Proton pump inhibitors may reduce the efficacy of ribociclib and palbociclib in metastatic breast cancer patients

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Abstract

Introduction

Approximately 20-33% of all cancer patients are treated with acid reducing agents (ARAs), most commonly proton pump inhibitors (PPIs), to reduce gastro esophageal reflux disease symptoms. Palbociclib and ribociclib are weak base so their solubility depends on different pH. The solubility of palbociclib dramatically decreases to <0.5 mg/ml when pH is above 4.5 but ribociclib's solubility decreases when pH increases above 6.5. In the current study, we aimed to investigate the effects of concurrent PPIs on palbociclib and ribociclib efficacy in terms of progression free survival in metastatic breast cancer (mBC) patients.

Patients and methods

We enrolled hormone receptor-positive, HER2-negative mBC patients treated with endocrine treatment (Letrozole or fulvestrant) combined palbociclib or ribociclib alone or with PPI accompanying our observational study. During palbociclib/ribociclib therapy, patients should be treated with "concurrent PPIs" defined as all or more than half of treatment with palbociclib/ribociclib, if no PPI was applied, it was defined as 'no concurrent PPI', those who used PPI but less than half were excluded from the study. All data collected from real life retrospectively.

Results

Our study included 217 patients, 105 of whom received palbociclib and 112 received ribociclib treatment. Of 105 patients who received Palbociclib, 65 were on concomitant PPI therapy, 40 were not. Of the 112 patients who received ribociclib, 61 were on concomitant PPI therapy, 51 were not. In the palbociclib group, the PFS of the patients using PPI was shorter than the PFS of the patients not using (13.04 months vs. unreachable, p<0.0001). it was determined that taking PPI was an independent predictor of shortening PFS (p<0.001) in the multivariate analysis, In the ribociclib group, the PFS of the patients using PPI was shorter than the PFS of the patients not using (12.64 months vs. unreachable, p=0.003). It was determined that taking PPI was single statistically independent predictor of shortening PFS (p=0.003, univariate analysis).

Conclusions

Our study demonstrated that concomitant usage of PPIs was associated with shorter PFS in mBC treated with both ribociclib and especially palbociclib. If it needs to be used, PPI selection should be made carefully and low-strength PPI or other ARAs (eg H2 antagonists, antacids) should be preferred.

Introduction

Targeted drugs such as tyrosine kinase inhibitors and cyclin-dependent kinase inhibitors are widely used in cancer patients. These drugs are oral medications, so, gastric pH has a significant effect on drug efficacy. There were many influencing factors on gastric pH, such as feeding and concomitant medications. These drugs can be dissolved well when the appropriate pH is established. Approximately 20-33% of all cancer
patients are treated with acid reducing agents (ARAs), most commonly proton pump inhibitors (PPIs), to reduce gastroesophageal reflux disease symptoms. PPIs also interact via the pharmacological and solubility pathways [1]. For this reason, drug-drug interactions (DDIs) at the time of absorption should be considered as one of the causes of treatment failure in cancer patients [2]. In fact, gastric pH elevation by PPIs reduces the oral bioavailability of many drugs used in cancer. This situation is demonstrated to be significant especially in those with exponentially decreasing solubility in the pH range 1-4. [3, 4]. The type of anticancer drugs determines the clinical occurrence of these changes [5]. It has been reported that long-term acid suppression by PPIs reduces the antitumor efficacy of pazopanib and capecitabine, while this effect of PPIs has not been found in clinical outcomes on patients treated with epidermal growth factor receptor (EGFR) tyrosine kinase inhibitors [6-8].

Ribociclib and palbociclib are oral cyclin-dependent kinase 4 and 6 (CDK4/6) inhibitors that arrest the cell cycle by inhibiting DNA synthesis inhibition[9]. The clinical efficacy of the combination of a CDK 4/6 inhibitor with aromatase inhibitors or fulvestrant has been acknowledged as the invention therapy in the first-second line treatment of human epidermal growth factor receptor 2 (HER2) negative hormone receptor positive premenopausal and postmenopausal women with advanced breast cancer [10-13]. Palbociclib is a weak base so its solubility depends on pH. The solubility of palbociclib dramatically decreases to <0.5 mg/ml when the pH is above 4.5 (i.e. gastric pH typically achieved by PPI). Ribociclib is also a weak base and its solubility decreases when the pH increases above 6.5. Medicines are usually taken with 200-250 ml of water. The in vitro solubility of ribociclib was investigated in biorelevant media consisting of simulated feeding (pH 5.0) and hungry (pH 6.5) intestinal fluid. The maximum dose of ribociclib (600 mg) was completely dissolved in 250 mL of biorelevant media [14]. Concomitant use of palbociclib with rabeprazole was found to reduce the mean area under the concentration time curve (AUC) – maximum plasma concentration (Cmax) 62% - 80% at fasting and 13% - 41% at fed. [15]. Clinical trial data and population pharmacokinetics showed that ribociclib absorption was similar at various stomach pH values that occur after food intake or concomitant use of PPIs [14, 16]. According to our knowledge to date, there are insufficient data on DDIs between palbociclib and PPIs other than rabeprazole.

In the current study, we aimed to investigate the effects of concurrent PPIs on palbociclib and ribociclib efficacy in terms of progression free survival in patients with estrogen-positive, HER2-negative metastatic breast cancer (mBC) treated with palbociclib/ribociclib as a first line or subsequent line of treatment.

**Patients And Methods**

We enrolled hormone receptor-positive, HER2-negative mBC patients treated with palbociclib or ribociclib alone or with PPI accompanying our observational study. Tumors with estrogen receptors in patients with metastatic breast cancer if expression is >10%, we defined hormone receptor positive as HER2-negative as a score of 0 or 1+ by immunohistochemistry and negative staining by SISH (silver in situ hybridization)/FISH (fluorescent in situ hybridization) in those with a score of 2+ in immunohistochemistry. During palbociclib/ribociclib therapy, patients were treated with "concurrent PPIs" defined as all or more than half of treatment with palbociclib/ribociclib, If no PPI was applied, it was defined as 'no simultaneous PPI'. Those who used PPIs but less than half were excluded from the study. Based on previous endocrine time response, those with endocrine sensitivity (if relapsed at least 12 months after completion of adjuvant endocrine therapy or de novo metastatic
breast cancer) or those who are endocrine resistant (relapse while receiving adjuvant therapy or recurrence within 12 months of discontinuation of adjuvant endocrine therapy) [17].

All clinicians in our study performed pharmacological and clinical interventions in real life according to clinical practice. One course of treatment was given as 28 days, consisting of 21 consecutive days of full and 7 days of blank treatment. Specifically, palbociclib orally at a dose of 125 mg, ribociclib orally at a dose of 600 mg/21 days on and 7 days off), 28-day full cycle plus fulvestrant or letrozole were administrated. Ribociclib dose reduction was made to 400 mg, and palbociclib dose reduction was made to 100 mg based on the toxicity profile. No lower dose was used in any patient. PPIs (lansoprazole 30 mg, esomeprazole 40 mg, omeprazole 40 mg, pantoprazole 40 mg, rabeprazole 20 mg dose) were recommended to take 30 minutes before breakfast. Ribociclib was used whether hungry or full, and palbociclib it was used with lunch. Strong inhibitors or inducers of cytochrome P450 3A4 (CYP3A4) while taking both drugs should be avoided. The doctors who wrote the prescription followed the patients' condition in accordance with the recommendations. Toxicity was assessed according to the World Health Organization (WHO) criteria classification. Approval was obtained from the local ethics committee in accordance with the Declaration of Helsinki, and the study was conducted accordingly.

**Statistical Analysis**

Eastern Cooperative Oncology Group (ECOG) performance status, hormone sensitivity, premenopausal versus postmenopausal status, treatment lines, visceral versus bone disease, and the number of tumor sites in absolute and median and relative frequencies and quantitative factors are categorical variables. The time from initiation of CDK combination therapy to progression was defined as PFS. For calculating PFS, generating survival curves and log-rank testing, the Kaplan-Meier method was used. Independent risk factors for PFS were determined with the Cox hazard regression method.

**Results**

Our study included 217 patients, 105 of whom received palbociclib and 112 of whom received ribociclib treatment. Of 105 patients who received palbociclib, 65 were on concomitant PPI therapy, and 40 were not. Of the 112 patients who received ribociclib, 61 were on concomitant PPI therapy, and 51 were not.

Forty-nine patients received, palbociclib combined with letrozole as first-line endocrine therapy (endocrine sensitive) and 56 endocrine refractory patients used fulvestrant as first- or subsequent line treatment. Of the patients treated with palbociclib, 49 (46.7%) received the 125 mg dose, and 56 (53.3%) received the 100 mg dose. Ribociclib was used as first line endocrine therapy (endocrine sensitive) with letrozol in 66 patients, and combined with fulvestrant as a first or subsequent line of therapy in 46 endocrine refractory patients. For patients treated with ribociclib, 51 (45.6%) patients received the 600 mg dose, and 61 (54.4%) patients received the 400 mg dose. There was no significant difference between the patients who took PPIs and those who did not in either the palbociclib group or the ribociclib group. The clinical characteristics for both drug groups separately are shown in **Table 1**.

In the palbociclib group, the PFS of the patients using PPIs was shorter than the PFS of the patients not using PPIs (13.04 months vs. unreachable, p<0.0001, respectively; Figure 1A). After a mean follow-up of 13 months, 83% of the patients who did not take PPIs did not progress. Univariate analysis included age, CDK combination,
number of metastatic sites, ECOG, menopausal status, dose reduction, metastasis diagnosis time, and CDK starting interval (CDK interval). Age, number of metastatic sites, ECOG PS, and menopausal status were found to be significantly associated with PFS (p=0.001, p=0.006, p=0.048, p=0.008, respectively; **Table 2**). As a result of multivariate analysis, it was determined that taking PPIs was an independent predictor of shortening PFS (hazard ratio 5.60; 95% confidence interval: 1.98-15.85; p=<0.001; **Table 2**). When we analysed the effective role of PPI use on PFS separately in the letrozole (hormone sensitive) and fulvestrant (hormone resistant) groups, the PFS was significantly shorter in patients using PPIs in both groups (p=0.006, p=0.021 **Figure2A, 2B**).

In the ribociclib group, the PFS of the patients using PPIs was shorter than the PFS of the patients not using PPIs (12.64 months vs. unreachable, p=0.003, respectively; **Figure 1B**). After a mean follow-up of 15 months, 65% of the patients who did not take PPIs did not progress. Univariate analysis included age, CDK combination, number of metastatic sites, ECOG, menopausal status, dose reduction, metastasis diagnosis time, and CDK starting interval (CDK interval). No statistical significance was found in any of the univariate analyses. Only PPI use was found to have a significant effect on PFS in patients receiving ribociclib (hazard ratio 2.9; 95% confidence interval: 1.38-6.40 ;p=0.003; **Table 2**). When we analysed the effective role of PPI use on PFS separately in the letrozole (hormone sensitive) and fulvestrant (hormone resistant) groups, the PFS was significantly shorter in patients using PPIs in the letrozole group (p=0.014, p=0.141 **Figure3A, 3B**).

In both the palbociclib group and the ribociclib group, there was no statistically significant difference in grade 3-4 adverse events requiring dose reduction between the patient groups taking and not taking PPIs (p=0.224, p=0.254; **Table 1**).

**Discussion**

Among different factors such as fast, feeding, concomitant drugs, gastric pH increase, etc., the pH solubility of the drug is considered to be the most relevant influencing drug absorption [18]. When stomach pH increases, the effectiveness of oral anticancer drugs with weak base properties decreases due to decreased bioavailability [3, 19]. To our knowledge, our study was the first to show that concomitant usage of PPIs with palbociclib/ribociclib in patients with mBC had a detrimental effect on PFS. We concluded that increasing gastric pH induced by PPIs may occur through lowering palbociclib plasma concentrations, which affects treatment efficacy and results in shorter progression-free survival. Palbociclib is a weak pH-dependent base with gradually increasing solubility when the pH rises above 4.5. Rabeprazole-induced changes in post-fed status on palbociclib pharmacokinetics were not considered clinically significant, and no restrictions for concomitant use of PPIs have been reported in palbociclib labelling [15, 20]. However, the clinical consequences of rabeprazole’s ability to reduce efficacy it were not investigated in the study performed by Sun et al. [15]. Additionally, while investigating the effect of rabeprazole on palbociclib pharmacokinetics, giving just 6 days may not have been enough, because in short-term treatment with PPIs, intragastric pH may not be increased throughout the 24-hour interval [13, 21]. In our study, PPIs (mainly pantoprazole, rabeprazole, esomeprazole) were given at no less than half of all palbociclib therapy for a greater and steady rise in intragastric pH.

The PFS of palbociclib with letrozole in paloma 2 and fulvestrant in paloma 3 was 27.6 months and 9.2 months, respectively, and the PFS of ribociclib with letrozole in monalisa 2 and fulvestrant in monalisa 3 was
25.3 months and 20.5 months, respectively [10-13]. According to our evaluation, the reason why PFS was lower in paloma 3 than in monalisa 3 was that some patients who received 1 step of chemotherapy in paloma 3 were included in the study, while those who received chemotherapy in monalisa 3 were not included in the study. In our study, some of the patients who received letrozole combination or fulvestrant combination had a history of chemotherapy in metastatic disease; therefore, the PFS of our study may be shorter. In the study by Re et al., PFS was 14 months versus 37.9 months in patients who received and did not receive concomitant PPIs with palbociclib, respectively. Additionally, no other significant variable affecting PFS was detected in the multivariate analysis [22]. In the results we presented, PFS was similar to that in this trial in patients who received PPIs, but PFS could not be reached yet in patients who did not receive PPIs.

When below the absolute threshold level, although it is not known at this time that the activity of palbociclib may be affected, palbociclib cell potency in vitro (IC50) with free mean steady-state concentration (Css) is comparable with a Css/IC50 ratio of 0.94 [23]. The findings of the present study support the following hypothesis: prolonged treatment with PPIs may reduce palbociclib to plasma levels below the threshold of the minimum effective concentration, thus reducing its effectiveness to some extent. Failure to evaluate the pharmacokinetic changes induced by PPIs in palbociclib is a limitation of our study. Additionally, studies have shown that short-term treatment with rabeprazole reduces fasting palbociclib Cmax by 80% and 41% at fasting and fed, respectively [15].

There is a little evidence in the literature suggesting that agents that alter gastric pH have no effect on ribociclib absorption [14, 16]. Samant et al. examined the steady-state pharmacokinetics of ribociclib (600 mg) during PPI use and found no differences in AUC and Cmax between the PPI-using and non-PPI-using groups [14]. However, that is not specified in this study is whether these patients used the drug when they were hungry or when they were full. The different behaviors of ribociclib and palbociclib in acidic media may be due to the difference in their dissolution strength. Consistent with this information, the solubility of ribociclib is >2.4 mg/ml at pH 4.5 and 0.8 mg/ml at pH>6.8, while that of palbociclib is >0.5 mg/ml at pH <4.5 only [14, 15]. Examining the in vitro solubility of ribociclib by simulating fasting intestinal fluid (pH 6.5) and postprandial intestinal fluid (pH 5.0) in biorelevant media, 600 mg was dissolved in 250 ml of fluid [14]. This feature of ribociclib makes it less affected by PPIs, but its absorption may be affected in environments where stomach acid is potently inhibited, especially in fasting conditions. Therefore, it may be more beneficial to take ribociclib with meals in patients taking ribociclib plus PPIs. Saman et al. reported that trough concentration mean ribociclib values (Ctrough) were 597 and 711 ng/ml in patients with or without PPI at 600 mg dose, respectively [14]. On average, free Css expressing a broad therapeutic index a reduction in ribociclib Ctrough is unlikely, as it greatly exceeds in vitro cell potency (Css/IC50 ratio >25) [23]. But in real life, almost half of the patients use Ribociclib at a dose of 400 mg. Therefore, Ctrough values may fall below effective levels. With respect to abemaciclib, this drug also shows pharmacokinetic similarities when compared to other CDK4/6 inhibitors. Notable features are saturable absorption with twice daily administration due to smaller volume of distribution and shorter half-life than ribociclib and palbociclib [24].

Whether P-glycoprotein (P-gp) inhibition has an effect on the PFS of PPIs observed in this study is another issue to be answered. According to our knowledge, palbociclib and ribociclib are P-gp substrates and are moderately inhibited by PPIs [25, 26]. Additionally, tyrosine kinase inhibitor (TKI) pharmacokinetics were found to be altered by pantaprazole through the influence of breast cancer resistance protein (BCRP) and P-gp [21]. If
the main mechanism of DDI is P-gp had it been inhibited by PPIs, fewer side effects would have been expected in PPI users due to the effect caused by the increase in gastric pH. In the presence or absence of PPIs, as the differences in adverse drug reactions were not statistically significant, so this hypothesis is not compatible with our data. Accordingly, rabeprazole is known to inhibit P-gp activity at appropriate concentrations, and its clinical net effect reduces palbociclib exposure [15]. However, this effect is great at fasting, in environments where the pH is higher. Therefore, gastric pH changes due to PPIs appear to be the main mechanism of interaction with drugs that require an acidic microenvironment for dissolution and absorption [27].

Studies to date have reported other instances of DDIs between PPIs and TKIs (i.e. pazopanib, sunitinib, gefitinib, and erlotinib) [7, 28-33]. A meta-analysis of 16 retrospective studies involving various solid tumors with a total of 372418 patients demonstrated that PPI therapy had a significant impact on survival outcomes in patients receiving oral anticancer drugs [34]. The effect of concomitant PPI administration on overall survival and treatment discontinuation, 90 days and 1 year after discontinuation, on overall survival in another 12 538 patients retrospective study with solid and haematological tumours evaluated. This study was performed retrospectively in patients treated with TKIs, and PPI use has been shown to be associated with an increased risk of death [35].

There were some limitations of our study. First, the adverse event profile can be underestimated because of the retrospective nature of our study. However, in the current study, dose reductions of CDK inhibitors were performed more than in other clinical trials. We generally used CDK inhibitors in the COVID-19 pandemic because the labelling time of palbociclib and ribociclib by health authorities in our country was May 2020, so physicians are sensitive to dose reduction when grade 3-4 neutropenia develops. Despite these limitations, we collected soluble and reliable data with satisfactory sample sizes. It was clearly demonstrated that concomitant usage of PPIs was associated with shorter PFS. We recommend caution in the long-term use of PPIs in this specific population and the benefits-risks of coadministration of anticancer drugs whose solubility and absorption depend on pH and strong acid-reducing agents should be evaluated and decided together. If used, PPI selection should be made carefully. For example, rabeprazole may provide more and longer acid suppression than other drugs in the same class; in treatment management H2-antagonists should also be considered instead of PPIs. Increasing the dose of palbociclib in patients using PPIs may theoretically make sense, but in clinical practice it is probably not an effective strategy due to possible off-label effects. If it is necessary to use PPI together with ribociclib, it should be used on a fed.

**Declarations**

The authors declare no competing interests.

**References**


17. Cardoso, F., et al., *4th ESO–ESMO International Consensus Guidelines for Advanced Breast Cancer (ABC 4)*† † These guidelines were developed by the European School of Oncology (ESO) and the European Society for Medical Oncology (ESMO). Annals of Oncology, 2018. 29(8): p. 1634-1657.


**Tables**

Table 1: Clinical characteristics of patients and their distribution across PPI groups.
<table>
<thead>
<tr>
<th>Characteristic</th>
<th>PALBOCICLIB</th>
<th>RIBOCICLIB</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Total number of patients (n=105)</td>
<td>Total number of patients (n=112)</td>
</tr>
<tr>
<td></td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>Age, median (Range)</td>
<td>59</td>
<td>58</td>
</tr>
<tr>
<td>Menopausal status, n(%)</td>
<td>32(30.4)</td>
<td>14(35.0)</td>
</tr>
<tr>
<td>Premenopause</td>
<td>73(69.5)</td>
<td>26(65.0)</td>
</tr>
<tr>
<td>Postmenopause</td>
<td>49(45.1)</td>
<td>34(66.7)</td>
</tr>
<tr>
<td>ECOG PS, n (%)</td>
<td>0</td>
<td>19(18)</td>
</tr>
<tr>
<td>1</td>
<td>74(70.4)</td>
<td>31(77.5)</td>
</tr>
<tr>
<td>2</td>
<td>12(11.4)</td>
<td>1(2.5)</td>
</tr>
<tr>
<td>Disease site, n (%)</td>
<td>63(60)</td>
<td>22(55.0)</td>
</tr>
<tr>
<td>Visceral</td>
<td>42(40)</td>
<td>18(45.0)</td>
</tr>
<tr>
<td>Nonvisceral</td>
<td>49(46.6)</td>
<td>23(57.5)</td>
</tr>
<tr>
<td>Endocrine therapy, n (%)</td>
<td>56(53.3)</td>
<td>17(42.5)</td>
</tr>
<tr>
<td>Letrozole</td>
<td>25(49.0)</td>
<td>36(59.0)</td>
</tr>
<tr>
<td>Fulvestrant</td>
<td>49(46.7)</td>
<td>21(52.5)</td>
</tr>
<tr>
<td>Dose Reduction, n (%)</td>
<td>65(61.9)</td>
<td>27(67.5)</td>
</tr>
<tr>
<td>Yes</td>
<td>56(53.3)</td>
<td>19(47.5)</td>
</tr>
<tr>
<td>No</td>
<td>49(46.7)</td>
<td>21(52.5)</td>
</tr>
<tr>
<td>CDK inhibitor interval</td>
<td>40(38)</td>
<td>13(32.5)</td>
</tr>
<tr>
<td>&lt;18 months</td>
<td>65(61.9)</td>
<td>27(67.5)</td>
</tr>
<tr>
<td>≥18 months</td>
<td>56(53.3)</td>
<td>19(47.5)</td>
</tr>
<tr>
<td>PPI , n (%)</td>
<td>Pantoprazole</td>
<td>28(43.1)</td>
</tr>
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Table 2. Univariate and multivariate analysis for progression-free survival.

<table>
<thead>
<tr>
<th>Medicine</th>
<th>Univariate</th>
<th>Multivariate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Esomeprazole</td>
<td>8 (12.3)</td>
<td>18 (29.5)</td>
</tr>
<tr>
<td>Lansoprazole</td>
<td>9 (13.8)</td>
<td>3 (4.9)</td>
</tr>
<tr>
<td>Omeprazole</td>
<td>3 (4.6)</td>
<td>12 (19.7)</td>
</tr>
<tr>
<td>Variables</td>
<td>PALBOCICLIB</td>
<td></td>
</tr>
<tr>
<td>------------------------------------------------</td>
<td>------------------------------</td>
<td>----------</td>
</tr>
<tr>
<td></td>
<td>Univariate</td>
<td>Multivariate</td>
</tr>
<tr>
<td></td>
<td>HR (95%) Cl</td>
<td>P value</td>
</tr>
<tr>
<td>Age</td>
<td>0.94 (0.91-0.97)</td>
<td>0.001</td>
</tr>
<tr>
<td>Number of metastatic sites</td>
<td>3.8 (1.46-10.04)</td>
<td>0.006</td>
</tr>
<tr>
<td>CDK inhibitor combination</td>
<td>1.8 (0.91-3.67)</td>
<td>0.089</td>
</tr>
<tr>
<td>ECOG PS</td>
<td>0.215 (0.04-0.98)</td>
<td>0.048</td>
</tr>
<tr>
<td>Pre/Post-menopause</td>
<td>0.394 (0.25-2.03)</td>
<td>0.008</td>
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<tr>
<td>Visseral-nonvisseral disease</td>
<td>0.58 (0.28-1.11)</td>
<td>0.130</td>
</tr>
<tr>
<td>Dose reduction</td>
<td>1.22 (0.62-2.37)</td>
<td>0.550</td>
</tr>
<tr>
<td>CDK inhibitor interval</td>
<td>1.92 (0.99-3.71)</td>
<td>0.054</td>
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<tr>
<td>Concomitant use of PPIs</td>
<td>5.60 (1.98-15.85)</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>

Figures
Figure 1

PFS curves of Palbociclib and Ribociclib combined endocrine therapy with or without PPIs. (Kaplan meier estimates) ET: endocrine treatment, PFS: progression free survival, PPI: proton pomp inhibitör, CI: confidence interval, HR: hazard ratio
Figure 2

PFS curves of Palbociclib+Al and Palbociclib+Fulvestrant with or without PPIs. (Kaplan meier estimates) Al: aromatase inhibitor, PFS: progression free survival, PPI: proton pomp inhibitör
PFS curves of Ribociclib+AI and Ribociclib+Fulvestrant with or without PPIs. (Kaplan meier estimates) AI: aromatase inhibitor, PFS: progression free survival, PPI: proton pomp inhibitör