AdvML Lab3 Group Report

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Question 1: STATE SPACE MODELS - PARTICLE FILTER

The purpose of the lab is to put in practice some of the concepts covered in the lectures. To do so, you are asked to implement the particle filter for robot localization. For the particle filter algorithm, please check Section 13.3.4 of Bishop's book and/or the slides for the last lecture on state space models (SSMs). The robot moves along the horizontal axis according to the following SSM:

$$p(z_{t} \mid z_{t-1}) = \left(\mathcal{N}(z_{t} \mid z_{t-1}, 1) + \mathcal{N}(z_{t} \mid z_{t-1} + 1, 1) + \mathcal{N}(z_{t} \mid z_{t-1} + 1, 1) \right) / 3$$
 Transition model
$$p(x_{t} \mid z_{t}) = \left(\mathcal{N}(x_{t} \mid z_{t}, 1) + \mathcal{N}(x_{t} \mid z_{t} - 1, 1) + \mathcal{N}(x_{t} \mid z_{t} + 1, 1) \right) / 3$$
 Emission model
$$p(z_{1}) = Uniform(0, 100)$$
 Initial model

1.1 Simulate from a SSM. Robot Localization via Particle Filter

Implement the SSM above. Simulate it for T = 100 time steps to obtain $z_{-}(1:100)$ (i.e., states) and x1:100 (i.e., observations). Use the observations (i.e., sensor readings) to identify the state (i.e., robot location) via particle filtering. Use 100 particles. Show the particles, the expected location and the true location for the first and last time steps, as well as for two intermediate time steps of your choice.

```
library(gridExtra)
library(ggplot2)
```

```
# plot robot locations for each timestep
plot_positions = function(x_matrix, true, predicts, title){
 n = length(true)
 index = c()
 point = c()
 for(i in 1:n){
   index = c(index, rep(i, n))
   point = c(point, x_matrix[i, ])
 df plot = data.frame(index = index,
                     particle = point)
 plot = ggplot() +
   geom_point(aes(x = df_plot$index, y = df_plot$particle),
            size = 2, shape = "-", alpha=0.25, col = \#e0fff6") +
   geom_line(aes(x = 1:n, y = true, col = "True"),
            size=0.4, alpha=0.7) +
   geom_line(aes(x = 1:n, y = predicts, col = "Predicted"),
             size=0.4, alpha=0.7) +
   geom_point(aes(x = 1:n, y = true, col = "True"),
             size=0.75, alpha=0.8) +
   geom_point(aes(x = 1:n, y = predicts, col = "Predicted"),
             size=0.75, alpha=0.8) +
   scale_color_manual(values = c("Predicted" = "#efdf00",
                                   "True" = \#3c5bc1")) +
   labs(title = "Position Comparison",
        subtitle = title,
```

```
col = "Type",
         x = "Timestep", y = "Position") +
    theme dark() +
   theme(plot.background = element rect(fill = "#d0d0d0"),
          legend.background = element rect(fill = "#d0d0d0"),
          legend.position = "bottom")
 return(plot)
#The transition model: NOTE - It is not the average of three normal!
#It is a mixture model. Each distribution has the same coeficient
#That means that it is equal likely to generate from one distribution between 3
#For that reason, we sample an index between in order to know from which
#distribution we are going to generate
#After that we generate from that one
#The difference between the distr is that we add a different constant in the mean
#For that reason we sample that constant with equal probability
TransModel = function(size,mu , sd_trans, coef){
  #sample equally from 3 constants
  index = sample(x = c(0,1,2), size = size, replace = TRUE, prob = coef)
  #Sample from the normal adding each time the constant
  #In this way, we sample from the micture distribution
  smple = rnorm(n = size, mean = mu + index, sd = 1)
 return(smple)
}
#The emission Model
#Every we say above holds also for the emission model
EmissModel = function(size, mu , sd_emi,coef){
  #Here the constant we add is different though
 index = sample(x = c(0,-1,1), size = size, replace = TRUE, prob = coef)
 smple = rnorm(n = size, mean = mu + index, sd = sd_emi)
 return(smple)
#The initial model: Just a uniform distribution: No clue where the stupid robot is
InitModel = function(init,size,...){
 res = init(size,...)
 return(res)
}
#Function to simulate the SSM
#In this function
```

```
SSM_simulation = function(steps,size,InitMl,sd_trans,sd_emi,...){
  #generate from the initial model
  #Use the function IniModel in order to use it for different models
  initValues = InitModel(init = InitMl, size, ...)
  True_state = rep(0,steps)
  True_state[1] = initValues
  Observations = rep(0, steps)
  Observations[1] = EmissModel(size = 1,True_state[1],sd_emi = sd_emi,
                               coef = rep(1/3,3))
  #a for loop to generate from the transition and emission model
  for (i in 2:steps) {
    True_state[i] = TransModel(size = size,mu = True_state[i-1],
                               sd_trans = sd_trans,coef = rep(1/3,3))
    Observations[i] = EmissModel(size = size,True_state[i],sd_emi = sd_emi,
                                 coef = rep(1/3,3))
    }
  return(list(True_state,Observations))
}
#function to evaluate the particles( compute the weights)
#Here we can caclulate the densities of the three normal and average them
#We also have to normalize the weights
evalParticles = function(Observation, particls, s){
  dens = (dnorm(x = Observation, mean = particls, sd = s)+
  dnorm(x = Observation, mean = particls -1, sd = s)+
  dnorm(x = Observation, mean = particls +1, sd = s))/3
  norm_dens = dens/sum(dens)
  #return the average of the densities
  return(norm_dens)
}
#######Particle filter########
PartFIlter = function(steps, #How many steps we will do
                       partSize,#How many particles
                       initMdl, #what is my initial model
                       sd_trans, #sd of transition model
                       sd_emi, #sd of emition model
                       correction = TRUE, #True if we want to calculate weights
                       method, #which method the user wants,
                       observ,
                       ...)#... for some extra parametres
```

```
{
#method first
if(method == 1){
  #Initially generate partSize of particles
  #The first step is to generate from the initial model ZO
 particles = InitModel(init = initMdl,size = partSize,...)
 N = length(particles)
  #Store all the particles in each step to plot them
 all_particles = matrix(0,nrow = steps,ncol = partSize)
  #vector to store the expected positions
 predictions = rep(0,N)
  Weights = rep(0,N) #store the weights
 for (i in 1:(steps)) {
    #sample from transition model one step ahead using the previous particles
    #So we think If i am in this position now, where i will be in the next
    #Time step. Of course this is just my belief.
    \# I am doing that for ALL particles without using the weights
    belief = TransModel(size = N,mu = particles,sd_trans = sd_trans,
                        coef = rep(1/3,3))
    #I calculate the weights now. The weights are how likely is to have
    #an observation with that position.
    #We know Z_t (observation) and we condition to the belief we did before.
    weights = evalParticles(Observation = observ[i],
                              particls = belief,
                              s = sd_{emi}
    #If we do not want a correction, we convert all weights into 1/N
    if(correction == FALSE){
     weights = rep(1/N,N)
    #Now the Resampling starts. We have the belief, but we know that not all
    #the particles are needed, in the sense that some of them are closer to
    #the real state. Therefore we have to weight them and use the weights
    #In this step we sample from our belief but we also use the weights.
    #More specific, we sample the same number of particles BUT using the
    #weights will give us many times the same particles.
    particles = sample(x = belief, size = N, replace = TRUE, prob = weights)
    all_particles[i,] = particles
   predictions[i] = mean(particles) #make the prediction
 }
```

```
return(list(predictions,all_particles))
#second method
}else{
  #We store the particles at each step
 particles = matrix(0,nrow = steps,ncol = partSize)
  #Initially generate partSize of particles
 init_particles = InitModel(init = initMdl, size = partSize,...)
 N = ncol(particles)
  #vector to store the expected positions
 predictions = rep(0,N)
 Weights = matrix(0,nrow = steps,ncol = partSize) #store the weights
  #In this method, we first evaluate the weights, using the function evalParticles
  #We calculate the weights for each particle in every step
  #This will give us the probability of my predictions
  #Now we use the trainsition model
  #First we sample from the particles that we already have,
  #But we use the weights in order to sample the most probable
  #In the initial step, all weights are equal
 temp_particles = sample(x = init_particles, size = N, replace = TRUE,
                           prob = rep(1,N))
      #After that we sent them to the transition model in order to make a prediction
  particles[1,] = TransModel(size = N,mu = temp_particles,
                               sd_trans = sd_trans, coef = rep(1/3,3))
  #finally we update the weights
  Weights[1,] = evalParticles(Observation = observ[1],
                                particls = particles[1,],
                                s = sd_{emi}
  if(correction == FALSE){
       Weights[1,] = rep(1/N,N)
 }
  #Store the prediction
 predictions[1] = sum(Weights[1,]*particles[1,])
  #We now do the same for ALL steps, each time the same procedure
 for (i in 2:(steps)) {
    #So first we have to propagate the particles one step ahead
```

```
#We sample using the weights N particles in order to use the most likely
      temp_particles = sample(x = particles[i-1,],size = N,replace = TRUE,
                              prob = Weights[i-1,] )
      #We propagate those particles using the transition model
      particles[i,] = TransModel(size = N,mu = temp_particles,
                                  sd_trans = sd_trans, coef = rep(1/3,3))
      #Now we update the weights for each particle
      Weights[i,] = evalParticles(Observation = observ[i],
                                particls = particles[i,],
                                 s = sd_{emi}
       #If we do not need a correction, we just assign equal weights
      if(correction == FALSE){
        Weights[i,] = rep(1/N,N)
      }
      #Here our prediction is the sum of the multiplication between the
      #current weight and the particles
      predictions[i] = sum(Weights[i,]*particles[i,])
      }
 }
    #The function returns the predictions and the particles for each time step
    return(list(predictions,particles))
}
# Function to take the legend of a plot
g_legend <- function(a.gplot){</pre>
  tmp <- ggplot_gtable(ggplot_build(a.gplot))</pre>
 leg <- which(sapply(tmp$grobs, function(x) x$name) == "guide-box")</pre>
 legend <- tmp$grobs[[leg]]</pre>
 return(legend)
}
#A function to plot the particles in different times steps
#Also we plot the true and the expected location
plot_function = function(particles,true_pos,exp_pos,plots){
 plot_list = list()
 plt_N = 1
```

```
legend_taken = FALSE
  for(i in plots){
    p = eval(substitute(
      ggplot() +
        geom_histogram(mapping = aes(x = particles[i,], y = ..density..),
                       bins = 50,fill = "lightblue",color = "blue")+
        geom_density(mapping = aes(x = particles[i,]),
                     color = "yellow", size = 0.5, lty = 2)+
        geom_point(mapping = aes(x = exp_pos[i], y = 0,
                                 col = "Expected\nPosition"), size = 2.5 )+
        geom_point(mapping = aes(x = true_pos[i], y = 0,
                                 col = "True\nPosition") ,size = 2.5)+
        scale_color_manual(values = c("True\nPosition" = "red",
                                      "Expected\nPosition" = "blue"),
                           name = "Position")),list(i = i))+
      labs(subtitle = paste("Time step = ",i),x = "Position",
           y = "Density")+
      theme_bw()
    # Make a common legend for every plot
    if(!legend_taken) {
      # Take the legend
      p = p +
      theme(legend.text = element_text(margin = margin(b = 7.5, unit = "pt")))
      my_legend <- g_legend(p)</pre>
      legend_taken <- T</pre>
    }
    \# Remove the legend from the individual plot and add it to the plot list
    p = p + theme(legend.position = "none",
    plot.margin = unit(c(5.5,22,5.5,5.5), "pt"))
    #plot_list = rlist::list.append(plot_list, p)
    plot_list[[plt_N]] = p
    plt_N = plt_N +1
    }
  grid.arrange(do.call("arrangeGrob", c(plot_list, ncol=2)), my_legend,
                ncol = 2, widths = c(8, 1))
}
#Plot the residuals
plot_res = function(resid, #vector of the residuals
```

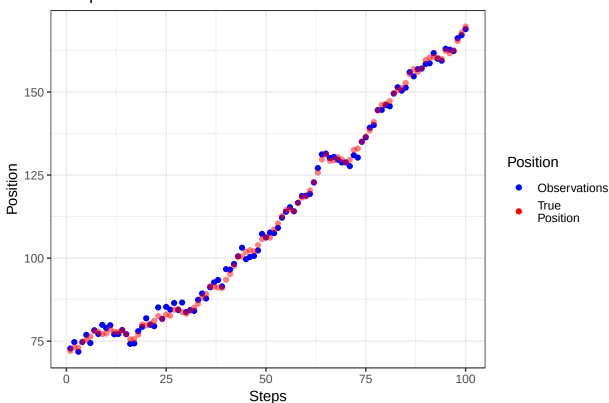
```
sd) # For what sd we run the filter in order to print it
  {
  #plot the residuals
  res_plot = ggplot()+
  \#geom\_histogram(mapping = aes(resid, y = ..density..), fill = "green", bins = 50) + (aesity..)
  geom_density(mapping = aes(resid), color= "black", alpha = 0.5,
               fill = "red", lty = 2) +
  geom_vline(xintercept = mean(resid), color = "yellow", lty = 2, size = 1)+
  labs(subtitle = paste("sd =",sd),
       x = "Residuals", y = "Density")+
 theme_light()
  return(res_plot)
}
#plot the predictions in the last step
plt_predict = function(exp_pos,true_pos, sd){
  steps = seq(1,nrow(exp_pos),1)
 plot_list = list()
 plt_N = 1
  legend_taken = FALSE
 for(i in sd){
    p = eval(substitute(
   ggplot()+
    geom_point(mapping = aes(x = steps, y = exp_pos[,plt_N],
                           col = "Expected\nPosition"), alpha = 0.5)+
    geom_line(mapping = aes(x = steps, y = true_pos,
                           col = "True\nPosition"), lty = 2)+
     scale_color_manual(values = c("True\nPosition" = "red",
                                       "Expected\nPosition" = "blue"),
                           name = "Position")),list(i = i))+
      labs(subtitle = paste("sd = ", sd[plt_N]),
           x = "Position",
           y = "Density")+
      theme bw()
     # Make a common legend for every plot
    if(!legend taken) {
      # Take the legend
     p = p +
      theme(legend.text = element_text(margin = margin(b = 7.5, unit = "pt")))
      my_legend <- g_legend(p)</pre>
      legend_taken <- T</pre>
    # Remove the legend from the individual plot and add it to the plot list
    p = p + theme(legend.position = "none",
```

```
plot.margin = unit(c(5.5,22,5.5,5.5), "pt"))
   #plot_list = rlist::list.append(plot_list, p)
   plot_list[[plt_N]] = p
   plt_N = plt_N + 1
   }
  grid.arrange(do.call("arrangeGrob", c(plot_list, ncol=2)), my_legend,
               ncol = 2, widths = c(8, 1), top = "Filtering distributions")
}
#Simulate the data
set.seed(12345) #set the seed for same results
#Simulate 100 steps
simulation_SSM = SSM_simulation(steps = 100, size = 1, InitMl = runif, 1, 1, 0, 100)
true_states = simulation_SSM[[1]] #store the true positions
Obs_states = simulation_SSM[[2]] #store the observations
#plot true positions and the observations of the robot in order to get an idea
plot_true = ggplot()+
   geom_point(mapping = aes(x = c(1:100), y = Obs_states,
                         col = "Observations"))+
   geom_point(mapping = aes(x = c(1:100), y = true_states,
                         col = "True\nPosition"), alpha = 0.5)+
    scale_color_manual(values = c("True\nPosition" = "red",
                                    "Observations" = "blue"),
                         name = "Position")+
   labs(title = "True position and observations",
                      x = "Steps",
                      v = "Position")+
     theme_bw()
#Run the filter for emission standard deviaton equal to 1
part_Filt_sd1 = PartFIlter(steps = 100,
                         partSize = 100,
                         initMdl = runif,
                         sd_trans = 1,
                         sd_{emi} = 1,
                         correction = TRUE,
                         method = 2,
                         observ = Obs_states,
                         0,100)
predict_sd1 = part_Filt_sd1[[1]] #Store the predictions
particles = part_Filt_sd1[[2]]
```

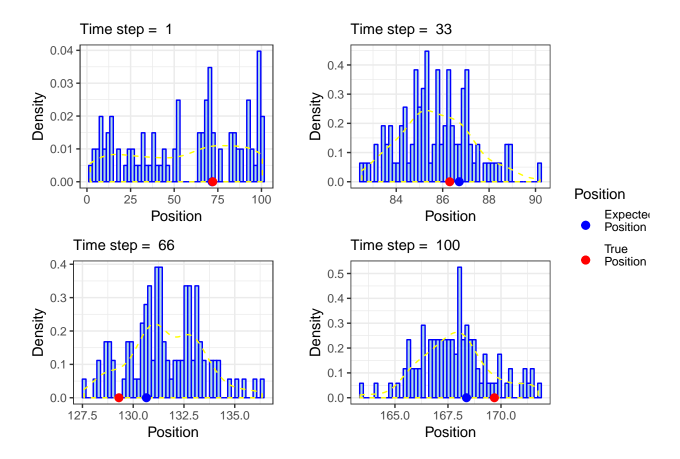
```
residuals = predict_sd1 - true_states #calculate the residuals
abs_error = mean(abs(residuals)) #calculate the mean of the resid error
```

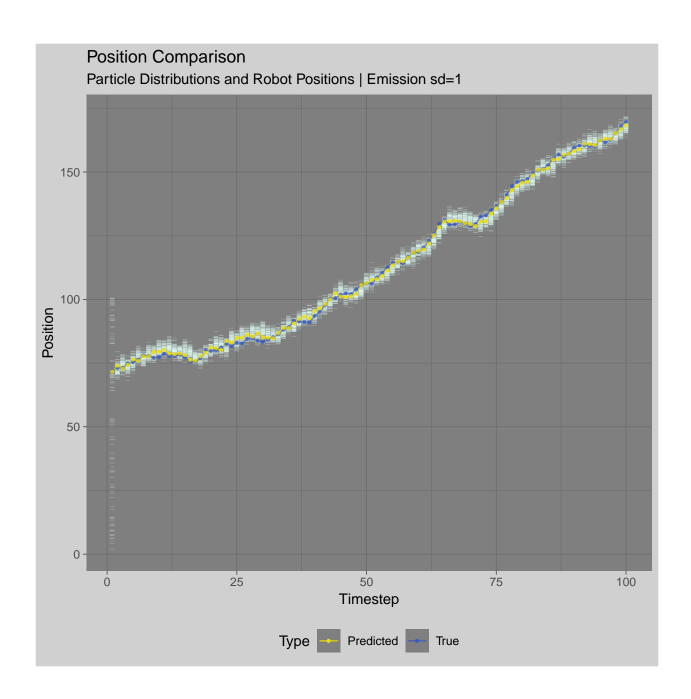
We first plot the observations and the true position of the robot. It is obvious that the sensor gives as pretty accurate results, which means that the proposal distribution(emission model) seems reasonable. Of course we could comment that from just looking to the two models(transition and emission), which are both a mixture of three Normal distributions with equal linear coefficients.

True position and observations



The plots below present the filtering distribution $P(Z_t|X_{1:t})$ for different time steps. They also show the true location and the expected location of the robot. As we can see the predictions seem both reasonable and accurate.





1.2 Repeat the procedure with different emission standard deviation

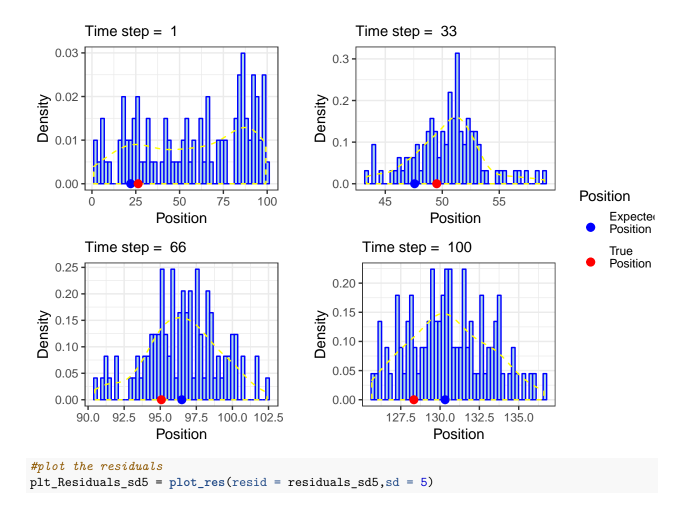
Repeat the exercise above replacing the standard deviation of the emission model with 5 and then with 50. Comment on how this affects the results.

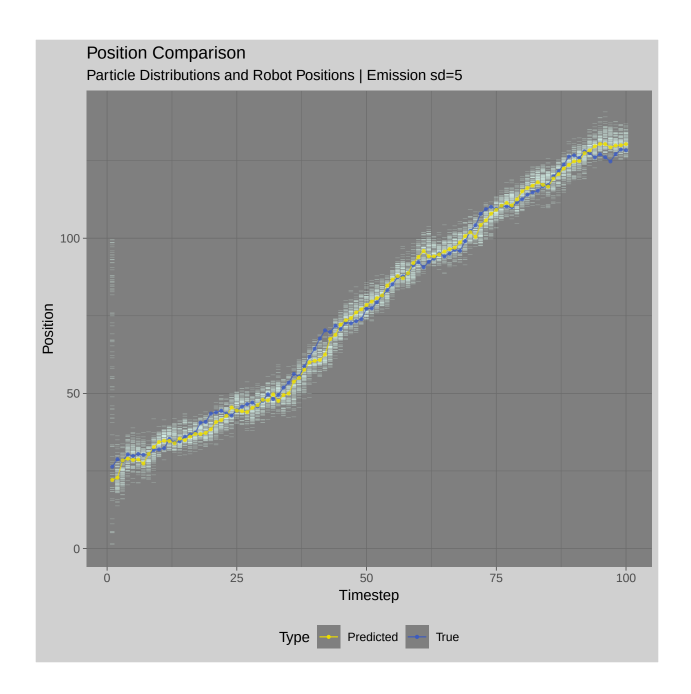
In this task we run the particle filter again but this time we change the standard deviation of emission model. We observe that increasing standard deviation of emission model causes an unaccurate predictions. This can be explained like we do not have a good sensor that gives the observations with confident. This uncertainty makes our weighted sums very smooth. We can see the particle distributions in the comparison plots and they are very wide also when we compare our predicted locations. And the true locations, we see that predicted locations cannot capture the noise in the true locations so it estimates it like a more straight line. This effect is more visible when we have bigger sd.

sd = 5

As we can see the results are slighly worse than the run with sd = 1. In this case the probability that a value outside of the target distribution will be selected is higher than before, because the proposal distribution is wider. Therefore, the particles that are close to the mean of will have a huge probability to be selected while other particles may never selected. As a result, the predictions will follow a linear form. We also plot the predictions in the task 1.3 below for each time step in order to comment on that assumption.

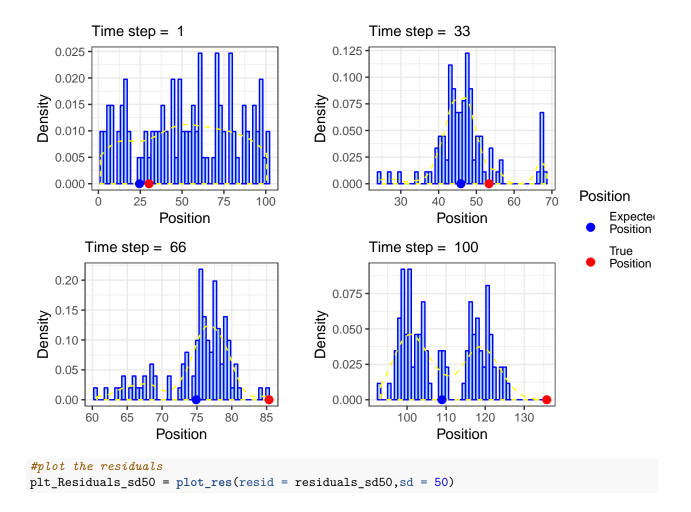
```
##### change the variance
#Simulate 100 steps
simulation SSM sd5 = SSM simulation(steps = 100, size = 1, InitMl = runif, 1, sd emi = 5,0,100)
true states sd5 = simulation SSM sd5[[1]] #store the true positions
Obs_states_sd5 = simulation_SSM_sd5[[2]]
#Here we run the particle filter with sd = 5
part_Filt_sd5 = PartFilter(steps = 100,partSize = 100,initMdl = runif,
                           sd_trans = 1,sd_emi = 5,correction = TRUE,method = 2,
                           observ = Obs_states_sd5,
                           0,100)
predict_sd5 = part_Filt_sd5[[1]] #Store the predictions
particles sd5 = part Filt sd5[[2]]
residuals_sd5 = predict_sd5 - true_states_sd5 #calculate the residuals
abs error sd5 = mean(abs(residuals sd5)) #calculate the abs error
plt_sd5 = plot_function(particles = particles_sd5,true_pos = true_states_sd5,
                        exp_pos = predict_sd5, plots = c(1,33,66,100))
```

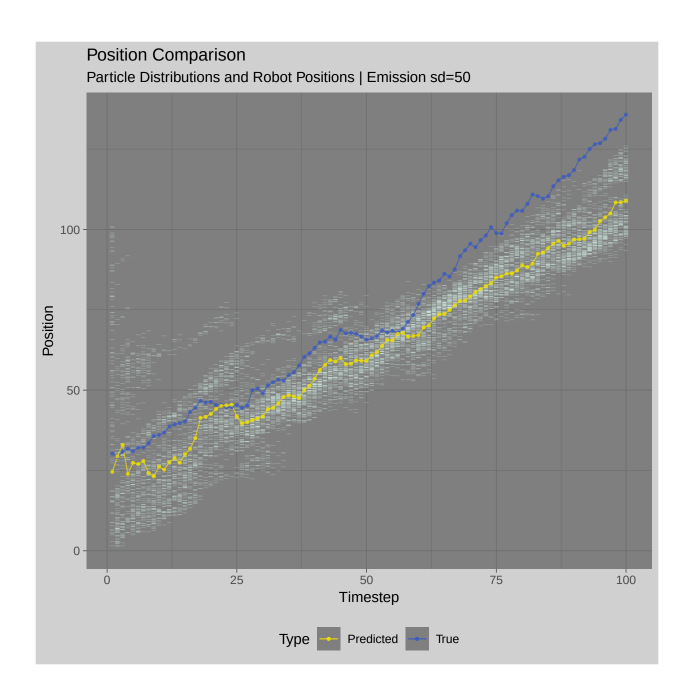




sd = 50

It is obvious that when we run the filter for sd=50, we do not have good results. The reason for that is because we generate the particles using a really huge range. That means that we do not have much information for the relationship between the obserations and the true position. Therefore, the sensor is not giving valiable information about the position of the robot. One solution to that could be to increase the number of particles. By the law of large numbers we know that, while $N \to \infty$, the expected value of the estimator will converge to the real value.

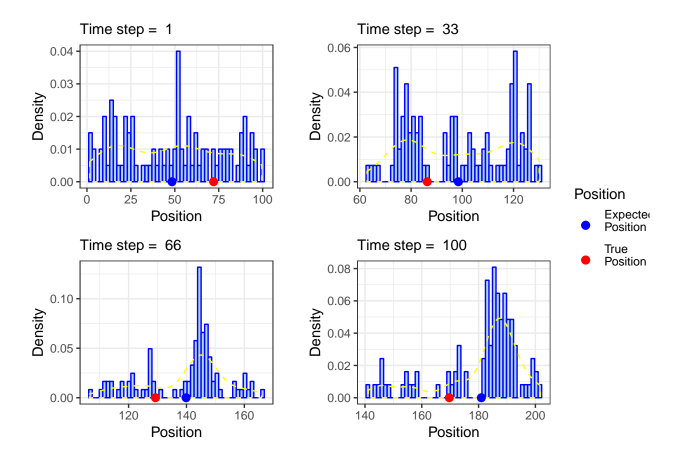


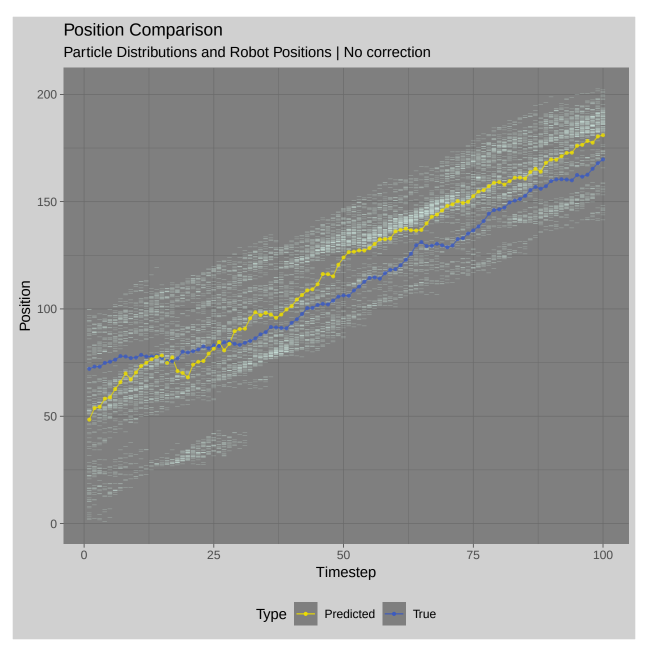


1.3 Weights of Filter equal to 1

Finally, show and explain what happens when the weights in the particle filter are always equal to 1, i.e. there is no correction.

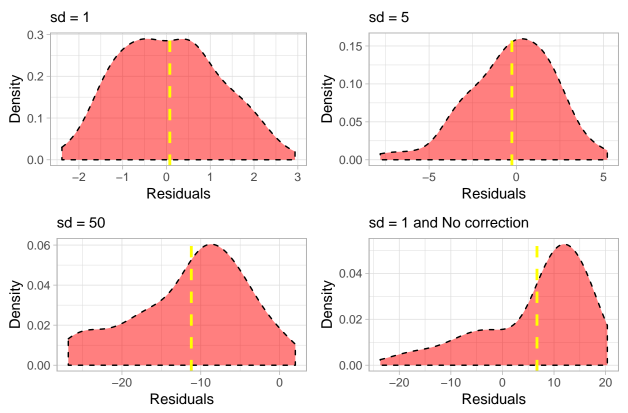
Finally, we run the algorithm without the correction part. That means that when we resample we do not use the emition model in order to compute the weights. It is obvious that this cause bad results. The entire idea of the *Particle Samlper* is that we weight the performance of each particle in order to avoid generating particles that are not possible to be found in the data. If you exclude this step, the algorithm is only using the transition model, which means that we do not actually use all information available. The filter is only simulating another state spaced model on the particles from the beginning, without taking into account any observations. Since the initial sampling is coming from the uniform distribution, it is like applying successively the transition model to the initial expected value. That kind of sampler is not a particle filter but one regular Importance sampler. Again, as before, if we increase the number of particles, the expected value wil converge to the true value but it will be really computational expensive.





Finally, we also plot the residuals distribution for all 4 methods above. For the residuals, we can see that will we increase the standard deviation, the distribution of the residuals is actually becoming wider and the mean of the residuals is of course increasing.

Residuals for different standard deviation



#Plot in the same figure all the predictions for diff sd # $df_pred = data.frame(predict_sd1,predict_sd5,predict_sd50,predict_noCorr)$ # $plt_all_predictions = plt_predict(exp_pos = df_pred,true_states,$ # $sd = c(1,5,50,"1 \ and \ No \ correction"))$