# MSc Statistical Programming 2023 Assessed Practical Assignment

P151

December 7, 2023

# 1 British House Prices

#### 1.1 Exploratory Data Analysis

Figure 1 below displays the evolution of House Prices in England over time.

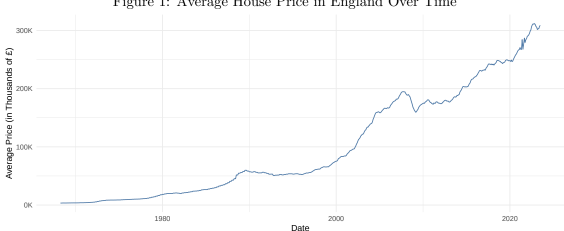


Figure 1: Average House Price in England Over Time

Figure 2 below additionally displays the evolution of House Prices in the Oxford Regions as well as England as a whole over time.

Table 1 below shows that of the ten regions with the highest median of the ratio regional average house price average house price in England. Perhaps unsurprisingly, we can see that the ten most expensive regions by this metric are geographically concentrated in London and its suburbs, and that no Oxford Regions are included.

Figure 3 below compares the month-on-month percentage increase in House Prices in England to monthly inflation, defined as the percentage increase in the monthly Consumer Price Index over the period in which data for both is available. We can see that like in most countries, the increase in the average price of a house has drastically outpaced the general cost-of-living increase since the year 2000.



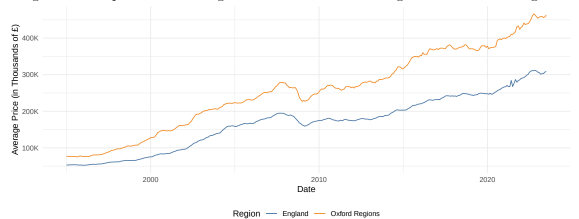


Table 1: Ten Highest Median Ratios of Average House Prices

	Region Name	Median Ratio	Initial Price	Final Price	% Increase	Oxford?
1	Kensington and Chelsea	4.59	182694.83	1344669.00	636.02	FALSE
2	City of Westminster	3.24	133025.28	954356.00	617.42	FALSE
3	Camden	2.92	120932.89	828389.00	585.00	FALSE
4	Hammersmith and Fulham	2.79	124902.86	798537.00	539.33	FALSE
5	City of London	2.56	91448.98	905489.00	890.16	FALSE
6	Richmond upon Thames	2.51	109326.12	770107.00	604.41	FALSE
7	Elmbridge	2.34	106524.00	678279.00	536.74	FALSE
8	Islington	2.32	92516.49	710181.00	667.63	FALSE
9	Wandsworth	2.15	88559.04	632895.00	614.66	FALSE
10	Mole Valley	1.98	101899.76	563083.00	452.59	FALSE

# 2 Chromosome Painting

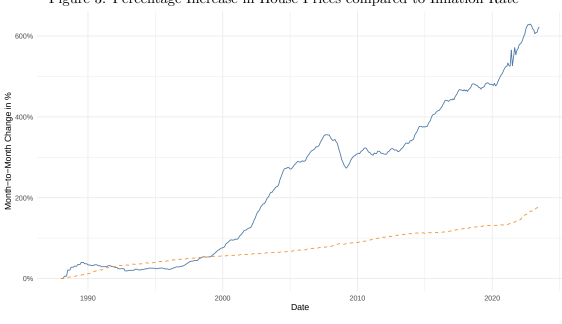
#### 2.1 Implementation of forward Algorithm

```
# Function to calculate emission probability (eq. 3)
emission_probability <- function(observed, reference, error = 0.1) {
   return ((1 - error)^(observed == reference) * error^(observed != reference))
}

# matrices alpha and beta , both with K rows and T columns
forward <- function(haps, hap, error = 0.1) {
   K <- nrow(haps)
   T <- ncol(haps)

# Initialize alpha matrix
alpha <- matrix(0, nrow = K, ncol = T)

# Compute initial and emission probabilities
pi <- 1 / K # (eq. 1)
for (k in 1:K) { # (eq. 4)
   alpha[k, 1] <- pi * emission_probability(hap[1], haps[k, 1], error)</pre>
```



Metric — House Prices in England — Inflation Rate

Figure 3: Percentage Increase in House Prices compared to Inflation Rate

```
# Induction step to compute (eq. 5)
for (t in 2:T) {
   for (k in 1:K) {
      transition_sum <- 0
      for (i in 1:K) {
      A_ik <- ifelse(i == k, (1 - 0.999) / K + 0.999, (1 - 0.999) / K) # (eq. 2)
      transition_sum <- transition_sum + alpha[i, t - 1] * A_ik
   }
   b_kt <- emission_probability(hap[t], haps[k, t], error)
   alpha[k, t] <- transition_sum * b_kt
   }
}
return(alpha)
</pre>
```

## 2.2 Unit Test for forward Algorithm Implementation

```
test_that("alpha matrix has expected form when haps and hap always match", {
   K <- 10  # Number of rows in haps
   T <- 15  # Number of columns in haps = length of hap
   e <- 0.1  # Error rate (= default value)</pre>
```

```
# Create haps and hap such that they always match (both all 0)
haps <- matrix(0, nrow = K, ncol = T)
hap <- rep(0, T)

# Run the forward function
alpha <- forward(haps, hap, error = e)

# Check first column
expected_first_column <- rep((1 - e) / K, K)
expect_equal(alpha[, 1], expected_first_column)

# Check all other columns
for (t in 2:T) {
    expect_equal(alpha[, t], alpha[, t - 1] * (1 - e), tolerance = 1e-5)
}
})

## Test passed</pre>
```

#### 2.3 Implementation of backward Algorithm

```
backward <- function(haps, hap, error = 0.1) {</pre>
  K <- nrow(haps)</pre>
  T <- ncol(haps)
  # Initialize beta matrix (eq. 6)
  beta <- matrix(0, nrow = K, ncol = T)</pre>
  beta[, T] <- 1 # Set last column to 1
  # Induction step (eq. 7)
  for (t in (T-1):1) {
      for (k in 1:K) {
        sum_transition <- 0</pre>
        for (i in 1:K) { # (eq. 2)
          A_ki \leftarrow ifelse(k == i, (1 - 0.999) / K + 0.999, (1 - 0.999) / K)
           b_i_t_plus_1 <- emission_probability(hap[t+1], haps[i, t+1], error)</pre>
           sum_transition <- sum_transition +</pre>
               A_{ki} * b_{i_tplus_1} * beta[i, t+1]
        beta[k, t] <- sum_transition</pre>
  return(beta)
```

#### 2.4 Implementation of gamma Function

```
gamma <- function(haps, hap, error = 0.1) {
    K <- nrow(haps)
    T <- ncol(haps)

# Compute alpha and beta matrices
    alpha <- forward(haps, hap, error)
    beta <- backward(haps, hap, error)

# Compute normalization factor (denominator)
    norm_factor <- sum(alpha[, T])

# Initialize gamma matrix
gamma_matrix <- matrix(0, nrow = K, ncol = T)

# Update gamma values (eq. 8)
for (t in 1:T) {
    for (k in 1:K) {
        gamma_matrix[k, t] <- (alpha[k, t] * beta[k, t]) / norm_factor
    }
}
return(gamma_matrix)
}</pre>
```

#### 2.5 Unit Test for gamma Function Implementation

```
test_that("column sums of gamma matrix all equal 1", {
   set.seed(42)  # For reproducibility
   K <- 10  # Number of rows in haps
   T <- 15  # Number of columns in haps = length of hap
   e <- 0.1  # Error rate (= default value)

# Create random haps and hap with 0 or 1 entries
   haps <- matrix(sample(0:1, K * T, replace = TRUE), nrow = K, ncol = T)
   hap <- sample(0:1, T, replace = TRUE)

# Run the gamma function
   gamma_matrix <- gamma(haps, hap, error = e)

# Check that each column sum is close to 1
   for (t in 1:T) {
        expect_equal(sum(gamma_matrix[, t]), 1, tolerance = 1e-5)
    }
}

## Test passed</pre>
```

## 2.6 Computational Complexity of forward and backward Algorithms

- 1. The forward algorithm has a time complexity of  $\mathcal{O}(K^2 \cdot T)$  as for each time step t, it iterates over K states, and within each state again iterates over K states to compute the transition probabilities.
- 2. The backward algorithm has an identical time complexity of  $\mathcal{O}(K^2 \cdot T)$  as it performs the same iterations, just in a different oder.

Figures 4 and 5 below show benchmark results for both the forward and backward algorithms, executed 10 times each for all  $K \times T$  with  $K, L = 5, \ldots, 20$ . The fitted lines for the median execution time for each value of K and T, respectively, look quadratic in K and linear in T.

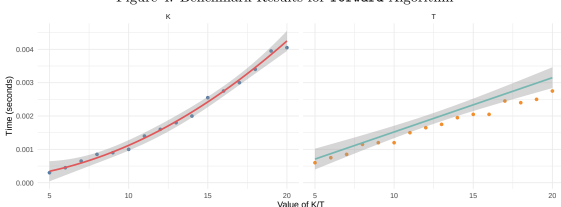


Figure 4: Benchmark Results for forward Algorithm

Median execution time required for varying values of K and T. Each point represents the median execution time across all 10 simulations for a given K or T.

The red and turquoise lines are smoothed fits on all data points indicating the trend of time complexity, with fitting quadratic in K and linear in T

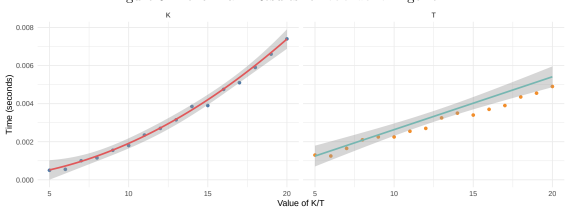


Figure 5: Benchmark Results for backward Algorithm

Median execution time required for varying values of K and T. Each point represents the median execution time across all 10 simulations for a given K or T.

The red and turquoise lines are smoothed fits on all data points indicating the trend of time complexity, with fitting quadratic in K and linear in T

#### 2.7 Implementation of forward2 Algorithm

```
forward2 <- function(haps, hap, error = 0.1) {
    K <- nrow(haps)
    T <- ncol(haps)
    alpha <- matrix(0, nrow = K, ncol = T)
    pi <- 1 / K

for (k in 1:K) {
        alpha[k, 1] <- pi * emission_probability(hap[1], haps[k, 1], error)
    }

for (t in 2:T) {
        phi <- sum(alpha[, t - 1]) * (1 - 0.999) / K
        for (k in 1:K) {
            A_ik <- ifelse(k == k, (1 - 0.999) / K + 0.999, (1 - 0.999) / K)
            b_kt <- emission_probability(hap[t], haps[k, t], error)
            alpha[k, t] <- (phi + 0.999 * alpha[k, t - 1]) * b_kt
        }
    }

    return(alpha)
}</pre>
```

```
test_that("forward and forward2 produce the same outputs", {
    set.seed(42)  # For reproducibility
    K <- 10  # Number of rows in haps
    T <- 15  # Number of columns in haps = length of hap
    e <- 0.1  # Error rate (= default value)

# Generate example data
    haps <- matrix(sample(0:1, K * T, replace = TRUE), nrow = K)
    hap <- sample(0:1, T, replace = TRUE)

# Run both functions
    alpha_forward <- forward(haps, hap, e)
    alpha_forward2 <- forward2(haps, hap, e)

# Check if the outputs are equal
    expect_equal(alpha_forward, alpha_forward2, tolerance = 1e-5)
})

## Test passed</pre>
```

The forward2 algorithm has a time complexity of  $\mathcal{O}(K \cdot T)$  as it computes the initial probabilities  $(\mathcal{O}(K))$  and then for each time step t, it sums over all K states  $(\mathcal{O}(K))$  and computes equation (14)  $(\mathcal{O}(1))$ . Since the main loop runs for each time step T, and within each time step, two  $\mathcal{O}(K)$  operations are performed (one for summing alpha and one for

the loop over K), the overall complexity of the forward2 function is  $\mathcal{O}(K \cdot T)$ . Figure 6 below displays this improved performance.

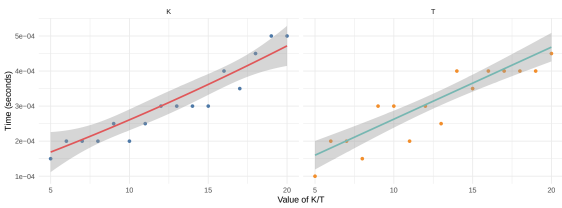


Figure 6: Benchmark Results for forward2 Algorithm

Median execution time required for varying values of K and T. Each point represents the median execution time across all 10 simulations for a given K or T.

The red and turquoise lines are smoothed fits on all data points indicating the trend of time complexity, with fitting quadratic in K and linear in T

#### 2.8 Implementation of backward2 Algorithm

```
backward2 <- function(haps, hap, error = 0.1) {</pre>
  K <- nrow(haps)</pre>
  T <- ncol(haps)
  # Initialize beta matrix (eq. 6)
  beta <- matrix(0, nrow = K, ncol = T)</pre>
  beta[, T] <- 1 # Set last column to 1
  # Induction step (eq. 14)
  for (t in (T-1):1) {
    # Calculate the phi term which is constant for all k (eq. 13)
    phi <- sum(beta[, t+1] *
                  sapply(1:K, function(i) emission_probability(hap[t+1],
                               haps[i, t+1], error))) * (1 - 0.999) / K
    for (k in 1:K) { # (eq. 14)
      beta[k, t] \leftarrow phi + 0.999 * beta[k, t+1] *
        emission_probability(hap[t+1], haps[k, t+1], error)
  return(beta)
```

```
test_that("backward and backward2 produce the same outputs", {
    set.seed(42)  # For reproducibility
    K <- 10  # Number of rows in haps
    T <- 15  # Number of columns in haps = length of hap
    e <- 0.1  # Error rate (= default value)

# Generate example data
    haps <- matrix(sample(0:1, K * T, replace = TRUE), nrow = K)
    hap <- sample(0:1, T, replace = TRUE)

# Run both functions
beta_backward <- backward(haps, hap, e)
beta_backward2 <- backward2(haps, hap, e)

# Check if the outputs are equal
    expect_equal(beta_backward, beta_backward2, tolerance = 1e-5)
})

## Test passed</pre>
```

The backward2 algorithm has a time complexity of  $\mathcal{O}(K \cdot T)$  as it computes the initial probabilities  $(\mathcal{O}(K))$  and then for each time step t, it sums over all K states  $(\mathcal{O}(K))$  and computes equation (14)  $(\mathcal{O}(1))$ . Since the main loop runs for each time step T, and within each time step, two  $\mathcal{O}(K)$  operations are performed (one for summing alpha and one for the loop over K), the overall complexity of the forward2 function is  $\mathcal{O}(K \cdot T)$ . Figure 7 below displays this improved performance.

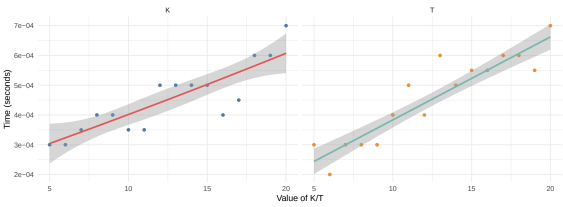


Figure 7: Benchmark Results for backward2 Algorithm

Median execution time required for varying values of K and T. Each point represents the median execution time across all 10 simulations for a given K or T.

The red and turquoise lines are smoothed fits on all data points indicating the trend of time complexity, with fitting quadratic in K and linear in T

#### 2.9 Implementation of gamma2 Function

```
gamma2 <- function(haps, hap, error = 0.1) {
  K <- nrow(haps)
  T <- ncol(haps)</pre>
```

```
# Compute alpha and beta matrices using forward2 and backward2
alpha <- forward2(haps, hap, error)
beta <- backward2(haps, hap, error)

# Compute normalization factor (denominator)
norm_factor <- sum(alpha[, T])

# Initialize gamma matrix
gamma_matrix <- matrix(0, nrow = K, ncol = T)

# Update gamma values (eq. 8)
for (t in 1:T) {
   for (k in 1:K) {
      gamma_matrix[k, t] <- (alpha[k, t] * beta[k, t]) / norm_factor
   }
}
return(gamma_matrix)
}</pre>
```

```
test_that("gamma and gamma2 produce the same outputs", {
    set.seed(42)  # For reproducibility
    K <- 10  # Number of rows in haps
    T <- 15  # Number of columns in haps = length of hap
    e <- 0.1  # Error rate (= default value)

# Generate example data
    haps <- matrix(sample(0:1, K * T, replace = TRUE), nrow = K)
    hap <- sample(0:1, T, replace = TRUE)

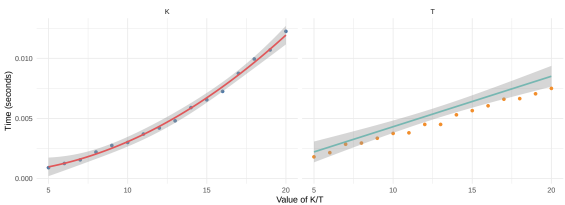
# Run both functions
    gamma_matrix <- gamma(haps, hap, e)
    gamma_matrix2 <- gamma2(haps, hap, e)

# Check if the outputs are equal
    expect_equal(gamma_matrix, gamma_matrix2, tolerance = 1e-5)
})

## Test passed</pre>
```

Figures 8 and 9 below show that the empirical time complexity of gamma and gamma2 indeed matches the reasoning laid out above.

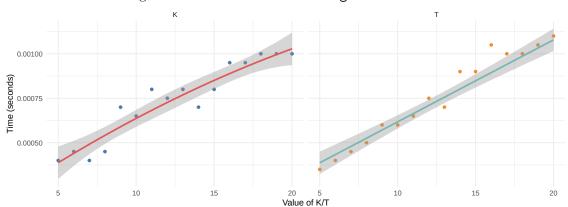
Figure 8: Benchmark Results for gamma Function



Median execution time required for varying values of K and T. Each point represents the median execution time across all 10 simulations for a given K or T.

The red and turquoise lines are smoothed fits on all data points indicating the trend of time complexity, with fitting quadratic in K and linear in T

Figure 9: Benchmark Results for gamma2 Function



Median execution time required for varying values of K and T. Each point represents the median execution time across all 10 simulations for a given K or T.

The red and turquoise lines are smoothed fits on all data points indicating the trend of time complexity, with fitting quadratic in K and linear in T

```
# Extract the first 5 target haplotypes
first_5_haps <- samples_raw[1:5, ]

paint_and_sum <- function(target_hap, ref_panel) {
    gamma_results <- gamma2(ref_panel, target_hap, error = 0.1)

# Sum contributions from YRI and CEU haplotypes
    yri_sum <- rowSums(gamma_results[grepl("YRI", rownames(ref_panel)), ])
    ceu_sum <- rowSums(gamma_results[grepl("CEU", rownames(ref_panel)), ])

return(list(YRI = yri_sum, CEU = ceu_sum))
}

# Apply function to first 5 target haplotypes (parallelized)
first_5_painted <- mclapply(1:nrow(first_5_haps), function(i) {
    paint_and_sum(first_5_haps[i, ], refpanel_raw)</pre>
```

```
}, mc.cores = detectCores())
compile_results_matrix <- function(painted_results) {</pre>
  results_matrix <- matrix(nrow = length(painted_results), ncol = 3)</pre>
  colnames(results_matrix) <- c("Sum_YRI", "Sum_CEU", "Percentage_YRI")</pre>
  rownames(results_matrix) <- rownames(first_5_haps)</pre>
  # Loop through each list of results and calculate the sum of YRI, CEU,
  # percentage of YRI
  for (i in seq_along(painted_results)) {
    sum_yri <- sum(painted_results[[i]]$YRI)</pre>
    sum_ceu <- sum(painted_results[[i]]$CEU)</pre>
    total_sum <- sum_yri + sum_ceu
    percentage_yri <- (sum_yri / total_sum) * 100</pre>
    results_matrix[i, ] <- c(sum_yri, sum_ceu, percentage_yri)</pre>
  colnames(results_matrix) <- c("Sum YRI", "Sum CEU",</pre>
                                   "Percentage of YRI in Total")
  return(results_matrix)
painted <- compile_results_matrix(first_5_painted)</pre>
```

Table 2 below shows that among the first 5 target haplotypes, haplotypes 2 and 5 show a high contribution of YRI haplotypes. Target haplotype 5 suggests an entirely African genetic background over the investigated chromosome.

Table 2: Sums and Proportion of YRI and CEU Ancestry in the first 5 Target Haplotypes

	Sum YRI	Sum CEU	Percentage of YRI in Total
ASW-NA19818	46.12	70.88	39.42
ASW-NA19819	98.58	18.42	84.26
ASW-NA19834	30.18	86.82	25.80
ASW-NA19835	71.69	45.31	61.27
ASW-NA19900	116.46	0.54	99.54