

Health Insurance Legislation Amendment (2018 Measures No. 1) Regulations 2018

I, General the Honourable Sir Peter Cosgrove AK MC (Ret’d), Governor‑General of the Commonwealth of Australia, acting with the advice of the Federal Executive Council, make the following regulations.

Dated 19 April 2018

Peter Cosgrove

Governor‑General

By His Excellency’s Command

Greg Hunt

Minister for Health

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1 Name

 This instrument is the *Health Insurance Legislation Amendment (2018 Measures No. 1) Regulations 2018*.

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.

| Commencement information |
| --- |
| Column 1 | Column 2 | Column 3 |
| Provisions | Commencement | Date/Details |
| 1. The whole of this instrument | 1 May 2018. | 1 May 2018 |

Note: This table relates only to the provisions of this instrument as originally made.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 the *Health Insurance Act 1973.*

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 Each instrument that is specified in a Schedule to this instrument is amended or repealed as set out in the applicable items in the Schedule concerned, and any other item in a Schedule to this instrument has effect according to its terms.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

Health Insurance (Diagnostic Imaging Services Table) Regulations 2017

1 Subclause 2.4.2(1) of Schedule 1

Omit “61646”, substitute “61647”.

2 Schedule 1 (item 61369)

Repeal the item, substitute:

|  |  |  |
| --- | --- | --- |
| 61369 | Indium‑labelled octreotide study (including single photon emission tomography when undertaken), if:(a) a gastro‑entero‑pancreatic endocrine tumour is suspected on the basis of biochemical evidence with negative or equivocal conventional imaging; or(b) both:(i) a surgically amenable gastro‑entero‑pancreatic endocrine tumour has been identified on the basis of conventional techniques; and(ii) the study is to exclude additional disease sites(R) (K) | 2,015.75 |

3 Schedule 1 (after item 61646)

Insert:

|  |  |  |
| --- | --- | --- |
| 61647 | Whole body 68Ga‑DOTA‑peptide PET study (including any associated computed tomography scans for anatomic localisation and attenuation correction), if:(a) a gastro‑entero‑pancreatic neuroendocrine tumour is suspected on the basis of biochemical evidence with negative or equivocal conventional imaging; or(b) both:(i) a surgically amenable gastro‑entero‑pancreatic neuroendocrine tumour has been identified on the basis of conventional techniques; and(ii) the study is for excluding additional disease sites(R) | 1,053.00 |

4 Schedule 1 (item 61671)

Repeal the item, substitute:

|  |  |  |
| --- | --- | --- |
| 61671 | Indium‑labelled octreotide study (including single photon emission tomography when undertaken), if:(a) a gastro‑entero‑pancreatic endocrine tumour is suspected on the basis of biochemical evidence with negative or equivocal conventional imaging; or(b) both:(i) a surgically amenable gastro‑entero‑pancreatic endocrine tumour has been identified on the basis of conventional techniques; and(ii) the study is to exclude additional disease sites(R) (NK) | 1,007.90 |

5 Subclause 2.5.1(1) of Schedule 1

Omit “Items 63001 to 63467, 63487 to 63490, 63470 to 63486 and 63740 to 63747”, substitute “The items in Subgroups 1 to 21”.

6 Subclause 2.5.1(2) of Schedule 1

Omit “Items 63457, 63458, 63464 to 63467, 63487 to 63490, 63470 to 63484 and 63740 to 63747”, substitute “Items 63395 to 63398 and the items in Subgroups 19, 20 and 21 (other than items 63455 and 63461)”.

7 Subclause 2.5.1(3) of Schedule 1

Omit “Items 63491 to 63497”, substitute “The items in Subgroup 22”.

8 Subclause 2.5.1(4) of Schedule 1

Omit “Items 63507 to 63561”, substitute “The items in Subgroups 33 and 34”.

9 Clause 2.5.4 of Schedule 1

Repeal the clause, substitute:

2.5.4 MRI and MRA services—eligible provider

 A person mentioned in column 2 of an item of the following table is an ***eligible provider*** for an MRI or MRA service mentioned in column 1 of the item.

| Eligible providers |
| --- |
| Item | Column 1MRA or MRA service | Column 2Person |
| 1 | A service to which none of items 63395 to 63398 apply | A person who:(a) is a specialist in diagnostic radiology; and(b) satisfies the Chief Executive Medicare that the specialist is a participant in the Royal Australian and New Zealand College of Radiologists’ Quality and Accreditation Program |
| 2 | A service to which any of items 63395 to 63398 apply | A person who is:(a) a specialist in diagnostic radiology or a consultant physician; and(b) recognised by the Conjoint Committee for Certification in Cardiac MRI |

10 Clause 2.5.9 of Schedule 1

Repeal the clause, substitute:

2.5.9 MRI or MRA services—application of items to related services provided in same period

 An MRI or MRA item does not apply to a service provided to a person if:

 (a) the MRI or MRA item is specified in column 1 of an item (the ***table item***) of the following table; and

 (b) during the period (the ***limitation period***):

 (i) specified in column 2 of the table item; and

 (ii) ending immediately before the service is provided;

 the person was provided with one or more services (the ***earlier services***) to which any of the MRI or MRA items mentioned in the table item applied; and

 (c) the number of earlier services provided to the person in the limitation period was equal to the maximum number specified in column 3 of the table item.

| Related services |
| --- |
| Item | Column 1MRI or MRA items | Column 2Limitation period | Column 3Maximum number of services |
| 1 | 63040 to 63085 | 12 months | 3 |
| 2 | 63101 and 63104 | 12 months | 3 |
| 3 | 63125 to 63136 | 12 months | 3 |
| 4 | 63161 to 63194 | 12 months | 3 |
| 5 | 63219 to 63265 | 12 months | 3 |
| 6 | 63271 to 63285 | 12 months | 3 |
| 7 | 63322 to 63348 | 12 months | 3 |
| 8 | 63361 and 63364 | 12 months | 2 |
| 9 | 63385 to 63394 | 12 months | 2 |
| 10 | 63395 and 63396 | 12 months | 1 |
| 11 | 63397 and 63398 | 36 months | 1 |
| 12 | 63401 to 63408 | 12 months | 3 |
| 13 | 63416 and 63419 | 12 months | 1 |
| 14 | 63425 to 63433 | 12 months | 2 |
| 15 | 63455 to 63467 | 12 months | 1 |
| 16 | 63547 and 63548 | patient’s lifetime | 1 |
| 17 | 63482 and 63486 | 12 months | 3 |
| 18 | 63507 to 63523 and 63551 to 63561 | 12 months | 3 |

11 Schedule 1 (at the end of Subgroup 14 of Group I5)

Add:

|  |  |  |
| --- | --- | --- |
| 63395 | MRI—scan of cardiovascular system for assessment of myocardial structure and function involving:(a) dedicated right ventricular views; and(b) 3D volumetric assessment of the right ventricle; and(c) reporting of end‑diastolic and end‑systolic volumes, ejection fraction and BSA‑indexed values;if the request for the scan indicates that:(d) the patient presented with symptoms consistent with arrhythmogenic right ventricular cardiomyopathy (ARVC); or(e) investigative findings in relation to the patient are consistent with ARVC(R) (K) (Anaes.) (Contrast) | 855.20 |
| 63396 | MRI—scan of cardiovascular system for assessment of myocardial structure and function involving:(a) dedicated right ventricular views; and(b) 3D volumetric assessment of the right ventricle; and(c) reporting of end‑diastolic and end‑systolic volumes, ejection fraction and BSA‑indexed values;if the request for the scan indicates that:(d) the patient presented with symptoms consistent with arrhythmogenic right ventricular cardiomyopathy (ARVC); or(e) investigative findings in relation to the patient are consistent with ARVC(R) (NK) (Anaes.) (Contrast) | 427.60 |
| 63397 | MRI—scan of cardiovascular system for assessment of myocardial structure and function involving:(a) dedicated right ventricular views; and(b) 3D volumetric assessment of the right ventricle; and(c) reporting of end‑diastolic and end‑systolic volumes, ejection fraction and BSA‑indexed values;if the request for the scan indicates that the patient:(d) is asymptomatic; and(e) has one or more first degree relatives diagnosed with confirmed arrhythmogenic right ventricular cardiomyopathy (ARVC)(R) (K) (Anaes.) (Contrast) | 855.20 |
| 63398 | MRI—scan of cardiovascular system for assessment of myocardial structure and function involving:(a) dedicated right ventricular views; and(b) 3D volumetric assessment of the right ventricle; and(c) reporting of end‑diastolic and end‑systolic volumes, ejection fraction and BSA‑indexed values;if the request for the scan indicates that the patient:(d) is asymptomatic; and(e) has one or more first degree relatives diagnosed with confirmed arrhythmogenic right ventricular cardiomyopathy (ARVC)(R) (NK) (Anaes.) (Contrast) | 427.60 |

12 Schedule 1 (at the end of Subgroup 19 of Group I5)

Add:

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| --- | --- | --- |
| 63547 | MRI—scan of both breasts for the detection of cancer, if:(a) a dedicated breast coil is used; and(b) the request for scan identifies that:(i) the patient has a breast implant in situ; and(ii) anaplastic large cell lymphoma has been diagnosed(R) (K) (Anaes.) (Contrast) | 690.00 |
| 63548 | MRI—scan of both breasts for the detection of cancer, if:(a) a dedicated breast coil is used; and(b) the request for scan identifies that:(i) the patient has a breast implant in situ; and(ii) anaplastic large cell lymphoma has been diagnosed(R) (NK) (Anaes.) (Contrast) | 345.00 |

13 Clause 3.1 of Schedule 1 (definition of *scan*)

Omit “63567”, substitute “63561 and 63740 to 63747”.

Health Insurance (General Medical Services Table) Regulations 2017

14 Subclause 1.2.5(1) of Schedule 1

After “11724,”, insert “11728,”.

15 Schedule 1 (at the end of Subgroup 6 of Group D1)

Add:

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| --- | --- | --- |
| 11728 | Implanted loop recording for the investigation of atrial fibrillation if the patient to whom the service is provided has been diagnosed as having had an embolic stroke of undetermined source, including reprogramming when required, retrieval of stored data, analysis, interpretation and report, other than a service to which item 38288 appliesFor any particular patient—applicable not more than 4 times in any 12 months | 34.75 |

16 Schedule 1 (item 15565, column headed “Description”, subparagraphs (b)(iii) and (iv))

Repeal the subparagraphs, substitute:

(iii) validating the accuracy of the derived IMRT dosimetry plan; and

17 Schedule 1 (items 30481 to 30483)

Repeal the items, substitute:

|  |  |  |
| --- | --- | --- |
| 30481 | Percutaneous gastrostomy (initial procedure):(a) including any associated imaging services; and(b) excluding the insertion of a device for the purpose of facilitating weight loss(Anaes.) | 357.00 |
| 30482 | Percutaneous gastrostomy (repeat procedure):(a) including any associated imaging services; and(b) excluding the insertion of a device for the purpose of facilitating weight loss(Anaes.) | 253.85 |
| 30483 | Gastrostomy button, caecostomy antegrade enema device (chait etc.) or stomal indwelling device:(a) non‑endoscopic insertion of; or(b) non‑endoscopic replacement of;on a person 10 years of age or over, excluding the insertion of a device for the purpose of facilitating weight loss (Anaes.) | 177.05 |

18 Schedule 1 (items 32520 to 32526)

Repeal the items, substitute:

|  |  |  |
| --- | --- | --- |
| 32520 | Varicose veins, abolition of venous reflux by occlusion of a primary or recurrent great (long) or small (short) saphenous vein of one leg (and major tributaries of saphenous veins as necessary), using a laser probe introduced by an endovenous catheter, if it is documented by duplex ultrasound that the great or small saphenous vein (whichever is to be treated) demonstrates reflux of 0.5 seconds or longer:(a) including all preparation and immediate clinical aftercare (including excision or injection of either tributaries or incompetent perforating veins, or both); and(b) not including radiofrequency diathermy, radiofrequency ablation or cyanoacrylate embolisation; and(c) not provided on the same occasion as a service described in any of items 32500, 32501, 32504 and 32507(Anaes.) | 533.60 |
| 32522 | Varicose veins, abolition of venous reflux by occlusion of a primary or recurrent great (long) and small (short) saphenous vein of one leg (and major tributaries of saphenous veins as necessary), using a laser probe introduced by an endovenous catheter, if it is documented by duplex ultrasound that the great and small saphenous veins demonstrate reflux of 0.5 seconds or longer:(a) including all preparation and immediate clinical aftercare (including excision or injection of either tributaries or incompetent perforating veins, or both); and(b) not including radiofrequency diathermy, radiofrequency ablation or cyanoacrylate embolisation; and(c) not provided on the same occasion as a service described in any of items 32500, 32501, 32504 and 32507(Anaes.) | 793.30 |
| 32523 | Varicose veins, abolition of venous reflux by occlusion of a primary or recurrent great (long) or small (short) saphenous vein of one leg (and major tributaries of saphenous veins as necessary), using a radiofrequency catheter introduced by an endovenous catheter, if it is documented by duplex ultrasound that the great or small saphenous vein (whichever is to be treated) demonstrates reflux of 0.5 seconds or longer:(a) including all preparation and immediate clinical aftercare (including excision or injection of either tributaries or incompetent perforating veins, or both); and(b) not including endovenous laser therapy or cyanoacrylate embolisation; and(c) not provided on the same occasion as a service described in any of items 32500, 32501, 32504 and 32507(Anaes.) | 533.60 |
| 32526 | Varicose veins, abolition of venous reflux by occlusion of a primary or recurrent great (long) and small (short) saphenous vein of one leg (and major tributaries of saphenous veins as necessary), using a radiofrequency catheter introduced by an endovenous catheter, if it is documented by duplex ultrasound that the great and small saphenous veins demonstrate reflux of 0.5 seconds or longer:(a) including all preparation and immediate clinical aftercare (including excision or injection of either tributaries or incompetent perforating veins, or both); and(b) not including endovenous laser therapy or cyanoacrylate embolisation; and(c) not provided on the same occasion as a service described in any of items 32500, 32501, 32504 and 32507(Anaes.) | 793.30 |
| 32528 | Varicose veins, abolition of venous reflux by occlusion of a primary or recurrent great (long) or small (short) saphenous vein of one leg (and major tributaries of saphenous veins as necessary), using cyanoacrylate adhesive, if it is documented by duplex ultrasound that the great or small saphenous vein (whichever is to be treated) demonstrates reflux of 0.5 seconds or longer:(a) including all preparation and immediate clinical aftercare (including excision or injection of either tributaries or incompetent perforating veins, or both); and(b) not including radiofrequency diathermy, radiofrequency ablation or endovenous laser therapy; and(c) not provided on the same occasion as a service described in any of items 32500, 32501, 32504 and 32507(Anaes.) | 533.60 |
| 32529 | Varicose veins, abolition of venous reflux by occlusion of a primary or recurrent great (long) and small (short) saphenous vein of one leg (and major tributaries of saphenous veins as necessary), using cyanoacrylate adhesive, if it is documented by duplex ultrasound that the great and small saphenous veins demonstrate reflux of 0.5 seconds or longer:(a) including all preparation and immediate clinical aftercare (including excision or injection of either tributaries or incompetent perforating veins, or both); and(b) not including radiofrequency diathermy, radiofrequency ablation or endovenous laser therapy; and(c) not provided on the same occasion as a service described in any of items 32500, 32501, 32504 and 32507(Anaes.) | 793.30 |

19 Schedule 1 (item 34103, column headed “Description”)

Omit “or 32526”, substitute “, 32526, 32528 or 32529”.

20 Schedule 1 (after item 38287)

Insert:

|  |  |  |
| --- | --- | --- |
| 38288 | Implantable loop recorder, insertion of, for diagnosis of atrial fibrillation, if:(a) the patient to whom the service is provided has been diagnosed as having had an embolic stroke of undetermined source; and(b) the bases of the diagnosis included the following:(i) the medical history of the patient;(ii) physical examination;(iii) brain and carotid imaging;(iv) cardiac imaging;(v) surface ECG testing including 24‑hour Holter monitoring; and(c) atrial fibrillation is suspected; and(d) the patient:(i) does not have a permanent indication for oral anticoagulants; or(ii) does not have a permanent oral anticoagulants contraindication;including initial programming and testing (Anaes.) | 192.90 |

21 Schedule 1 (after item 42651)

Insert:

|  |  |  |
| --- | --- | --- |
| 42652 | Corneal collagen cross linking, on a person with a corneal ectatic disorder, with evidence of progression—per eye (Anaes.) | 1,200.00 |

Health Insurance Regulations 1975

22 Subregulation 20C(1) (table item 12, column 2)

Omit “61646”, substitute “61647”.

23 Subregulation 20C(1) (table items 13 and 14, column 3)

Omit “61646”, substitute “61647”.