

公告罕見疾病名單暨ICD-10-CM編碼一覽表(依疾病分類排序)

中華民國109年10月13日

| 分類 | 序號 | 中文病名(僅供參考) | 英文病名(縮寫) | ICD-10-CM 診斷代碼 |
|--|----|-----------------------|---|--------------------------------|
| A.先天性代謝異常 | | | | |
| ◎A1尿素循環代謝異常 Urea cycle disorders (高血氮症) | | | | |
| A1 | 01 | 先天性尿素循環代謝障礙 | Congenital Urea cycle disorders | E72.20 |
| | 02 | 瓜胺酸血症 | Citrullinemia | E72.23 |
| | 03 | 乙醯穀胺酸合成酶缺乏症 | Nitroacetylglutamate synthetase deficiency, NAG synthetase deficiency | E72.29 |
| | 04 | 鳥胺酸氨甲醯基轉移酶缺乏症 | Ornithine transcarbamylase deficiency | E72.4 |
| | 05 | 高鳥胺酸血症-高氮血症-高瓜胺酸血症症候群 | Hyperornithinemia-Hyperammonemia-Homocitrullinuria Syndrome | E72.4 |
| ◎A2 胺基酸/有機酸代謝異常 Amino acid metabolic disorders / Organic acidemias | | | | |
| A2 | 01 | 胺基酸代謝疾病 | Amino acid metabolic disorders(Aminoacidopathies) | E72.8 |
| | 02 | 高胱胺酸血症 | Homocystinuria | E72.11 |
| | 03 | 高甲硫胺酸血症 | Hypermethioninemia | E72.19 |
| | 04 | 非酮性高甘胺酸血症 | Nonketotic hyperglycinemia | E72.51 |
| | 05 | 苯酮尿症 | Phenylketouria | E70.0 |
| | 06 | 四氫基喋呤缺乏症 | Tetrahydrobiopterin deficiency | E70.1 |
| | 07 | 遺傳性高酪胺酸血症 | Hereditary tyrosinemia | E70.21 |
| | 08 | 楓糖尿症 | Maple syrup urine disease | E71.0 |
| | 09 | 有機酸血症 | Organic acidemias | E71.118 |
| | 10 | 異戊酸血症 | Isovaleric acidemia | E71.110 |
| | 11 | 戊二酸尿症, 第一型、第二型 | Glutaric aciduria type I、II | type I:E72.3 type11:E71.313 |
| | 12 | 丙酸血症 | Propionic acidemia | E71.121 |
| | 13 | 甲基丙二酸血症 | Methylmalonic acidemia | E71.120 |

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| | 14 | 3-氨基-3-甲基戊二酸血症 | 3-Hydroxy-3-methyl-glutaric acidemia | E71.118 |
| | 15 | 典型苯酮尿症合併蔗糖酶同麥芽糖酶缺乏症 | PAH type PKU combine with Sucrase-isomaltase deficiency | E74.31+E70.0 |
| | 16 | 高離氨基酸血症 | Hyperlysinemia | E72.3 |
| | 17 | 組胺酸血症 | Histidinemia | E70.41 |
| | 18 | 三甲基巴豆醯輔酶 A 羧化酵素缺乏症 | 3-Methylcrotonyl-CoA carboxylase deficiency | E71.19 |
| | 19 | 多發性羧化酶缺乏症 | Multiple carboxylase deficiency | D81.819 |
| | 20 | 高脯胺酸血症 | Hyperprolinemia | E72.59 |
| | 21 | 芳香族 L-胺基酸類脫羧基酶缺乏症 | Aromatic L-amino acid decarboxylase deficiency | E70.9 |
| | 22 | 酪胺酸羥化酶缺乏症 | Tyrosine hydroxylase deficiency | E70.20 |
| | 23 | 甲基丙二酸血症併高胱胺酸血症 (Cbl C 型) | Cobalamin C defect (Methylmalonic aciduria and Homocystinuria, cbl C type) | E71.120+E72.11 |
| ◎ A3 脂質儲積 | | | | |
| A3 | 01 | 高雪氏症 | Gaucher's disease | E75.22 |
| | 02 | GM1/GM2 神經節苷脂儲積症 | GM1/GM2 gangliosidosis | GM1: E75.19 GM2: E75.00 |
| | 03 | Fabry 氏症 | Fabry disease | E75.21 |
| | 04 | Niemann-Pick 氏症，鞘髓磷脂儲積症 | Niemann-Pick disease | E75.240: Type A E75.241: Type B E75.242: Type C E75.243: Type D E75.248: other E75.249: unspecified |
| | 05 | MLD 症候群 | Metachromatic Leukodystrophy (MLD) | E75.25 |

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| | 06 | 球細胞腦白質失養症 | Globoid Cell Leukodystrophy (Krabbe's disease) | E75.23 |
| | 07 | 嬰兒型溶酶體酸性脂肪酶缺乏症 (又稱伍爾曼氏症) | Infantile form Lysosomal Acid Lipase Deficiency (Wolman Disease) | E75.5 |
| ◎A4碳水化合物代謝異常 | | | | |
| A4 | 01 | 半乳糖血症 | Galactosemia | E74.21 |
| | 02 | 肝醣儲積症 | Glycogen storage disease | E74.09:type 0 E74.01:Type I E74.02:type II E74.03:type III E74.09:type IV E74.04:type V E74.09:type VI-XI E74.01:Von Gierke's |
| | 03 | 腦血管屏障葡萄糖輸送缺陷 | Glut (Glucose Transport) 1 deficiency syndrome | E74.8 |
| ◎ A5脂肪酸氧化異常 | | | | |
| | 01 | 脂肪酸氧化作用缺陷 | Fatty acid oxidation defect | E71.30 E71.310 E71.311 E71.312 E71.313 |
| | 02 | 原發性肉鹼缺乏症 | Carnitine deficiency syndrome, primary | E71.41 |
| | 03 | 中鏈脂肪酸去氫酵素缺乏症 | Medium-chain acyl-coenzyme A dehydrogenase deficiency (MCAD) | E71.311 |
| | 04 | 短鏈脂肪酸去氫酶缺乏症 | Short-chain acyl-CoA dehydrogenase deficiency | E71.312 |
| ◎ A6粒線體代謝異常 | | | | |
| A6 | 01 | 粒線體缺陷 | Mitochondrial defect | E88.40 |
| | 02 | Kearns-Sayre 氏症候群 | Kearns-Sayre syndrome | H49.811 H49.812 H49.813 H49.819 |
| | 03 | Leigh 氏童年期腦脊髓病變 | Leigh disease | G31.82 |

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| | 04 | MELAS 症候群 | MELAS | E88.41 |
| | 05 | MNGIE 症候群粒線體性神經胃腸腦病變症候群 | Mitochondrial Neurogastrointestinal Encephalopathy Syndrome | E88.49 |
| | 06 | 丙酮酸鹽脫氫酶缺乏症 | Pyruvate dehydrogenase deficiency | E74.4 |
| | 07 | 巴氏症候群 | Barth Syndrome | E78.71 |
| ◎ A7溶小體代謝異常 | | | | |
| A7 | 01 | 胱氨酸血症 | Cystinosis | E72.04 |
| | 02 | 黏多糖症 | Mucopolysaccharidoses | Type1: E76.01 E76.02 E76.03 Type2:E76.1 other : E76.210 E76.211 E76.219 E76.22 E76.29 Unspecified:E76.3 |
| | 03 | 岩藻糖代謝異常 (儲積症) | Fucosidosis | E77.1 |
| | 04 | 涎酸酵素缺乏症 | Sialidosis | E77.1 |
| | 05 | 黏脂質症 | Mucopolipidosis | type I:E77.1 type II、III E77.0 type IV:E75.11 |
| | 06 | 神經元蠟樣脂褐質儲積症 | Neuronal ceroid lipofuscinosis | E75.4 |
| | 07 | 多發性硫酸脂酶缺乏症 | Multiple Sulfatase deficiency | E75.29 |
| ◎ A8膽固醇及脂質代謝異常 Cholesterol and Lipid metabolism | | | | |
| A8 | 01 | 同合子家族性高膽固醇血症 | Homozygous familial hypercholesterolemia | E78.0 |
| | 02 | 家族性高乳糜微粒血症 | Familial Hyperchylomicronemia | E78.3 |
| | 03 | 豆固醇血症 (植物性) | Sitosterolemia | E78.0 |
| ◎ A9礦物離子缺陷 | | | | |
| A9 | 01 | 威爾森氏症 | Wilson's disease | E83.01 |
| | 02 | Menkes 症候群 | Menkes syndrome | E83.09 |
| | 03 | 鉬輔酶缺乏症 | Molybdenum cofactor deficiency | E61.5 |
| ◎ A10過氧化體代謝異常 | | | | |

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| A10 | 01 | Zellweger 氏症候群 | Zellweger syndrome | E71.510 |
| | 02 | 腎上腺腦白質失養症 | Adrenoleukodystrophy | E71.511 E71.520 E71.521 E71.528 E71.529 |
| | 03 | 肢近端型點狀軟骨發育不良 | Rhizomelic Chondrodysplasia Punctata | E71.540 |

◎ A11 其他代謝異常

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| | 01 | 紫質症 | Porphyria | E80.20 E80.21 E80.29 |
| | 02 | Lesch-Nyhan 氏症候群 | Lesch-Nyhan syndrome | E79.1 |
| | 03 | 亞硫酸鹽氧化酶缺乏 | Sulfite oxidase deficiency | E72.19 |
| | 04 | 碳水化合物缺乏醣蛋白症候群 | Carbohydrate-deficiency glycoprotein syndrome | E77.8 |
| | 05 | 三甲基胺尿症 | Trimethylaminuria | E72.52 |
| | 06 | 先天性全身脂質營養不良症 | Congenital generalized lipodystrophy | E88.1 |
| | 07 | 腦腱性黃瘤症 | Cerebrotendinous Xanthomatosis | E75.5 |
| | 08 | 低磷酸酯酶症 | Hypophosphatasia | E83.39 E83.31 |
| | 09 | Beta 硫解酶缺乏症 | Beta-Ketothiolase Deficiency | E71.19 |
| | 10 | 生物素酶缺乏症 | Biotinidase Deficiency | D81.810 |

B 腦部或神經系統病變

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| B1 | 01 | 多發性硬化症 | Multiple sclerosis | G35 |
| | 02 | 肌萎縮性側索硬化症 | Amyotrophic lateral sclerosis (ALS) | G12.21 |
| | 03 | 共濟失調微血管擴張症候群 | Ataxia telangiectasia | G11.3 |
| | 04 | 亨丁頓氏舞蹈症 | Huntington disease(又稱 Huntington's chorea) | G10 |
| | 05 | 雷特氏症 | Rett syndrome | F84.2 |

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| 06 | 脊髓性肌肉萎縮症 | Spinal muscular atrophy | G12.9 |
| 07 | 脊髓小腦退化性動作協調障礙 | Spinocerebellar ataxia | G11.1 |
| 08 | 結節性硬化症 | Tuberous sclerosis | Q85.1 |
| 09 | 先天性痛不敏感症合併無汗症 | Congenital insensitivity to pain with anhidrosis (CIPA) | L74.4 |
| 10 | 神經纖維瘤症候群第二型 | Neurofibromatosis type II | Q85.02 |
| 11 | Alexander 氏病 | Alexander disease | E75.29 |
| 12 | 僵體症候群 | Stiffperson syndrome | G25.82 |
| 13 | 遺傳性痙攣性下身麻痺 | Hereditary spastic paraplegia | G11.4 |
| 14 | Joubert 氏症候群 (家族性小腦蚓部發育不全) | Joubert syndrome | Q04.3 |
| 15 | Pelizaeus-Merzbacher 氏症 (慢性兒童型腦硬化症) | Pelizaeus-Merzbacher Disease | E75.29 |
| 16 | Charcot Marie Tooth 氏症 (進行性神經性腓骨萎縮症) | Charcot Marie Tooth Disease | G60.0 |
| 17 | 甘迺迪氏症 (脊髓延髓性肌肉萎縮症) | Kennedy Disease | G12.20 G12.21 G12.22 G12.29 |
| 18 | 家族性澱粉樣多發性神經病變 | Familial Amyloidotic Polyneuropathy | E85.1 |
| 19 | Moebius 症候群 | Moebius syndrome | Q87.0 |
| 20 | Mcleod 症候群 | Mcleod syndrome | Q97.8 Q98.8 |
| 21 | Aicardi-Goutieres 症候群 | Aicardi-Goutieres syndrome | G31.89 |
| 22 | 普洛提斯症候群 | Proteus Syndrome | Q87.3 |
| 23 | MECP2 綜合症候群 | Methyl CpG binding protein 2 Duplication Syndrome (MECP2 Duplication Syndrome) | Q99.8 |
| 24 | 腦肋小頷症候群 | Cerebro-Costo-Mandibular Syndrome | Q87.89 |
| 25 | Dravet 症候群 | Dravet Syndrome, DS | G40.311 |

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| | 26 | 腦白質消失症 | Vanishing White Matter Disease | G37.8 |
| | 27 | 泛酸鹽激酶關聯之神經退化性疾 | Pantothenate Kinase Associated Neurodegeneration (PKAN) | G23.0 |
| | 28 | 磷脂質脂解酶 A2 關聯之神經退化性疾 | Phospholipase A2-associated neurodegeneration(PLAN) | G23.0 |
| C 呼吸循環系統病變 | | | | |
| C1 | 01 | 特發性嬰兒動脈硬化症 | Idiopathic Infantile Arterial Calcification | Q28.8 |
| | 02 | 囊狀纖維化症 | Cystic fibrosis | E84.9 |
| | 03 | 特發性或遺傳性肺動脈高壓 | Idiopathic or Heritable pulmonary arterial hypertension (IPAH or HPAH) | I27.0 |
| | 04 | Holt-Oram 氏症候群 | Holt-Oram Syndrome | Q87.2 |
| | 05 | Andersen 氏症候群 (心節律障礙暨週期性麻痺症候群; 鉀離子通道病變) | Andersen syndrome | E74.09 |
| | 06 | 遺傳性出血性血管擴張症 | Hereditary Hemorrhagic Telangiectasia | I78.0 |
| | 07 | 窒息性胸腔失養症 | Asphyxiating thoracic dystrophy | Q77.2 |
| | 08 | 先天性中樞性換氣不足症候群 | Congenital Central Hypoventilation Syndrome | G47.35 |
| D 消化系統病變 | | | | |
| | 01 | 進行性家族性肝內膽汁滯留症 | Progressive intrahepatic cholestasis, PFIC | K83.1 |
| | 02 | 先天性膽酸合成障礙 | Inborn errors of bile acid synthesis | E78.70 |
| | 03 | α 1-抗胰蛋白酶缺乏症 | α 1-Antitrypsin deficiency | E88.01 |
| | 04 | 先天性 Cajal 氏間質細胞增生合併腸道神經元發育異常 | Congenital Interstitial Cell of Cajal Hyperplasia with Neuronal Intestinal Dysplasia | Q43.8 |
| | 05 | 阿拉吉歐症候群 | Alagille Syndrome | Q44.7 |
| E 腎臟泌尿系統病變 | | | | |
| E1 | 01 | Lowe 氏症候群 | Lowe syndrome | E72.03 |
| | 02 | Bartter 氏症候群 | Bartter's syndrome | E26.81 |
| | 03 | 體染色體隱性多囊性腎臟疾病 | Autosomal recessive polycystic kidney disease | Q61.19 |
| F 皮膚病變 | | | | |
| F1 | 01 | 遺傳性表皮分解性水泡症 | Hereditary epidermolysis bullosa | Q81.9 |
| | 02 | 層狀魚鱗癬 (自體隱性遺傳型) | Ichthyosis, lamellar recessive | Q80.2 |

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| | 03 | 膠膜兒 | Collodion baby | Q80.2 |
| | 04 | 斑色魚鱗癬 | Harlequin ichthyosis | Q80.4 |
| | 05 | 水泡型先天性魚鱗癬樣紅皮症（表皮鬆解性角化過度症） | Bullous Congenital ichthyosiform erythroderma (epidermolytic hyperkeratosis) | Q80.3 |
| | 06 | 外胚層增生不良症 | Ectodermal Dysplasias | Q82.4 |
| | 07 | Meleda 島病 | Meleda disease | Q82.8 |
| | 08 | Darier 氏症（毛囊角化病） | Darier's disease | Q82.8 |
| | 09 | 先天性角化不全症 | Dyskeratosis Congenita | Q82.8 |
| | 10 | 皮膚過度角化症雅司病 | Diffuse Non-epidermolytic Palmoplantar Keratoderma type Unna-Thost | Q82.8 |
| | 11 | 色素失調症 | Incontinentia Pigmenti | Q82.3 |
| | 12 | Netherton 症候群 | Netherton Syndrome | Q80.3 |
| G 肌肉病變 | | | | |
| G1 | 01 | 裘馨氏肌肉失養症 | Duchenne muscular dystrophy | G71.0 |
| | 02 | Nemaline 線狀肌肉病變 | Nemaline Rod Myopathy | G71.2 |
| | 03 | Schwartz Jampel 氏症候群 | Schwartz Jampel syndrome | G71.13 |
| | 04 | 肌肉強直症 | Myotonic dystrophy | G71.11 |
| | 05 | 面肩胛肱肌失養症 | Facioscapulohumeral muscular dystrophy | G71.0 |
| | 06 | 肌小管病變 | Myotubular Myopathy | G71.2 |
| | 07 | 貝克型肌肉失養症 | Becker Muscular Dystrophy | G71.0 |
| | 08 | Freeman-Sheldon 氏症候群 | Freeman-Sheldon syndrome | Q87.0 |
| | 09 | 肢帶型肌失養症 | Limb-girdle muscular dystrophy | G71.0 |
| | 10 | 先天性肌失養症 | Congenital Muscular Dystrophy | G71.0 |
| | 11 | 中心軸空肌病 | Central Core Disease | G71.2 |
| | 12 | 多微小軸空肌病 | Multiminicore Disease | G71.2 |
| | 13 | Emery-Dreifuss 肌失養症 | Emery-Dreifuss Muscular Dystrophy (EDMD) | G71.0 |
| | 14 | GNE 遠端肌病變 | GNE myopathy | G71.8 |
| | 15 | 史托摩根症候群 | Stormorken syndrome | D69.8 |
| H 骨及軟骨病變 | | | | |

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| H1 | 01 | 軟骨發育不全症 | Achondroplasia | Q77.4 | |
| | 02 | 成骨不全症 | Osteogenesis imperfecta | Q78.0 | |
| | 03 | 原發性變形性骨炎 | Primary Paget disease | M88.0 M88.1 M88.811 M88.812 M88.819 M88.821 M88.822 M88.829 M88.831 M88.832 M88.839 M88.841 M88.842 M88.849 | M88.851 M88.852 M88.859 M88.861 M88.862 M88.869 M88.871 M88.872 M88.879 M88.88 M88.89 M88.9 |
| | 04 | 鎖骨顱骨發育異常 | Cleidocranial dysplasia | Q74.0 | |
| | 05 | 進行性骨化性肌炎 | Fibrodysplasia Ossificans Progressiva | M61.10 M61.111 M61.112 M61.119 M61.121 M61.122 M61.129 M61.131 M61.132 M61.139 M61.141 M61.142 M61.143 M61.144 M61.145 M61.146 M61.151 | M61.152 M61.159 M61.161 M61.162 M61.169 M61.171 M61.172 M61.173 M61.174 M61.175 M61.176 M61.177 M61.178 M61.179 M61.18 M61.19 |
| | 06 | 裂手裂足症 | Split-hand/ Split-foot malformation (SHFM) | Q71.60 Q71.61 Q71.62 Q71.63 | Q72.70 Q72.71 Q72.72 Q72.73 |
| | 07 | 骨質石化症 | Osteopetrosis | Q78.2 | |
| | 08 | 假性軟骨發育不全 | Pseudoachondroplastic dysplasia | Q77.8 | |
| | 09 | 多發性骨骺發育不全症 | Multiple Epiphyseal Dysplasia | Q78.3 | |
| I 結締組織病變 | | | | | |
| I1 | 01 | 先天結締組織異常第四型 | Ehlers Danlos syndrome IV | Q79.6 | |
| J 血液疾病 | | | | | |

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| | 01 | 重型海洋性貧血 | Thalassemia major | D56.0 D56.1 |
| | 02 | 血小板無力症 | Thrombasthenia | D69.1 |
| | 03 | 同基因合子蛋白質 C 缺乏症 | Homozygous protein C deficiency | D68.59 |
| | 04 | 陣發性夜間血紅素尿症 | Paroxysmal Nocturnal Hemoglobinuria | D59.5 |
| | 05 | 非典型性尿毒溶血症候群 | Atypical Hemolytic Uremic Syndrome | D59.3 |
| K 免疫疾病 | | | | |
| K1 | 01 | 原發性慢性肉芽腫病 | Chronic primary granulomatous disease | D71 |
| | 02 | 先天性高免疫球蛋白 E 症候群 | Congenital Hyper IgE syndrome | D82.4 |
| | 03 | 布魯頓氏低免疫球蛋白血症 | Bruton's agammaglobulinemia | D80.0 |
| | 04 | Wiskott- Aldrich 氏症候群 | Wiskott- Aldrich Syndrome | D82.0 |
| | 05 | 嚴重複合型免疫缺乏症 | Severe combined immunodeficiency | D81.0 D81.1 D81.2 D81.9 |
| | 06 | 補體成份 8 缺乏症 | Complement Component 8 deficiency | D84.1 |
| | 07 | IPEX 症候群 | IPEX Syndrome | E31.0 |
| | 08 | 高免疫球蛋白 M 症候群 | Hyper-IgM syndrome | D80.5 |
| | 09 | γ 干擾素受體 1 缺陷 | Interferon γ receptor 1 deficiency | D84.8 |
| | 10 | 遺傳性血管性水腫 | Hereditary Angioedema (HAE) | D84.1 |
| L 內分泌疾病 | | | | |
| L1 | 01 | Kenny-Caffey 氏症候群 | Kenny-Caffey syndrome | Q87.1 |
| | 02 | 假性副甲狀腺低能症 | Pseudohypoparathyroidism | E20.1 |
| | 03 | 性連遺傳型低磷酸鹽佝僂症 | X-linked hypophosphatemic rickets | E83.31 |
| | 04 | Laron 氏侏儒症候群 | Laron syndrome (Laron Dwarfism) | E34.3 |
| | 05 | Bardet-Biedl 氏症候群 | Bardet-Biedl syndrome | Q87.89 |
| | 06 | Alstrom 氏症候群 | Alstrom Syndrome | Q87.89 |
| | 07 | 持續性幼兒型胰島素過度分泌低血糖症 | Persistent hyperinsulinemic hypoglycemia of infancy (PHHI) | E16.1 |
| | 08 | Wolfram 氏症候群 | Wolfram syndrome, DIDMOAD | E88.9 |

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| | 09 | McCune Albright 氏症候群 | McCune Albright syndrome | Q78.1 |
| | 10 | 短指發育不良及性別顛倒 | Campomelic dysplasia with autosomal sex reversal | Q99.8 |
| | 11 | 腎上腺皮促素抗性 | ACTH resistance | E27.49 |
| | 12 | 第一型遺傳性維生素 D 依賴型佝僂症 | 25-Hydroxyvitamin D 1-Alpha-Hydroxylase Deficiency | E83.32 |
| | 13 | 先天性腎上腺發育不全 | Congenital adrenal hypoplasia | Q89.1 |
| | 14 | Kallmann 氏症候群 | Kallmann syndrome | E23.0 |
| | 15 | 永久性新生兒糖尿病 | Permanent Neonatal Diabetes Mellitus | P70.2 |
| M 先天畸形症候群 | | | | |
| M 1 | 01 | Aarskog-Scott 氏症候群 | Aarskog-Scott syndrome | Q87.1 |
| | 02 | 瓦登伯格氏症候群 | Waardenburg syndrome | E70.8 |
| | 03 | 愛伯特氏症 | Apert syndrome | Q87.0 |
| | 04 | Smith-Lemli-Opitz 氏症候群 | Smith-Lemli-Opitz syndrome | E78.72 |
| | 05 | Larsen 氏症候群 (顎裂-先天性脫位症候群) | Larsen syndrome | Q74.8 |
| | 06 | Beckwith Wiedemann 氏症候群 | Beckwith Wiedemann syndrome | Q87.3 |
| | 07 | Crouzon 氏症候群 | Crouzon syndrome | Q75.1 |
| | 08 | Fraser 氏症候群 | Fraser syndrome | Q87.0 |
| | 09 | 多發性翼狀膜症候群 | Multiple pterygium syndrome | Q79.8 |
| | 10 | Cornelia de Lange 氏症候群 | Cornelia de Lange syndrome | Q87.1 |
| | 11 | 海勒曼-史德萊夫氏症候群 | Hallerman-Streiff Syndrome | Q87.0 |
| | 12 | 歌舞伎症候群 | Kabuki syndrome | Q89.8 |
| | 13 | 耳-齶-指 (趾) 症候群 | Oto-Palato-Digital syndrome | Q87.0 |
| | 14 | Conradi-Hunermann 氏症候群 | Conradi-Hunermann syndrome | Q77.3 |
| | 15 | Treacher Collins 氏症候群 | Treacher Collins Syndrome | Q75.4 |
| | 16 | Robinow 氏症候群 | Robinow Syndrome | Q87.1 |
| | 17 | Pfeiffer 氏症候群 | Pfeiffer syndrome | Q87.0 |

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| 18 | (修正為 B1-27) | | |
| 19 | 指(趾)甲發育症候群 | Nail-Patella Syndrome | Q87.2 |
| 20 | CFC 症候群 | Cardiofaciocutaneous Syndrome | Q87.89 |
| 21 | Peters-Plus 症候群 | Peters-Plus syndrome | Q13.4 |
| 22 | Nager 症候群 | Nager Syndrome | Q75.4 |
| 23 | CHARGE 症候群 | CHARGE Syndrome | Q89.8 |
| 24 | 懷特-薩頓症候群 | White-Sutton syndrome | Q99.8 F84.8 F78 |
| 25 | 克斯提洛氏彈性蛋白缺陷症 | Costello syndrome | Q87.89 |
| 26 | Ayme-Gripp 症候群 | Ayme-Gripp syndrome | Q87.89 |
| 27 | Coffin-Lowry 症候群 | Coffin-Lowry Syndrome | Q89.8 |
| N 染色體異常 | | | |
| N1 | 01 | Angelman 氏症候群 | Angelman syndrome Q93.5 |
| | 02 | DiGeorge's 症候群 | DiGeorge's syndrome D82.1 |
| | 03 | Prader-Willi 氏症候群 | Prader-Willi syndrome Q87.1 |
| | 04 | 威爾姆氏腫瘤、無虹膜、性器異常、智能障礙症候群 (WAGR 症候群) | WAGR syndrome (Wilms' tumor-Aniridia-Genitourinary Anomalies-mental Retardation) Q87.89 |
| | 05 | Miller Dieker 症候群 | Miller Dieker syndrome Q93.88 |
| | 06 | Rubinstein-Taybi 氏症候群 | Rubinstein-Taybi syndrome Q87.2 |
| | 07 | 威廉斯氏症候群 | Williams Syndrome Q93.89 |

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| | 08 | Von Hippel–Lindau 症候群 | Von Hippel–Lindau disease | Q85.8 |
| | 09 | Branchio-Oto-Renal Syndrome (BOR Syndrome) | Branchio-Oto-Renal 症候群 (BOR 症候群) | Q87.89 |
| Z 其他未分類或不明原因 | | | | |
| Z1 | 01 | Cockayne 氏症候群 | Cockayne syndrome | Q87.1 |
| | 02 | 早老症 | Hutchinson Gilford progeria syndrome | E34.8 |
| | 03 | 髮-肝-腸症候群 | Tricho-hepato-enteric syndrome | Q89.7 |
| | 04 | Stargardt's 氏症 | Stargardt's disease | H35.50 |
| | 05 | 隱匿性黃斑部失養症 | Occult Macular Dystrophy ; OMD | H35.50 |

備註：

1. 中華民國93年1月7日署授國字第092401548號公告「胰島母細胞瘤 (Nesidioblastosis)」因屬舊的病名用法，自即日起併入罕見疾病序號 L07號 Persistent hyperinsulinemic hypoglycemia of infancy (PHHI) 名單。
2. 中華民國95年9月12日署授國字第09504009072號公告「Tyrosinemia I、II、III (酪胺基酸血症第一型、第二型、第三型)」，自即日起併入罕見疾病序號 A207號 Hereditary tyrosinemia (遺傳性高酪胺基酸血症) 名單。
3. 中華民國96年8月22日署授國字第09604006002號修正已公告 Urea cycle disorders 為 Congenital Urea cycle disorders。
4. 中華民國98年7月3日署授國字第0980400742號公告 罕見疾病序號 B107號 Spinocerebellar ataxia，原中文病名：脊髓小腦性共濟失調，修正為：脊髓小腦退化性動作協調障礙。
5. 中華民國99年3月19日署授國字第0990400103號公告 罕見疾病序號 K102號 Congenital Hyper IgE syndrome (先天性高免疫球蛋白 E 症候群)，原 ICD-9-CM 編碼：「279.9」，修正為：「288.1」。