03_proc_Hawkes

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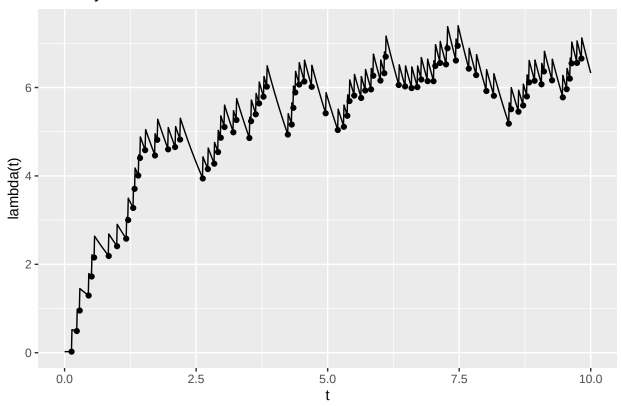
```
library(tidyverse)
library(ggplot2)
library(patchwork)
```

```
# Writing the thinning algorithm using James's way
# Initialize
\# Note that mu > 0 and 0 < alpha < beta ?? Yes.
mu = 0.025
alpha = 0.5
beta = 0.7
# Create the lambda(t) function
lmbda_fun <- function(time, obs){</pre>
 diff = time - obs
 diff = diff[diff > 0]
 a = sum(alpha * exp(-beta * diff))
 out = mu + a
  return(out)
}
# Apply the lmbda_fun function
lmbda_star <- sapply(X = t_vec, FUN = lmbda_fun, obs = t_vec)</pre>
```

```
set.seed(1)
lmbda <- median(lmbda_star) #use median instead of 10
prob_keep <- lmbda / lmbda_star</pre>
```

```
#t_keep <- t_vec[runif(length(prob_keep)) < min(prob_keep, 1)]</pre>
check <- runif(length(prob_keep)) < min(prob_keep, 1)</pre>
lmdba_keep <- lmbda_star[check]</pre>
t_keep <- t_vec[check]</pre>
# Check to see if HPP is thinned
# length(lmbda_star) #114
# length(t_vec) #114
# length(lmdba_keep) #87
# length(t_keep) #87
# Plot Hawkes
# Create a df
df_Hawkes = tibble(
 x = t_{keep}
  y = 1:(length(t_keep)), #0:(length(X) - 1)
 lmbda = lmdba_keep,
 lmbda2 = sapply(X = t_keep, FUN = lmbda_fun, obs = t_keep)
df_Hawkes2 = tibble(
 x = seq(0, 10, length.out = 1001),
 y = sapply(X = x, FUN = lmbda_fun, obs = t_keep)
)
p_Hawkes <- ggplot(data=df_Hawkes, mapping=aes(x=x, y=y)) +</pre>
  geom step() +
  labs(title = "Hawkes Process",
       x = "t",
       y = "N(t)")
#p_Hawkes
# Plot time plot
p_Hawkes_time <- ggplot(data=df_Hawkes, mapping=aes(x=x, ymin = -0.5, ymax = 0.5)) +
  geom_linerange() +
  geom_hline(aes(yintercept = 0), linetype = "dashed") +
  labs(title = "Corresponding Inter-Arrivial Times",
       x = "t"
       y = "") +
  scale_x_continuous(limits = c(0, 10), breaks = seq(0, 10, by = 1)) +
  theme(axis.text.y=element_blank(), axis.ticks.y=element_blank())
#p_Hawkes_time
# Plot lambda_star vs t
\# p\_Hawkes\_Int \leftarrow ggplot(df\_Hawkes, aes(x=x, y=lmbda2)) +
  geom_line() +
  labs(title = "Intensity Function",
#
         x = "t"
         y = "lambda(t)")
# p_Hawkes_Int
p_Hawkes_Int2 <- ggplot(df_Hawkes2, aes(x=x, y=y)) +</pre>
  geom_line() +
```

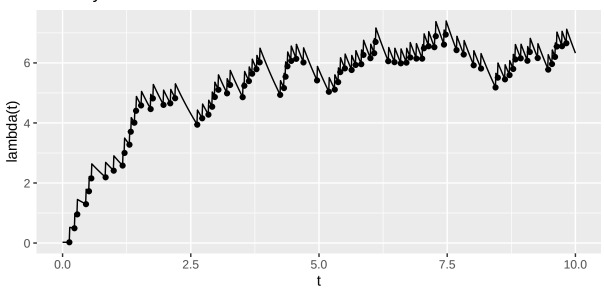
Intensity Function



```
# # Plot lambda (rate)
# p_Hawkes_hist <- ggplot(data=df_Hawkes, mapping=aes(x=x)) +
# geom_histogram(bins = 11, color = "white") +
# labs(title = "Intensity Function",
# x = "t",
# y = "lambda(t)") +
# xlim(0, 10) # so that scale lines up
# p_Hawkes_hist</pre>
```

```
# # Combine plots
# require(gridExtra)
# grid.arrange(p_Hawkes, p_Hawkes_time)
p_Hawkes_Int2 / p_Hawkes_time + plot_layout(heights = c(0.9, 0.1))
```

Intensity Function



Corresponding Inter-Arrivial Times



```
# Save output and adjust size
png(file = '/Users/franceslinyc/Hawkes-Process-2021/results/plot_1D_Hawkes.png', width = 450*2, height
p_Hawkes_Int2 / p_Hawkes_time + plot_layout(heights = c(0.9, 0.1))
```

```
# while(t <= t_max){
# if(runif(1) < min(lmbda/lmbda_star, 1)){
# X_keep <- c(X, t)
# }
# return(X_keep)
# }</pre>
```

```
# # Plot lmbda_star
# # Not thinned yet
# length(lmbda_star) #114
# x_max = length(lmbda_star) - 1
# plot(x = 0:x_max, y = lmbda_star)
# lmbda_star
```

```
# # Plot lmbda_star
# # Not thinned yet
# length(t_keep) # 75
# x_max = length(t_keep) - 1
# plot(x = 0:x_max, y = t_keep)
```