Positive Quality Intervention: CDK4/6 Inhibitors in HR+/HER2- Advanced Breast Cancer

Description: Cyclin-dependent kinase (CDK) 4/6 inhibitors have demonstrated significant advances in the treatment of hormone receptor (HR) positive and human epidermal growth factor receptor (HER2) negative advanced breast cancer. While current available data do not support the preferential use of one agent over the other, the dosing and safety monitoring differ and may help in treatment decision making with patients. Multidisciplinary care teams have key roles in education and supportive care management to optimize the use of these agents and disease response.1,2

Background: Patients with HR+/HER2- advanced breast cancer should be considered candidates for CDK inhibitors (CDKi) therapy (palbociclib, ribociclib, and abemaciclib) due to the progression free survival (PFS) advantages demonstrated in the PALOMA, MONALEESA, and MONARCH trials, summarized in Table 2 of Supplemental Information within this PQI. A recent meta-analysis of the trials evaluating CDKi in HR+/HER2- advanced breast cancer demonstrated a 50% reduction in the rate of disease progression when CDKIs were used in the first or second line setting irrespective of age, site of metastasis, and disease free interval.5 In addition, a publication of long term follow up of PALOMA-2 reported the safety profile remained favorable and with no differences in patient reported quality of life with the addition of palbociclib to endocrine therapy over time.11 CDKIs work at CDK 4/6 resulting in the blockade of phosphorylation of the retinoblastoma (Rb) protein which hinders the activation of transcription factors involved in S-phase entry, arresting the cell cycle progression at G1 phase.3

PQI Process: Upon receipt of an order for a CDK inhibitor:

- Ensure the patient has HR+/HER2- advanced breast cancer
- If patient had recently been on endocrine only therapy and experienced disease recurrence or progression, ensure endocrine therapy was changed along with prescribing of CDKi. New hormonal therapy can be started while access to new CDKi is in process
- There is no data to support continued CDKi use following disease progression; although switching from one to another due to tolerability is appropriate
- Assessment of drug interaction should be reviewed at baseline and throughout therapy as new medications are prescribed
  - CDKi are cytochrome P450 3A4 substrates. Medication lists should be reviewed and alternatives discussed for strong inhibitors/inducers. The use of moderate CYP3A4 inducers/inhibitors should be reviewed and adjustments in dose and/or monitoring discussed with patients and providers
- Due to the high rate of dose interruptions, care coordination is critical to ensure laboratory monitoring and assessments are occurring at appropriate times to avoid unnecessary holds while ensuring the safe monitoring. If a cycle is delayed, future labs should be adjusted from the true start of the current cycle
- A 7 day follow up should be conducted for all CDKis, especially abemaciclib for diarrhea management
- Midcycle CBC is recommended for ribociclib and palbociclib
- EKG is recommended every 2 weeks for ribociclib for the first 2 cycles

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Patient Centered Activities:

- Provided Oncology Chemotherapy Education (OCE) Sheet for CDKi and hormonal therapy to patient
- Ensure patients are aware their hormonal therapy does not stop when holding CDKi therapy
- Provide patient expectations of therapeutic assessment and laboratory monitoring when starting therapy, especially during first 8 weeks. Review more than half of patients will require dose interruption or reduction, although ADRs generally resolve rapidly
- Review with patients as they require dose reductions there is no dose response established with CDKi. Regardless of dose, patients can continue to experience efficacy
- Refer to *Table 1 of Supplemental Information within this PQI* for drug-specific dosing, monitoring, and counseling
- Provided instructions for use of loperamide and potentially provide a prescription for diphenoxylate/atropine for patients prescribed abemaciclib
  - Provide antidiarrheal management guidance per *Abemaciclib (Verzenio) Diarrhea Management PQI*
- All medications should be stored in a cool dry place at room temperature. If dispensed in blister packaging, CDKis should stay in original packaging
- Take medication at approximately the same time each day, do not double doses if one is forgotten
- **Patient assistance:**
  - Abemaciclib: [https://www.verzenio.com/savings-support/savings-card](https://www.verzenio.com/savings-support/savings-card)
  - Palbociclib: [https://www.ibrance.com/financial-support-resources](https://www.ibrance.com/financial-support-resources)
  - Ribociclib: [https://www.copay.novartisoncology.com/?name=kisqali](https://www.copay.novartisoncology.com/?name=kisqali)

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# Table 1: Dosing and Monitoring Guideline Summary

<table>
<thead>
<tr>
<th></th>
<th>Palbociclib</th>
<th>Ribociclib</th>
<th>Abemaciclib</th>
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<tbody>
<tr>
<td><strong>FDA Indication</strong></td>
<td>HR+/HER2- advanced or metastatic breast cancer in combination with</td>
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<td>• \textit{An AI} as initial endocrine-based therapy in postmenopausal</td>
<td>• \textit{An AI} as initial endocrine-based therapy for pre/peri or</td>
<td>• \textit{An AI} as initial endocrine-based therapy in postmenopausal</td>
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<td></td>
<td>women/men</td>
<td>postmenopausal women/men</td>
<td>women/men</td>
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<tr>
<td></td>
<td>• Fulvestrant in adults with disease progression following endocrine</td>
<td>Fulvestrant as initial endocrine-based therapy or following disease</td>
<td>Fulvestrant in adults with disease</td>
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<tr>
<td></td>
<td>therapy</td>
<td>progression following endocrine therapy</td>
<td>progression following endocrine therapy</td>
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<tr>
<td></td>
<td></td>
<td>for postmenopausal women</td>
<td>• Monotherapy after progression on ET and chemo in metastatic setting</td>
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<tr>
<td><strong>Dosing</strong></td>
<td>125 mg daily with food 21 days on/7 days off + continuous ET</td>
<td>600 mg daily qam without regards to meals 21 days on/7 days off + continuous ET</td>
<td>200 mg twice daily monotherapy OR 150 mg twice daily without regards to meals + continuous ET</td>
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<td><strong>Monitoring</strong></td>
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<td>CBC w/diff:</td>
<td>D 1&amp;15 cycles 1&amp;2, then prior to each cycle x 4, then Q3 months* if no grade 3/4 neutropenia within first 6 months</td>
<td></td>
<td>D 1&amp;15 cycles 1&amp;2, then prior to each cycle x 2</td>
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<tr>
<td>LFTs/Tbili:</td>
<td>NA</td>
<td>Every 2 weeks x 4, then monthly x 4, then every 3 months or as clinically indicated</td>
<td>Every 2 weeks x 4, then monthly x 2 (same as CBC), then every 3 months or as clinically indicated</td>
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<tr>
<td>Electrolytes (BMP)</td>
<td>NA</td>
<td>Monthly x 6</td>
<td>NA</td>
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<tr>
<td>EKG:</td>
<td>NA</td>
<td>Every 2 weeks x 3; &lt;450msec to initiate, refer to PI for management after initiation</td>
<td>NA</td>
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<td>Neutropenia Management</td>
<td>\textit{Day 1 of each cycle} must have ANC $\geq$ 1.0 to start \textit{D15 first 2 cycles}: ANC 0.5-1.0 continue and recheck day 22; if &lt;0.5 at day 22: start next cycle at reduced dose once improves to grade 2. At any time: ANC$&lt;1.0$ with fever or &lt;0.5: hold, resume at reduced dose once returns to grade $\geq2$</td>
<td>\textit{Day 1 of each cycle}: must have ANC $\geq$ 1.0 to start. Interrupt therapy until recovered, then resume at same dose if grade 3 ANC was experienced on Day 1 or at reduced dose if grade 4 or recurrent grade 3</td>
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<td>Hepatobiliary Management (onset: 2-6 months)</td>
<td>N/A</td>
<td>LFTs&gt;3-5xULN AND Tbili&lt;2xULN: interrupt until returns to baseline, restart at same dose; If recurrence: resume at dose reduction** LFTs&gt;5-20xULN AND Tbili&lt;2xULN: interrupt therapy until return to baseline, restart at dose reduction; Grade 3 recurrence, discontinue. LFTs&gt;20xULN: Discontinue</td>
<td>LFTs &gt;3-5xULN AND Tbili&lt;2xULN: no interruption; if recurrence: consider interruption and dose reduction** LFTs&gt;5-20xULN AND Tbili&lt;2xULN: interrupt therapy until resolution to baseline or grade 1, restart at dose reduction LFTs&gt;20xULN: Discontinue</td>
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<td><strong>Dose Reductions</strong></td>
<td>125 mg→100 mg→75 mg daily 21 days on/7 days off 75 mg 14 days on/14 days off has been reported</td>
<td>600 mg→400 mg→200 mg daily 21 days on/7 days off</td>
<td>200 mg→150 mg→100 mg→50 mg twice daily continuously (200 mg if monotherapy)</td>
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<td><strong>Adjustments at treatment initiation</strong></td>
<td>Hepatic: Child Pugh Class C: reduce to 75 mg daily 21 days on/7 days off</td>
<td>Hepatic: Child Pugh Class B or C: reduce to 400 mg daily 21 days on/7 days off</td>
<td>Hepatic: Child Pugh Class C: reduce to once daily dosing</td>
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<tr>
<td>Renal: Not studied &lt;15ml/min, not anticipated to impact metabolism</td>
<td>Renal: eGFR 15-30ml/min reduce to 200 mg; not studied &lt;15ml/min</td>
<td>Renal: Not studied &lt;30ml/min, not anticipated to impact metabolism. SCr increases anticipated after initiation, not anticipated to reflect change in renal function.¹ Assess along with severity of diarrhea</td>
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<td><strong>Drug-Drug Interaction:</strong> Strong CYP3A4 inhibitors and inducers should be avoided, including grapefruit. Moderate CYP3A4 inhibitors and inducers should be discussed with prescribers</td>
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<td><strong>Assessment to review:</strong></td>
<td>Fevers, chills, dizziness; SOB, weakness; Unusual bruising or bleeding; chest pain, tachypnea, tachycardia; adherence; laboratory values</td>
<td>At week 1: adherence and ensuring day 15 labs to be complete</td>
<td>At week 1: N/V, diarrhea, utilization of antidiarrheals</td>
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*grade 3 anemia can occur late, therefore continued periodic monitoring is recommended

**dose reductions are not warranted if baseline liver function tests are grade 2 at baseline; grade 2 AST/ALT with Tbili elevation (in absence of cholestasis) therapy should be discontinued
<table>
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<tr>
<th>Trial</th>
<th>Patient Population</th>
<th>Description</th>
<th>Outcome</th>
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| Paloma-1/Trio-18⁶,⁷ | Phase II 1st line Post-menopausal HR+/HER2- | Letrozole vs Letrozole + Palbociclib               | ORR: 39% vs 55%  
PFS: 10.2 vs 20.2 months  
OS: 33.3 vs 37.5 months* |
| Paloma-2⁴,¹⁰,¹¹ | Phase III 1st line Post-menopausal HR+/HER2- | Letrozole + Placebo vs Letrozole + Palbociclib    | ORR: 44% vs 55%  
PFS: 14.5 vs 27.6 months |
| Paloma-3¹⁶,¹⁷ | Phase III 2nd line or later Post- menopausal HR+/HER2- | Fulvestrant + placebo vs Fulvestrant + Palbociclib | PFS 4.6 vs 9.5 months  
OS: 28.0 vs 34.9 months  
OS: 29.7 vs 39.7 months (prior sensitivity to ET) |
| Monaleesa- 2¹⁴ | Phase III 1st line Post-menopausal HR+/HER2- | Letrozole + Placebo vs Letrozole + Ribociclib    | ORR: 39% vs 55%  
PFS: 16 vs 25.3 months |
| Monaleesa- 7²² | Phase III 1st line Pre/peri menopausal HR+/HER2- | OFS/AI or Tamoxifen + Ribociclib vs OFS/AI or Tamoxifen + Placebo | ORR: 36% vs 51%  
PFS: 13.8 vs 27.5 months  
OS: 47.7 vs 58.7 months |
| Monaleesa- 3¹⁹ | Phase III 1st line or 2nd line HR+/HER2- | Fulvestrant + Placebo vs Fulvestrant + Ribociclib | ORR: 36% vs 51%  
PFS: 18.3 vs NR*1st line ORR: 19 vs 41%  
PFS: 12.8 vs 20.5 months  
OS: 40 months vs NOT YET REACHED |
| MONARCH- 1²⁰ | Phase II 3rd line or later HR+/HER2- | Abemaclicib monotherapy                          | ORR: 20%  
PFS: 6.0 months  
OS: 22 months |
| MONARCH- 2²¹ | Phase III 2nd line** Pre/Peri/Post menopausal HR+/HER2- | Fulvestrant + Placebo vs Fulvestrant + abemaciclib | ORR: 21 vs 48%  
PFS: 9.3 vs 16.4 months  
OS: 37.3 vs 46.7 months |
| MONARCH-3¹⁸ | Phase III First Line Post-menopausal HR+/HER2- | NSAI + Placebo vs NSAI + Abemaciclib             | ORR: 44 vs 59%  
PFS: 14.7 vs 28.18 months |

*not statistically significant, potentially due to small population  
**Progression during neo-adjuvant/adjuvant ET, <12months from end of adjuvant ET or during 1st line ET for advanced breast cancer treatment
References:
4. Shah M, Nunes MR, Stearns V. CDK4/6 inhibitors: game changers in the management of hormone receptor-positive advanced breast cancer?