



# Medical Device Application

## Class IIb Status : Approved

### Application Change history

<b>Application Progress Date</b>	
Date received:	09/04/2005
<b>Review Information</b>	
Review flag:	
Auto review required:	No
<b>ARTG &amp; Product ID</b>	
ARTG ID	118430
Product ID	199040
<b>Application Details</b>	
Application identifier:	DV-20050405-DA-002081-2
Submission identifier:	DV-2005-1454
Sponsor's own reference:	hp/eska/ genius hip surface replacement 01
Application for:	
Will you be applying for listing of this product or procedure in the Medicare Benefit Schedule (MBS)?	<input type="radio"/> Yes <input type="radio"/> No
Will you be applying for listing of this product on the Prosthesis List?	<input type="radio"/> Yes <input type="radio"/> No
Will you be applying for listing of this product on the Co-dependent or hybrid technology application list?	<input type="radio"/> Yes <input type="radio"/> No
Cancel ARTG - product:	
<b>Sponsor Details</b>	
Agent name:	SPECTRUM TECHNOLOGY PTY LTD
Sponsor name:	Eska Australia
Contact details:	

Contact email:

### Class Details

Class: Class IIb

Intended purpose: A device used to resurface the articulating surfaces of the femoral head and acetabulum

### Device Product Characteristics

Is the device, or any form of the device, supplied sterile: Yes

Sterilisation Method:

Is the device intended to be invasive: Yes

Is the device, or any form of the device, intended for single use: Yes

Is the device an active device: No

Does the device contain material or ingredients of microbial origin: No

Does the device contain material or ingredients of recombinant origin: No

Does the device contain material or ingredients manufactured or formulated using a genetically modified organism: No

Does the device contain material or ingredients of Human Origin: No

Does the device contain Human Blood or its components: No

Does the device consist of: Single product only

Does the device contain material or ingredients of Animal Origin rendered non-viable: No

Animal Species:

Country of Origin:

Does any component in the procedure, kit or system contain material or ingredients of Animal Origin rendered non-viable: No

Is the device medicated: No

Is the device formulated: No

Does the product contain a medicine that is supplied separately in the Australian Market: No

Does the product contain a medical device which incorporates a medicine as an integral part and that has an action ancillary to the device: No

Does the device contain a metal on metal bearing:

I declare that this device contains only components that are medical devices which have been individually certified. No

### Manufacturer Details

Manufacturer evidence number: DV-20050329-MC-001787-2 : hp/eska implants/various 01

Manufacturer name: Eska Implants GmbH and Co (Germany)[45325]

Manufacturer address as on evidence:	GrapengieBerstraBe 34 Lubeck D-23556 Germany S [ 147473]

<b>GMDNS Code and Description</b>
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GMDNS code and description:	Prosthesis, internal, joint, hip, resurfacing[33717]
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<b>Device Category Terms</b>
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Device category 1:	Non-active implantable devices
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Device category 2:	
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Device category 3:	
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<b>Product Details</b>
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UPI (Unique product identifier):	
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Total number of devices covered:	
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Functional decription:	
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<b>Variant List</b>
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#	Variant type	Variant range
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<b>Standard Conditions</b>
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<b>Non Standard Conditions</b>
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Note: A non standard conditions must not contain semi colons.
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To remove, enter item #
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<b>Declaration</b>
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- (a) devices of the kind in question are medical devices; and
- (b) devices of that kind are intended for a specified purpose, as ascertained under The definition of a medical device; and
- (c) the kind of device is correctly classified according to the medical device classifications; and
- (d) devices of that kind comply with the essential principles; and
- (e) I:
- (i) have available sufficient information to substantiate that compliance with the essential principles; or

- (ii) have procedures in place, to ensure that such information can be obtained from the manufacturer within the period specified in the regulations; and
- (f) an appropriate conformity assessment procedure has been applied to devices of that kind; and
- (g) I:
- (i) have available sufficient information to substantiate the application of those conformity assessment procedures; or
- (ii) have procedures in place, to ensure that such information can be obtained from the manufacturer within the period specified in the regulations; and
- (h) devices of that kind comply with every requirement (if any) relating to advertising applicable under the regulations; and
- (i) devices of that kind do not contain substances that are prohibited imports for the purposes of the Customs Act 1901; and
- (ia) devices of that kind are not to be used exclusively for one or more of the purposes specified under section 41BEA; and
- (j) the information included in or with the application is complete and correct.

I understand the consequences of making a false declaration, as outlined below.

In electronically submitting this application to TGA, I hereby declare that in relation to this medical device the information given in this application and the above statements on this declaration form are current and correct.

**PLEASE NOTE:**

A false declaration will result in the device entry being removed/cancelled from the ARTG.

Signatory name of the person submitting the application.:

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History
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14/04/2005 10:49:34 AM Approved.

Review Completed - Accepted, 14/04/2005)

Record	Date
Fee 670	Date Paid 12/04/2005
	Date Decision 14/04/2005

Start Dates	Finish Dates	Working Days
Application Received 09/04/2005	Payment Received 12/04/2005	1
Payment Received 12/04/2005	Application Decision 14/04/2005	3
Total Working Days		4