proc_using_spatstat

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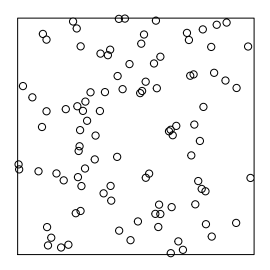
```
library(spatstat)
library(tidyverse)
library(here)

# Check to see how to use the function
?rpoispp
# Check to see how this is written
#View(rpoispp)

# Plot a homogeneous Poisson process
# p.1334 of https://mran.microsoft.com/snapshot/2016-04-25/web/packages/spatstat/spatstat.pdf
# https://spatstat.org/SSAI2017/solutions/solution04.html

par(mfrow=c(1, 1))
plot_2D_HPP <- plot(rpoispp(lambda = 100, win=square(1)), main = "HPP (rate = 100)")</pre>
```

HPP (rate = 100)



```
plot_2D_HPP
```

Symbol map with no parameters

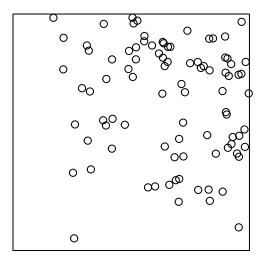
```
#plot(rpoispp(lambda = 50, win=square(1)),main = "HPP (rate = 50)")
#plot(rpoispp(lambda = 10, win=square(1)),main = "HPP (rate = 10)")

# Plot a homogeneous Poisson process
# p.33 of https://spatstat.org/resources/spatstatJSSpaper.pdf
```

```
#Imbda_function <- function(x, y) 400*x
#so that they have the same expected # of events but why ???

par(mfrow=c(1, 1))
plot_2D_NPP <- plot(rpoispp(lambda = function(x, y) 400*x*y, win=square(1)), main = "NPP (intensity = 1)</pre>
```

NPP (intensity = 100*x)

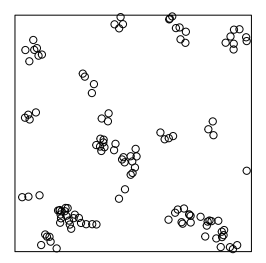


```
plot_2D_NPP
```

Symbol map with no parameters

plot_2D_Matern <- plot(rMatClust(kappa = 20, r = 0.05, mu = 5), main = "Matern (kappa = 20, r = 0.05, m

Matern (kappa = 20, r = 0.05, mu = 5)



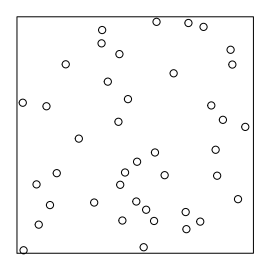
```
plot_2D_Matern
```

```
## Symbol map with constant values
## cols: #000000FA
```

```
?rMaternII
# kappa = intensity
# r = inhibition distance
```

```
# Plot a Matern I process
par(mfrow=c(1, 1))
plot(rMaternI(kappa = 100, r = 0.05), main = "Matern I (kappa = 100, r = 0.05)")
```

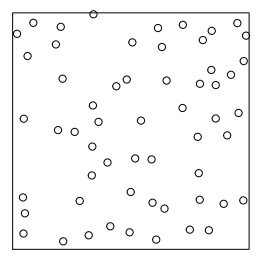
Matern I (kappa = 100, r = 0.05)



```
\#plot(rMaternI(kappa = 50, r = 0.05), main = "Matern I (kappa = 50, r = 0.05)") \\ \#plot(rMaternI(kappa = 10, r = 0.05), main = "Matern I (kappa = 10, r = 0.05)") \\
```

```
# Plot a Matern II process
par(mfrow=c(1, 1))
plot(rMaternII(kappa = 100, r = 0.05), main = "Matern II (kappa = 100, r = 0.05)")
```

Matern II (kappa = 100, r = 0.05)



```
#plot(rMaternII(kappa = 50, r = 0.05), main = "Matern II (kappa = 50, r = 0.05)")
#plot(rMaternII(kappa = 10, r = 0.05), main = "Matern II (kappa = 10, r = 0.05)")

# # Save out results
# write_rds(plot_2D_HPP, here("results", "plot_2D_HPP.jpeg"))

# # Error checking
# plot_2D_HPP <- readRDS(here("results", "plot_2D_HPP.jpeg"))
# plot_2D_HPP</pre>
```