



Criteria	Checklist
<p>The application meets the following general application criteria:</p> <ul style="list-style-type: none"> <li>• The COB evaluation package is complete (including the original dossier submitted to the COB), in English (or accompanied by a certified translation to English) and un-redacted.</li> <li>• The COB assessment package must have received full marketing approval following a <i>de novo</i> assessment made by a COB, and presents an independent assessment of data provided to that COB.</li> <li>• The application must not have been delayed, deferred, rejected, refused or withdrawn at any time, in any country.</li> <li>• The application should not be subject to any further restrictions or conditions following approval, that have not been identified in the report.</li> <li>• The COB evaluation report is not subject to any restrictions on use or disclosure by the TGA.</li> </ul>	<p>Yes <input type="checkbox"/></p> <p>No <input type="checkbox"/> not eligible for the COB report-based process.</p>
<p>For dossiers that include proprietary information from a third party, evidence to confirm that all proprietary information, including the COB evaluation and relevant communication between the third party and the COB, has been provided to the TGA.</p>	<p>Yes <input type="checkbox"/></p> <p>No <input type="checkbox"/> not eligible for the COB report-based process</p> <p>N/A <input type="checkbox"/></p>
<p>Nomenclature:</p> <ul style="list-style-type: none"> <li>• Does the COB report use <a href="#">Australian approved terminology</a> (e.g. AANs, ABNs, dosage forms)?</li> </ul>	<p>Yes <input type="checkbox"/></p> <p>No <input type="checkbox"/> provide further information.</p>
<p><b>Indications</b></p>	
<p>The proposed indications are equivalent to those approved by the COB, including equivalent dosing and administration details, directions for use, target population demographics, disease profiles, expected health outcomes and intent and meaning.</p>	<p>No <input type="checkbox"/> not eligible for the COB report-based process.</p> <p>Yes <input type="checkbox"/> specify approved indications in COB report.</p>
<p>Are any indications proposed beyond those approved by the COB?</p>	<p>Yes <input type="checkbox"/> Not eligible for the COB report-based process.</p> <p>No <input type="checkbox"/></p>

Criteria	Checklist
<p>For generic medicines:</p> <p>The proposed indications are identical to the indications approved for the Australian originator product.</p>	<p>Yes <input type="checkbox"/></p> <p>No <input type="checkbox"/> not eligible for the COB report-based process.</p>
<b>Medicine characteristics</b>	
<p>The proposed medicine is identical to that evaluated and approved within the COB assessment package with regard to all of the below:</p> <ul style="list-style-type: none"> <li>· Dosage form</li> <li>· Strengths</li> <li>· Formulation/s (active substance and excipients)</li> <li>· Quality aspects</li> </ul>	<p>Yes <input type="checkbox"/></p> <p>No <input type="checkbox"/> not eligible for the COB report-based process.</p>
<p>Are the manufacturers and manufacturing process (drug substance and drug product, including finished product container) identical to that evaluated and approved within the COB assessment package?</p>	<p>Yes <input type="checkbox"/></p> <p>No <input type="checkbox"/> specify and provide details.</p>
<p>Are additional manufacturing sites nominated in the application for Australia?</p>	<p>No <input type="checkbox"/></p> <p>Yes <input type="checkbox"/> specify and provide details.</p>
<b>Generic medicines</b>	
<p>Is your Australian application for a generic medicine?</p>	<p>No <input type="checkbox"/> skip to next section.</p> <p>Yes <input type="checkbox"/> complete the generic medicines section below.</p>
<p>Was the reference medicine used in the comparative studies in the COB dossier a medicine currently listed on the ARTG and sourced from Australia?</p>	<p>No <input type="checkbox"/> continue to next question.</p> <p>Yes <input type="checkbox"/> Insert ARTG number:</p>
<p>Did the dossier submitted to the COB include biopharmaceutic data?</p>	<p>Yes <input type="checkbox"/> continue to next question.</p> <p>No <input type="checkbox"/> specify and provide justification for not supplying biopharmaceutic data.</p>
<p>Is evidence provided that the reference product used in any evaluation of bioequivalence is identical to the Australian reference product?</p>	<p>Yes <input type="checkbox"/></p> <p>No <input type="checkbox"/> not eligible for the COB report-based process.</p>

Criteria	Checklist
<b>International alignment of guidelines and standards</b>	
The COB evaluation report cites international guidelines and standards consistent with those adopted by the TGA.  See <a href="#">Clinical Efficacy and Safety Guidelines</a>	Yes <input type="checkbox"/> No <input type="checkbox"/> provide justification where COB guidelines do not align with TGA adopted guidelines and submit a full copy of the relevant COB guidelines.
Have country-specific guidelines cited in the COB evaluation report that are not adopted by the TGA been justified?	Yes <input type="checkbox"/> specify where in your application No <input type="checkbox"/> specify and provide a justification N/A <input type="checkbox"/>
The COB evaluation report is consistent with the methodology used by the TGA e.g. contains information consistent with the Common Technical Document (CTD)	Yes <input type="checkbox"/> No <input type="checkbox"/> provide details and justification
The COB evaluation report contains all information necessary to meet the TGA's technical requirements.	Yes <input type="checkbox"/> No <input type="checkbox"/> provide details and justification
<b>Currency of clinical studies</b>	
Are any updates to pivotal studies or supportive studies available that were not considered in the COB approval supporting the proposed indications.	No <input type="checkbox"/> Yes <input type="checkbox"/> provide details.
Is there any additional information available relevant to the risk-benefit of the overseas medicine since COB approval?	No <input type="checkbox"/> Yes <input type="checkbox"/> provide details.
Have there been any variations or updates to the overseas medicine since it was evaluated by the COB (e.g. directions for use, ingredients, indications, label warning statements).	No <input type="checkbox"/> Yes <input type="checkbox"/> provide details.

By completing and signing this form I give permission for the TGA to contact the above-mentioned COB and share information relating to my application.

Name

--

Signature

	Date	
--	------	--