FAIR TRADING ACT 2010

The following instrument is published under the *Fair Trading Act 2010* section 21


I, General the Honourable David Hurley AC DSC (Retd), Governor General of the Commonwealth of Australia, acting with the advice of the Federal Executive Council, make the following regulations.


David Hurley,
Governor General.

By His Excellency’s Command

Karen Andrews
Minister for Industry, Science and Technology.
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Schedule 1—Amendments

Competition and Consumer Regulations 2010
1 Name

This instrument is the *Competition and Consumer Amendment (Australian Consumer Law—Country of Origin Representations) Regulations 2020*.

2 Commencement

(1) Each provision of this instrument specified in column 1 of the table commences, or is taken to have commenced, in accordance with column 2 of the table. Any other statement in column 2 has effect according to its terms.

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<th>Commencement information</th>
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<tr>
<td><strong>Column 1</strong></td>
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<td>Provisions</td>
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<td>1. The whole of this instrument</td>
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Note: This table relates only to the provisions of this instrument as originally made. It will not be amended to deal with any later amendments of this instrument.

(2) Any information in column 3 of the table is not part of this instrument. Information may be inserted in this column, or information in it may be edited, in any published version of this instrument.

3 Authority

This instrument is made under section 139G of the *Competition and Consumer Act 2010*.

4 Schedules

Each instrument that is specified in a Schedule to this instrument is amended or repealed as set out in the applicable items in the Schedule concerned, and any other item in a Schedule to this instrument has effect according to its terms.
Schedule 1—Amendments

_Competition and Consumer Regulations 2010_

1 Regulation 92AA

Repeal the regulation, substitute:

92AA Process substantially transforming medicines in Australia

(1) For the purposes of paragraph 255(2)(c) of the Australian Consumer Law, this regulation prescribes a process that medicines have undergone in Australia to be substantially transformed in Australia.

(2) This regulation applies to medicines that are complementary medicines (within the meaning of the _Therapeutic Goods Regulations 1990_) and are either:
   (a) listed goods; or
   (b) registered goods.

(3) The process is the carrying out of the last step (except one covered by subregulation (4)) in the manufacture of the dosage form of medicines that:
   (a) occurs at premises in Australia; and
   (b) is authorised by a licence to occur in relation to those medicines at those premises.

(4) This subregulation covers the following steps:
   (a) covering of the dosage form of medicines in containers;
   (b) packaging of the dosage form of medicines;
   (c) labelling of the dosage form of medicines;
   (d) storage of the dosage form of medicines (whether in packaging or not);
   (e) testing of the dosage form of medicines;
   (f) release for supply of the dosage form of medicines.

(5) A term (except “process”) used in this regulation and the _Therapeutic Goods Act 1989_ has the same meaning in this regulation as it has in that Act.

Note: Terms whose meaning is affected include “containers”, “dosage form”, “labelling”, “licence”, “listed goods”, “manufacture”, “medicines”, “packaging”, “premises”, “registered goods”, “release for supply”, “storage” and “testing.”