Malalignancy happens when changes called transformations occur in qualities that direct cell development. The transformations let the cells separate and increase in an uncontrolled manner. Bosom malignant growth is disease that creates in bosom cells. Regularly, the disease shapes in either the lobules or the conduits of the bosom. Lobules are the organs that produce milk, and channels are the pathways that carry the milk from the organs to the areola. Malignant growth can likewise happen in the greasy tissue or the stringy connective tissue inside your bosom.

The uncontrolled malignant growth cells frequently attack other sound bosom tissue and can head out to the lymph hubs under the arms. The lymph hubs are an essential pathway that help the malignancy cells move to different pieces of the body. See pictures and get familiar with the structure of the bosom. In its beginning times, bosom malignant growth may not bring on any manifestations. Much of the time, a tumor might be too little to even think about being felt, however a variation from the norm can in any case be seen on a mammogram. On the off chance that a tumor can be felt, the principal sign is normally another knot in the bosom that was not there previously. Be that as it may, not all bumps are cancer. Inflammatory bosom disease (IBC) is an uncommon however forceful sort of bosom malignant growth. IBC makes up just somewhere in the range of 1 and 5 percent of all bosom malignancy cases.

With this condition, cells obstruct the lymph hubs close to the bosoms, so the lymph vessels in the bosom can’t appropriately deplete. Rather than making a tumor, IBC makes your bosom swell, look red, and feel warm. A carcinogenic bosom may seem hollowed and thick, similar to an orange strip.

IBC can be forceful and can advance rapidly. Consequently, it’s critical to summon your PCP right on the off chance that you notice any indications. Discover progressively about IBC and the manifestations it can cause.

Objective: High and low risk genetic correlation to local and distant recurrence in early breast cancer at Edgardo Rebagliati Martins Hospital in the period of 2011-2013.

Secondary objective the correlation between lymphovascular invasion, the status of hormonal receptors, molecular type versus type according to genetic platform.

The percentage of high and low risk according to genetic type, the percentage of local and distant recurrence of early breast cancer, the percentage of patients according to molecular subtype.

Methodology: An observational, cross-sectional, study was carried out. Patients aged between 18 and 70 years were diagnosed with breast cancer with or without node involvement without distant metastases at diagnosis.

Results: At the follow-up of 4.9 years, none of the 22 patients presented recurrence either locally or distant, for either high or low risk. High-risk patients were treated with adjuvant chemotherapy and those at low risk did not receive adjuvant chemotherapy.

In the present study, 47.6% were obtained for low-risk patients and 52.4% for high-risk patients. The status of progesterone receptors was related to the type according to the genetic platform. Patients with a luminal molecular subtype B 54%, luminal A of 41%, and 5% for triple negative patients. High risk patients 60% versus 27.3% low risk in luminal A subtype, 63.6% versus 40% luminal B.

Conclusions: At the follow-up of 4.9 years no patient present local recurrence or distance, in favor of genetic study. That progesterone receptors would be related to the low-risk genetic profile. Patients with a diagnosis of early B-luminal breast cancer are in higher percentage of high risk and those luminal A are in higher percentage of low risk according to genetic platform.

Key words: Breast cancer EC I and II, local recurrence, Recurrence distance, genetic platform,