Quandrants")
for i in convert(Int64, (dat.n+3)/2):dat.n
              for j in (dat.n-i+2):i
                   dat.e[i,j] = conj(dat.e[dat.n+1-j, dat.n+1-i])
              end
          end
          #print_all && println("\tFinalising data")
          for i in 1:dat.n
              for j in 1:dat.n
                   if( j<= i)
                       dat.a[i,j] = dat.e[i,j]
                       dat.a[i,j] = conj(dat.e[j,i])
                   end
              end
          end
      end
      return
 end
```

runExperiment (generic function with 1 method)

```
function runExperiment(dat::Data)
   initExperimentValues(dat)

startPower = calculatePower(dat)
startCoh = calculateCoherence(dat, startPower)
startIntensity = getCentreIntensity(dat)
iterateOverSlits(dat)

endPower = calculatePower(dat)
endCoh = calculateCoherence(dat, endPower)
endIntensity = getCentreIntensity(dat)

return startPower, startCoh, startIntensity, endPower, endCoh, endIntensity
end
```

Main.workspace3.runMultipleExperiments

```
    Runs a set of different experiments, with all varaibles constant except

the total number of slits 'nr'.
# Arguments
- 'n::Integer' : The dimensionality to solve over
 - 'zmax::Float': The distance between soruce slit and detector
 - 'dd::Float' : Not really sure
function runMultipleExperiments(n, zmax, dd, min_nr, max_nr)
     results = zeros(max_nr-min_nr + 1, 7) * 1im
      for nr in min_nr:max_nr
         dat = Data(n,zmax, nr, dd)
          sPow, sCoh, sInt, ePow, eCoh, eInt = runExperiment(dat)
 #
          println(nr - min_nr + 1)
         results[nr - min_nr + 1, 1] = nr
         results[nr - min_nr + 1, 2] = sPow;
         results[nr - min_nr + 1, 3] = sCoh;
         results[nr - min_nr + 1, 4] = sInt;
         results[nr - min_nr + 1, 5] = ePow;
         results[nr - min_nr + 1, 6] = eCoh;
         results[nr - min_nr + 1, 7] = eInt;
     end
     return results
end
```

0.1

```
begin
    #currently using a small n=51, this should be increased to 201
    n = 51
    zmax = 0.5
    dd = 0.1
end
```

"Experiment run succesfully for slit counts between 1 and 20"

```
begin
results = runMultipleExperiments(n, zmax, dd, 1, 20);
"Experiment run successfully for slit counts between 1 and 20"
# we wish z_d = 0.7/ka^2
end
```

We have generated and stored our results as a 2d array. Each row represents a different experiment, with the collomns representing (from left to right):

- 1. NR The number of slits for this experiment
- 2. Start Power (Complex?)
- 3. Start Coherence
- 4. Start Intensity
- 5. End Power (Complex)
- 6. End Coherence
- 7. End Intensity

We can now try and plot these, such that we can compare them to the original paper.

```
md"""
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```

"Plot values successfully extracted"



