appropriate isolation and oseltamivir use in flu positive patients: 2018 = 27% (6/22); 2019 = 74% (17/23).

Conclusion and relevance Increased flu screening in 2019 despite a national fall in hospitalised flu cases compared with 2018 suggests that clinicians were more likely to consider influenza when rapid diagnostics were available on-site. On-site testing significantly reduced TAT, having a measurable impact on the appropriateness of isolation and oseltamivir use. The absence of isolation facilities in the coronary care unit represented a significant clinical risk of influenza exposure.

REFERENCES AND/OR ACKNOWLEDGEMENTS
No conflict of interest.

EFFECTIVENESS OF NEOADJUVANT TREATMENT IN LOCALLY ADVANCED BREAST CANCER
B Bertan De Lis*, M Mañes, I Soto, E Maroto, EP Gómez, J Solís, C Moriel. Hospital Universitario De Móstoles, Hospital Pharmacy, Móstoles, Spain
10.1136/ejhpharm-2020-eahpconf.438

Background and importance Neoadjuvant chemotherapy has become the standard treatment for patients with inoperable locally advanced tumours, and it is also optional for operable stages. The literature reports rates of 20–30% and 56–66% for pathological complete response (pCR) in patients treated with combinations of anthracyclines and taxanes, and dual blockade of human epidermal growth factor receptor 2 (HER2), respectively.

Aim and objectives To assess the effectiveness of neoadjuvant chemotherapy in stage II/III breast cancer according to expression of HER2 and hormonal receptors (HR) based on pCR.

Material and methods A retrospective observational study was carried out between January 2016 and December 2018 in a second level hospital. Through the Farmatools software, patients were identified. Clinical histories were obtained through Selene software to compile demographic (sex and age) and clinical (stage, HR and HER2, lymph nodes, treatment regimen and pCR results) data.

Results Within the study period, 35 patients received neoadjuvant chemotherapy regimens for breast cancer. Median age of the women was 50 years (IQR 18 years), 54.3% were diagnosed with stage II neoplasia and 62.9% had lymph nodes involved: 40% reached pCR. Patients were classified according to HER2 expression:

- 45.7% showed positive HER2 expression (HER2+), 50% of whom reached pCR after neoadjuvant treatment. In 81.25%, treatment was a docetaxel-carboplatin-trastuzumab regimen plus pertuzumab, obtaining pCR in 33.85%, and 18.75% received chemotherapy regimens based on anthracyclines+taxanes+trastuzumab+ pertuzumab, reaching a pCR of 33.33%.

- In the 54.3% of HER2 negative (HER2–) patients, 31.58% reached pCR: 94.74% received combinations of anthracyclines+taxanes, obtaining a pCR of 33.33%. Only one patient was treated with docetaxel-cyclofosfamide (TC), not achieving pCR. Within the HER2– group, 57.89% did not overexpress any receptor, qualifying as triple negative (TN). All of these patients received regimens based on anthracyclines and reached a pCR of 45.45%.

Conclusion and relevance pCR rates obtained in our centre were correlated with the results described in the literature, and slightly lower in HER2+ patients. In the case of the TN subgroup, a pCR rate greater than in reported data was found, despite being the subgroup with the worst prognosis.

REFERENCES AND/OR ACKNOWLEDGEMENTS
No conflict of interest.

Preliminary Clinical Response of Ribociclib as a Single Agent in Advanced Breast Cancer: In Search of New Therapeutic Indications
Y Cura*, A Sánchez Martín, C Pérez Ramírez, MDM Malaldonado Montoro, MDC Ramírez Tortosa, E Martínez Martínez, A Jiménez Morales, N Márquez Nette. 1University Hospital Virgen De Las Nieves, Pharmacogenetics Unit, Hospital Pharmacy Service, Granada, Spain; 2University Hospital Virgen Macarena, Pharmacogenetics Unit, Hospital Pharmacy Service, Sevilla, Spain; 3University Hospital San Cecilio, Clinical Analysis Service, Granada, Spain; 4University of Granada, Department of Biochemistry, Granada, Spain; 5University of Granada, Pharmaceutical Care Research Group, Granada, Spain
10.1136/ejhpharm-2020-eahpconf.439

Background and importance Ribociclib, an orally bioavailable CDK4/6 inhibitor, is currently approved in combination with an anatamastase inhibitor for the treatment of pre/perimenopausal women with HR positive, HER2 negative advanced breast cancer. Alterations in the CDK4/6-Rb-E2F pathway, which promotes cell proliferation, usually occur in human tumours. Thus ribociclib remains as an attractive therapeutic strategy for the treatment of other neoplasms in which this pathway is significantly dysregulated.

Aim and objectives To evaluate the preliminary clinical response of ribociclib as a single agent, in terms of best overall response (BOR) and progression free survival (PFS) in patients with Rb+ advanced solid tumours (AST) and lymphomas.

Material and methods A literature review was carried out of studies published during 2016–2019 in the electronic databases Medline, Embase and Cochrane Library. No restrictions in terms of language or publication year were applied. Search strategy terms were: ‘Ribociclib’, ‘clinical response’, ‘single agent’ and ‘advanced cancer’. Boolean operators were used to connect specific search keywords for each database and other free text terms.

Results Five clinical trials were found. A phase I study of single agent ribociclib in 132 patients from Europe and USA with Rb+ AST and lymphomas showed preliminary signs of clinical activity (NCT01237236): 3 patients achieved a partial response (PR), 43 a BOR of stable disease (SD) and 8 had PFS for >6 months. In another phase I trial in 17 Japanese patients with advanced ossephageal, breast, peritoneum and soft tissue tumours (NCT01898843), ribociclib exhibited a limited response, as no patient achieved a complete response (CR) or PR, and 4 achieved BOR on SD. In a phase I study in 32 paediatric patients with neuroblastoma and malignant rhabdoid tumours treated with single agent ribociclib (NCT01747876), BOR was SD in 9 patients and 5 achieved SD for more than 6, 6, 8, 12 and 13 cycles, respectively. The results of phase 0 and phase Ib studies that assessed the clinical response of ribociclib as monotherapy in glioblastoma (NCT02933736, NCT02345824), showed limited clinical efficacy and ineffectiveness, respectively. Both studies mentioned the presence of a significant increase in cells mTOR/PI3K signalling pathway activity.