```
import numpy as np
import pandas as pd
import numpy.random as rnd
from sklearn.datasets import load_breast_cancer

cancer = load_breast_cancer(as_frame = True)
df = cancer.frame
df
```

	mean radius	mean texture	mean perimeter	mean area	mean smoothness	mean compactness	mean concavity	mean concave points	mean symmetry	mean fractal dimension	•••	worst texture	worst perimeter	worst area
0	17.99	10.38	122.80	1001.0	0.11840	0.27760	0.30010	0.14710	0.2419	0.07871		17.33	184.60	2019.0
1	20.57	17.77	132.90	1326.0	0.08474	0.07864	0.08690	0.07017	0.1812	0.05667		23.41	158.80	1956.0
2	19.69	21.25	130.00	1203.0	0.10960	0.15990	0.19740	0.12790	0.2069	0.05999		25.53	152.50	1709.0
3	11.42	20.38	77.58	386.1	0.14250	0.28390	0.24140	0.10520	0.2597	0.09744		26.50	98.87	567.7
4	20.29	14.34	135.10	1297.0	0.10030	0.13280	0.19800	0.10430	0.1809	0.05883		16.67	152.20	1575.0
564	21.56	22.39	142.00	1479.0	0.11100	0.11590	0.24390	0.13890	0.1726	0.05623		26.40	166.10	2027.0
565	20.13	28.25	131.20	1261.0	0.09780	0.10340	0.14400	0.09791	0.1752	0.05533		38.25	155.00	1731.0
566	16.60	28.08	108.30	858.1	0.08455	0.10230	0.09251	0.05302	0.1590	0.05648		34.12	126.70	1124.0
567	20.60	29.33	140.10	1265.0	0.11780	0.27700	0.35140	0.15200	0.2397	0.07016		39.42	184.60	1821.0
568	7.76	24.54	47.92	181.0	0.05263	0.04362	0.00000	0.00000	0.1587	0.05884		30.37	59.16	268.6
569 rows × 31 columns														

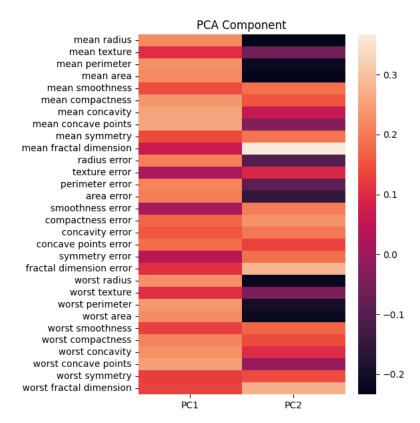
x = df[cancer['feature_names']]
y

	mean radius	mean texture	mean perimeter	mean area	mean smoothness	mean compactness	mean concavity	mean concave points	mean symmetry	mean fractal dimension	 worst radius	worst texture
0	17.99	10.38	122.80	1001.0	0.11840	0.27760	0.30010	0.14710	0.2419	0.07871	 25.380	17.33
1	20.57	17.77	132.90	1326.0	0.08474	0.07864	0.08690	0.07017	0.1812	0.05667	 24.990	23.41
2	19.69	21.25	130.00	1203.0	0.10960	0.15990	0.19740	0.12790	0.2069	0.05999	 23.570	25.53
3	11.42	20.38	77.58	386.1	0.14250	0.28390	0.24140	0.10520	0.2597	0.09744	 14.910	26.50
4	20.29	14.34	135.10	1297.0	0.10030	0.13280	0.19800	0.10430	0.1809	0.05883	 22.540	16.67
564	21.56	22.39	142.00	1479.0	0.11100	0.11590	0.24390	0.13890	0.1726	0.05623	 25.450	26.40
565	20.13	28.25	131.20	1261.0	0.09780	0.10340	0.14400	0.09791	0.1752	0.05533	 23.690	38.25
566	16.60	28.08	108.30	858.1	0.08455	0.10230	0.09251	0.05302	0.1590	0.05648	 18.980	34.12
567	20.60	29.33	140.10	1265.0	0.11780	0.27700	0.35140	0.15200	0.2397	0.07016	 25.740	39.42
568	7.76	24.54	47.92	181.0	0.05263	0.04362	0.00000	0.00000	0.1587	0.05884	 9.456	30.37

569 rows × 30 columns

```
x_mean = x.mean()
x_std = x.std()
z = (x-x_mean)/x_std
c = z.cov()
eigenvalues, eigenvectors = np.linalg.eig(c)
idx = eigenvalues.argsort()[::-1]
eigenvalues = eigenvalues[idx]
eigenvectors = eigenvectors[:,idx]
```

```
explained_var = np.cumsum(eigenvalues) / np.sum(eigenvalues)
explained_var
     array([0.44272026, 0.63243208, 0.72636371, 0.79238506, 0.84734274,
            0.88758796, 0.9100953 , 0.92598254, 0.93987903, 0.95156881,
            0.961366 , 0.97007138, 0.97811663, 0.98335029, 0.98648812,
             0.98915022, \ 0.99113018, \ 0.99288414, \ 0.9945334 \ , \ 0.99557204, 
            0.99657114, 0.99748579, 0.99829715, 0.99889898, 0.99941502,
            0.99968761, 0.99991763, 0.99997061, 0.99999557, 1.
n_components = np.argmax(explained_var >= 0.50) + 1
n_components
import matplotlib.pyplot as plt
import seaborn as sns
u = eigenvectors[:,:n_components]
pca_component = pd.DataFrame(u,
                             index = cancer['feature_names'],
                             columns = ['PC1','PC2']
# plotting heatmap
plt.figure(figsize =(5, 7))
sns.heatmap(pca_component)
plt.title('PCA Component')
plt.show()
```



Start coding or generate with AI.