

assignment1

February 6, 2021

1 Assignment 1 - LS on 2nd order polynomial

First, we will define the necessary functions and objects to carry out task 1.4. Then, we will examine each sub-case in task 1.4.

```
[1]: # Imports
      %matplotlib inline
      import numpy as np
      import matplotlib.pyplot as plt
      from scipy.stats import laplace, norm
      from typing import Callable
```

```
[2]: def arbitrary_poly(params: list) -> Callable:
      """
      Takes the parameters in params and makes them into a polynomial
       $\theta_0 + \theta_1 x + \theta_2 x^2 + \dots + \theta_n x^n$ 
      where  $n$  is model order and  $\theta_0$  is bias.
      Args:
          params: list on format [ $\theta_0$ ,  $\theta_1$ ,  $\theta_2$ , ...,  $\theta_n$ ]
      Returns:
          Function with single argument  $x$ .
      """
      poly_model = lambda x: sum([p*(x**i) for i, p in enumerate(params)])
      return poly_model
```

```
[3]: class GaussParams:
      def __init__(self, mean: float, std: float):
          self.mean = mean
          self.std = std
          self.var = std**2

          # aliases, these are the names used in scipy.stats functions
          self.loc = self.mean
          self.scale = self.std

      def __iter__(self): # to allow tuple unpacking
          return iter((self.mean, self.std))
```

```

class LaplaceParams:
    def __init__(self, mean: float, beta: float):
        self.mean = mean
        self.beta = beta

        # aliases, these are the names used in scipy.stats functions
        self.loc = self.mean
        self.scale = self.beta

    def __iter__(self): # to allow tuple unpacking
        return iter((self.mean, self.beta))

```

1.1 Measurement noise generation from a chosen distribution

Probability density functions (PDFs):

Gaussian pdf: $P_N(x) = \frac{1}{\sigma\sqrt{2\pi}} e^{-\frac{1}{2}(\frac{x-\mu}{\sigma})^2}$

Laplacian pdf: $P_L(x) = \frac{1}{2b} e^{-\frac{|x-\mu|}{b}}$

```

[4]: def generate_noise(N: int, magnitude: float, alpha: float, gauss_params:
    ↳GaussParams, laplace_params: LaplaceParams, seed: int = None) -> np.ndarray:
        """
        Generate noise based on the mixture model: alpha * gaussian + (1 - alpha) *
    ↳laplacian.
        Args:
            N: number of noise points to generate.
            magnitude: the noise amplitude/magnitude.
            alpha: fraction of gaussian and laplacian, should be in interval [0, 1].
            gauss_params: mean and std of gaussian distribution
            laplace_params: mean and beta of laplacian distribution
        Returns:
            A list of N noise points.
        """
        if seed is not None:
            np.random.seed(seed)

        if alpha == 1:
            assert gauss_params is not None
            noise = np.random.normal(*gauss_params, N)
        elif alpha == 0:
            assert laplace_params is not None
            noise = np.random.laplace(*laplace_params, N)
        else:
            assert gauss_params is not None
            assert laplace_params is not None

```

```

    assert 0 < alpha < 1
    noise = np.zeros((N,))
    for i in range(N):
        r = np.random.uniform(0,1)
        if r <= alpha:
            noise[i] = np.random.normal(*gauss_params)
        else:
            noise[i] = np.random.laplace(*laplace_params)
    noise *= magnitude

    return noise

```

1.2 Least squares regression - LS

1.2.1 Given the sampled data we can estimate the underlying polynomial model using LS regression

Solution to a least squares problem is given by the normal equation:

$$\hat{\theta} = (\mathbf{u}^T \cdot \mathbf{u})^{-1}(\mathbf{u}^T \cdot \mathbf{y})$$

1.2.2 Step 1 - rewrite the model in matrix form to get the data tensor \mathbf{u}

$$y_{\text{measured}} = \theta_0 + \theta_1 \cdot u^1 + \dots + \theta_n \cdot u^n + \text{noise} = [1 \ u^1 \dots u^n] \cdot [\theta_0 \ \theta_1 \dots \theta_n]^T + \text{noise}$$

$$\mathbf{u} = [1 \ u^1 \dots u^n]$$

1.2.3 Step 2

calculate $(\mathbf{u}^T \cdot \mathbf{u})^{-1}$

1.2.4 Step 3

calculating dot product calculate $(\mathbf{u}^T \cdot \mathbf{y})$

1.2.5 Step 4

solve for $\hat{\theta} = (\mathbf{u}^T \cdot \mathbf{u})^{-1}(\mathbf{u}^T \cdot \mathbf{y})$

```

[5]: def least_square(u: np.ndarray, y: np.ndarray, n_params: int) -> np.ndarray:
    """
    Uses least square (LS) to make the best fitting list of parameters.
    Args:
        u: data x values, shape = (N_samples,)
        y: data y values, shape = (N_samples,)
        n_params: number of parameters to find
    """
    N = u.shape[0]

```

```

# Create data tensor
u_tensor = np.ones((N,1))
for i in range(1, n_params):
    u_tensor = np.append(u_tensor, np.power(u.reshape((-1,1)), i), axis=1)

# Solve for LS params
u_dot_u = u_tensor.T @ u_tensor
u_dot_y = u_tensor.T @ y
# linalg.solve is more numerically stable than linalg.inv and is therefore
→preferred
ls_params = np.linalg.solve(u_dot_u, u_dot_y)

return ls_params

```

```

[6]: def ls_statistics(
    u: np.ndarray,
    y: np.ndarray,
    true_params: np.ndarray,
    ls_params_estimate: np.ndarray) -> tuple:
    """
    Calculate statistics on LS performance.
    Args:
        u: data points x value.
        y: measured data points (with noise).
        true_params: list of the true parameters.
        ls_params_estimate: list of the LS parameters estimates.
    Returns:
        Tuple on the form
            (RMSE, d)
        where d is the Euclidean distance between the true and estimated
        →parameters.
    """
    ls_model = arbitrary_poly(ls_params_estimate)
    rmse = np.sqrt(np.mean( (ls_model(u) - y)**2 ))
    d = np.linalg.norm( ls_params_estimate - np.array(true_params) )

    return rmse, d

```

1.3 Task 1.4: Set up global parameters for each case

```

[7]: # True model
true_params = [1,-8,4]
n_params = len(true_params)
y_model = arbitrary_poly(true_params)

# Hyperparameters for noise generation

```

```

magnitude = 1.2 # noise magnitude
N = 20          # number of samples

# Setup RNG
np.random.seed(0) # Non-random generation between code executions. Comment out
↳for true random

# Generate data points
range_low = -1
range_high = 1
u = np.sort(np.random.uniform(range_low, range_high, N))
y_true = y_model(u)

```

1.4 Case 1: $\alpha = 1$ and σ is varying

```

[8]: # Parameters
alpha = 1

# Add noise to the generated data points - thus simulating measurement
ls_data = []
sigma_max = 2
for sigma in np.linspace(0, sigma_max, 1000):
    noise = generate_noise(N, magnitude, alpha, GaussParams(0, sigma), None,
↳seed=None)
    y = y_true + noise

    ls_params = least_square(u, y, n_params)
    rmse, d = ls_statistics(u, y, true_params, ls_params)
    ls_data.append((sigma, ls_params, y, d))

```

```

[9]: # Plot
sigmas, ds = [], []
for sigma, _, _, d in ls_data:
    sigmas.append(sigma)
    ds.append(d)

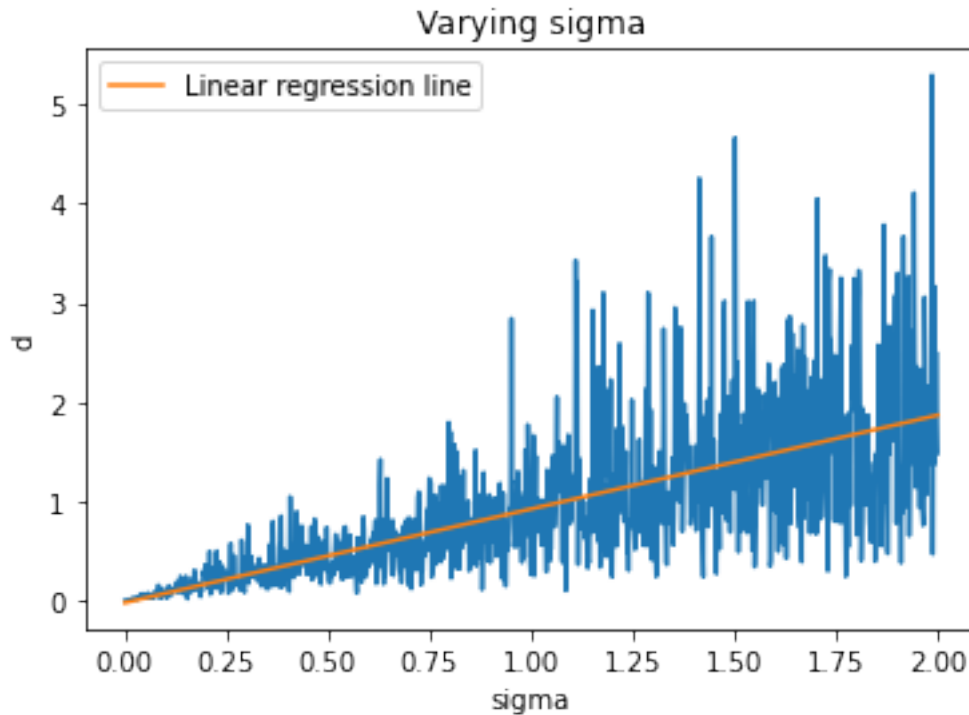
sigmas = np.array(sigmas)
lin_reg_line = least_square(sigmas, np.array(ds), 2)
print(lin_reg_line)

plt.figure()
plt.plot(sigmas, ds)
plt.plot(sigmas, arbitrary_poly(lin_reg_line)(sigmas), label="Linear regression,
↳line")
plt.xlabel("sigma")
plt.ylabel("d")

```

```
plt.title("Varying sigma")
plt.legend()
plt.show()
```

```
[-0.02346982  0.94533022]
```



1.4.1 Comments to case 1

σ is the standard deviation of the Gaussian distribution and determines how “wide” the distribution is. This can be seen in the graph above where σ is varying between 0 and 2. At low σ the LS estimates of the parameters are not spread out much. This tells us that low σ , meaning low noise, gives a good LS estimate. Increasing σ , we see clearly that the LS estimates become more varied in precision, meaning that they deviate more from the true parameters.

There is a clear upwards trend in the data with increasing σ , and the linear regression line confirms this point. This is in tune with what we would expect. Larger σ means larger spread in the noise which gives a more uncertain estimate.

1.5 Case 2: $\alpha = 0$ and β is varying

```
[10]: # Parameters
      alpha = 0

      # Add noise to the generated data points - thus simulating measurement
```

```

ls_data = []
beta_max = 2
for beta in np.linspace(0, beta_max, 1000):
    noise = generate_noise(N, magnitude, alpha, None, LaplaceParams(0, beta),
        seed=None)
    y = y_true + noise

    ls_params = least_square(u, y, n_params)
    rmse, d = ls_statistics(u, y, true_params, ls_params)
    ls_data.append((beta, ls_params, y, d))

```

```

[11]: # Plot
betas, ds = [], []
for beta, params, y, d in ls_data:
    betas.append(beta)
    ds.append(d)

betas = np.array(betas)
lin_reg_line = least_square(betas, np.array(ds), 2)
print(lin_reg_line)

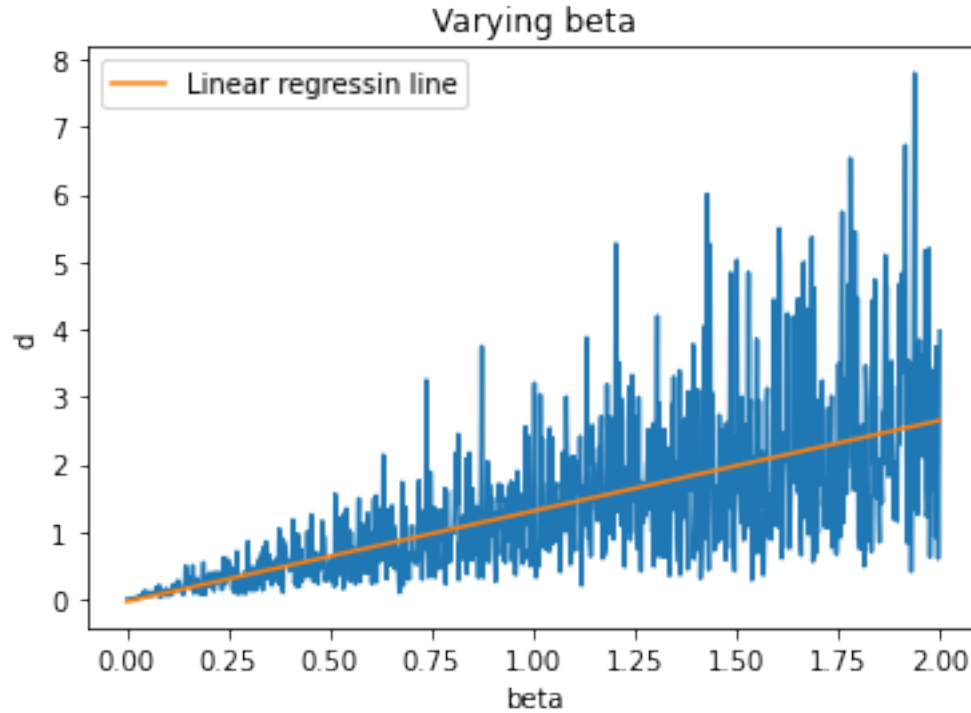
plt.figure()
plt.plot(betas, ds)
plt.plot(betas, arbitrary_poly(lin_reg_line)(betas), label="Linear regression line")
plt.xlabel("beta")
plt.ylabel("d")
plt.title("Varying beta")
plt.legend()
plt.show()

```

```

[-0.03159766  1.34124622]

```



1.5.1 Comments to case 2

Similarly to the Gaussian, the β parameter determines how wide the Laplacian distribution is. The same points as above can be made for the Laplacian, i.e. that larger β leads to poorer LS estimate. Again, we also see a clear upwards trend with increasing β , which also is expected as larger β means wider distribution which gives noise values that are more spread out.

An interesting note is that the slope of the linear regression line is here greater than for the Gaussian. This means that increasing β will affect the LS estimate more than the same increase in σ . A possible reason for this is that the covariance of the Gaussian is simply σ^2 , while for the Laplacian it is $2\beta^2$.

1.6 Case 3: $\sigma = \beta = 1$ and α is varying

```
[12]: # Parameters
gp = GaussParams(0, 1)
lp = LaplaceParams(0, 1)

# Add noise to the generated data points - thus simulating measurement
ls_data = []
for alpha in np.linspace(0,1,1000):
    noise = generate_noise(N, magnitude, alpha, gp, lp, seed=None)
    y = y_true + noise
```



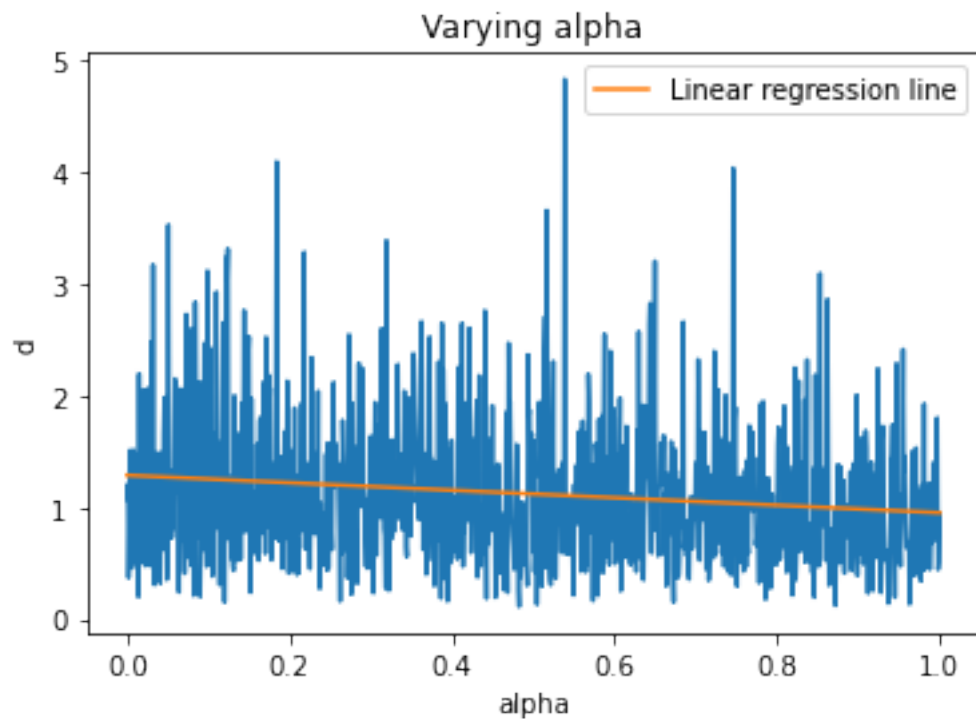
```
ls_params = least_square(u, y, n_params)
rmse, d = ls_statistics(u, y, true_params, ls_params)
ls_data.append((alpha, ls_params, y, d))
```

```
[13]: # Plot
alphas, ds = [], []
for alpha, _, _, d in ls_data:
    alphas.append(alpha)
    ds.append(d)

alphas = np.array(alphas)
lin_reg_line = least_square(alphas, np.array(ds), 2)
print(lin_reg_line)

plt.figure()
plt.plot(alphas, ds)
plt.plot(alphas, arbitrary_poly(lin_reg_line)(alphas), label="Linear regression ↵
↵line")
plt.xlabel("alpha")
plt.ylabel("d")
plt.title("Varying alpha")
plt.legend()
plt.show()
```

```
[ 1.28788062 -0.33508435]
```



1.6.1 Comments to case 3

Looking at the raw data (blue plot), it is hard to make out any clear trend, but the linear regression line has a negative slope. Since greater α means a greater probability of having more Gaussian noise, the negative slope means that the LS estimate becomes slightly better with more Gaussian noise than with more Laplacian noise. However, the difference is not too pronounced. Above we saw that pure Gaussian noise has better performance than pure Laplacian noise. Combined with the negative slope we see here, we can conclude that Gaussian noise is preferred since it gives a slightly better LS estimate (most of the time).

In fact, it is known that the LS estimate when we have Gaussian noise is the same as the Maximum Likelihood estimate, so this solidifies the conclusion that Gaussian noise is preferred.

1.6.2 General comments to all cases

In this assignment we only used 20 samples. It is possible that if we were to use more samples, the estimates would be better in some or all cases since. It could also be that we can be lucky/unlucky on where we draw the 20 samples, so that in some cases they align well with the model and in other cases it does not. This is likely why we see a lot of variation/“noise” in the plots. Where d is small, we got “lucky” and where it is large we got “unlucky”.