

UMC	Naam van Expertisecentrum	Cluster van / Specifieke aandoening(en)
AMC	Amsterdam Lysosome Center ("Sphinx")	Gaucher disease Fabry disease Niemann-Pick disease type A Niemann-Pick disease type B Niemann-Pick disease type C Mucopolysaccharidosis type 1 Mucopolysaccharidosis type 3 Mucopolysaccharidosis type 4 <i>Lysosomal Disease</i> <i>Cholesteryl ester storage disease</i>
AMC	Dutch Centre for Peroxisomal disorders	Peroxisome biogenesis disorder-Zellweger syndrome spectrum Disorder of peroxisomal alpha - beta- and omega-oxidation Rhizomelic chondrodysplasia punctata Non-syndromic pontocerebellar hypoplasia
AMC	Expertise center Vascular medicine	Homozygous familial hypercholesterolemia Familial lipoprotein lipase deficiency Tangier disease
AMC	Centre for Genetic Metabolic Diseases Amsterdam	Disorder of galactose metabolism Disorder of phenylalanine metabolism
AMC	Centre for Neuromuscular Diseases	Neuromuscular disease Motor neuron disease; amyotrophic lateral sclerosis, primary sclerosis and progressive muscular atrophy Idiopathic inflammatory myopathy, incl dermatomyositis, polymyositis, necrotizing autoimmune myopathy and inclusion body myositis Poliomyelitis Hereditary motor and sensory neuropathy Chronic inflammatory demyelinating polyneuropathy, incl. Guillain_Barre syndrome, CIDP, MMN
AMC	Centre for rare thyroid diseases	Congenital hypothyroidism
AMC	Centre for gastroenteropancreatic neuroendocrine tumors	Gastroenteropancreatic endocrine tumor
AMC	Centre for rare hypothalamic and pituitary diseases	Rare hypothalamic or pituitary disease
AMC	Hemophilia Comprehensive Care Treatment Centre	Rare hemorrhagic disorder due to a coagulation factors defect; hemophilia Rare hemorrhagic disorder due to a coagulation factors defect; von Willebrand Disease
AMC	Centre for Sickle Cell Disease	Hemoglobinopathy; incl Sickle cell disease and alpha or beta thalassemia.
AMC	Centre for Bone Marrow Failure	Rare constitutional medullar aplasia Shwachman Diamond disease Congenital neutropenia
AMC	Centre for pediatric thromboembolic events	Rare thrombotic disease of hematologic origin
AMC	Gastro-Intestinal Oncology Centre Amsterdam	Rare cystic pancreas tumors Rare hepatic and biliary tract tumor; incl gallbladder tumors, Ampulla of Vater carcinoma
AMC	Centre for Upper GI tumors Amsterdam	Gastro-esophageal tumor
AMC	Intestinal Failure Unit	Short bowel syndrome, also secondary Chronic intestinal failure

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AMC	Achalasia Center, part of Esophageal Center Amsterdam	Sporadic achalasia	
AMC	Center for immune-mediated and genetic cholestasis syndromes	Primary biliary cirrhosis	
		Primary sclerosing cholangitis	
		Chronic autoimmune hepatitis	*
		Familial intrahepatic cholestasis	*
		Benign recurrent intrahepatic cholestasis	*
		IgG4-related disease	*
		Crigler-Najjar syndrome	*
		Crigler-Najjar syndrome type 1	
		Crigler-Najjar syndrome type 2	
AMC	Centre for Immunodeficiencies	Rare immune disease; incl. primary immunodeficiencies	
AMC	National centre for primary hyperoxaluria	Primary hyperoxaluria	
AMC	Centre for Kawasaki Disease	Kawasaki disease	
AMC	Centre for Pediatric Rheumatic Diseases	Juvenile idiopathic arthritis	
AMC	Expertise Center Clinical Immunology and Rheumatology-Vasculitis	Vasculitis	
AMC	CAHAL (Center for Congenital Heart Disease Amsterdam-Leiden, adult CHD)	Congenital heart malformation; adult congenital heart disease	
AMC	Centre for Hereditary cardiac rhythm and function disorders	Familial isolated arrhythmogenic right ventricular dysplasia	
		Familial dilated cardiomyopathy with conduction defect due to LMNA mutation	
		Genetic cardiac rhythm disease	
		Familial dilated cardiomyopathy	
		Peripartum cardiomyopathy	
		Hypertrophic cardiomyopathy	
AMC	Centre for Marfan syndrome	Marfan syndrome	
AMC	Amsterdam Expert Center for Developmental Disorders	Cornelia de Lange syndrome	
		Rubinstein-Taybi syndrome	
		Pitt-Hopkins syndrome	
		Marshall-Smith syndrome	
		Hutchinson-Gilford progeria	*
AMC	Centre for Hereditary Angioedema	Hereditary angioedema	
AMC	Expertise center for genetic tumors of the digestive tract	Familial adenomatous polyposis	
		Hereditary nonpolyposis colon cancer	
		Hyperplastic polyposis syndrome	
AMC & VUmc	Cystic Fibrosis Centre Amsterdam	Cystic fibrosis	
AMC	Solvent Team	Rare intoxication	
AMC	Center for Idiopathic Nephrotic Syndrome	Idiopathic nephrotic syndrome	

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AMC	Vascular malformations and hemangiomas centre	Diffuse lymphatic malformation Mucocutaneous venous malformations Dandy-Walker malformation - facial hemangioma; all subs Familial multiple nevi flammei	
AMC	AMC Pulmonary Hypertension Center	Eisenmenger syndrome	
AMC	Amsterdam Multidisciplinary Lyme borreliosis Center	Lyme disease	
AMC	Center for Paediatric oncology	Osteosarcoma soft tissue sarcomas	
AMC	Expert Centre for congenital anomalies of the urinary tract EKZ-AMC	Non-syndromic renal or urinary tract malformation; CAKUT	
AMC & VUmc	Amsterdam Centre of congenital malformations	Hirschsprung Disease Esophageal atresia Anorectal malformation Sacrococcygeal teratoma Chronic intestinal pseudoobstruction	
AMC	Center for Osteochondral Defects of the Talus	Osteochondritis van tarsaal/metatarsaal bot	
AMC	Center for condylar hyperplasia	Temporomandibular joint anomaly Condylaire hyperplasie	*
AMC	Amsterdam expert center for bronchopulmonary dysplasia	Bronchopulmonary dysplasia	
AMC & VUmc	Amsterdam Center for ILD and sarcoidosis	Sarcoidosis	*
AMC & VUmc & LUMC	Center for Hereditary Retinal Diseases Leiden Amsterdam	Retinal dystrophy Centrale sereuze chorioretinopathie	
AMC	Center for rare movement disorders	Primary orthostatic tremor Benign adult familial myoclonic epilepsy	
AMC	Center for congenital nevi	Large congenital melanocytic nevus	*
AMC	Center for hematological immune diseases Amsterdam	Autoimmune hemolytic anemia (AIHA)	
AMC	Melioidosis Expertise Center	Melioidosis	*
AMC	LYMMCARE Amsterdam (Lymphoma and Myeloma CARE and REsearch)	Waldenstrom macroglobulinemia B-cell chronic lymphocytic leukemia Lymphoma Multiple myeloma	
AMC	Centre for graft versus host disease Amsterdam	Graft versus host disease (acuut en chronisch)	
Erasmus MC	Center for Lysosomal and Metabolic Diseases	Mucopolysaccharidosis Neuronal ceroid lipofuscinosis Glycoproteinosis Disorder of lysosomal amino acid transport Sphingolipidosis Primary bone dysplasia with defective bone mineralization	
Erasmus MC	Dutch Porphyria Center	Porphyria Erythropoietic protoporphyria Acute hepatic porphyria	
Erasmus MC	Center urea cycle disorders and organic acidurias	Disorder of urea cycle metabolism and ammonia detoxification Disorder of branched-chain amino acid metabolism	

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Erasmus MC	NeMo, expert centre for Neuromuscular and Mitochondrial Diseases	Mitochondrial disease
Erasmus MC	Center for Neuro-inflammatory disorders	Paraneoplastic neurologic syndrome Postinfectious encephalitis Limbic encephalitis CLIPPERS Inflammatory and autoimmune disease with epilepsy Morvan syndrome Isaac syndrome
Erasmus MC	Pompe Center	Glycogen storage disease due to acid maltase deficiency - infantile onset idem - juvenile onset idem- adult onset
Erasmus MC	Neuromuscular Center Erasmus MC	Neuromuscular disease Guillain-Bar syndrome Chronic inflammatory demyelinating polyneuropathy
Erasmus MC	MS center	Neuromyelitis optica Multiple sclerosis variant
Erasmus MC	ENCORE - Expertise Center for Neuro-developmental disorders	Neurofibromatosis type 1 Tuberous sclerosis Angelman syndrome Fragile X syndrome Sturge-Weber syndrome Central nervous system malformation Cardiofaciocutaneous syndrome Costello syndrome
Erasmus MC	Pick Centrum	Behavioral variant of frontotemporal dementia Semantic dementia Progressive non-fluent aphasia Frontotemporal dementia with motor neuron disease Classical progressive supranuclear palsy Corticobasal degeneration Amyotrophic lateral sclerosis-parkinsonism-dementia complex Transmissible spongiform encephalopathy
Erasmus MC	Centre of Oligodontia	Oligodontia
Erasmus MC	Center for pediatric laryngotracheal stenosis	Congenital subglottic stenosis Laryngo-tracheo-esophageal cleft Congenital tracheal stenosis
Erasmus MC	ErasmusMC centre for endocrine disorders	Rare thyroid disease Rare hypothalamic or pituitary disease Rare adrenal disease Endocrine tumor
Erasmus MC	Hemophilia treatment center (volwassen)	Hemophilia Von Willebrand disease Rare hemorrhagic disorder due to a coagulation factors defect, incl FXIII, FXI, FVII,FV deficiency and alpha2-antiplasmin def. Rare hemorrhagic disorder due to a platelet anomaly
Erasmus MC	Expertise Center Rare hemorrhagic disorders (=Hemophilia treatment center, kind)	Hemophilia Von Willebrand disease Rare hemorrhagic disorder due to a coagulation factors defect, incl FXIII, FXI, FVII,FV deficiency and alpha2-antiplasmin def. Rare hemorrhagic disorder due to a platelet anomaly
Erasmus MC	Sickle cell center, volwassen	Sickle cell disease and related diseases Beta-thalassemia and related diseases

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		Alpha-thalassemia and related diseases
Erasmus MC	Sickle cell center, kind	Sickle cell disease and related diseases Beta-thalassemia and related diseases Hemoglobinopathy
Erasmus MC	Leukemia and Stem cell transplantation center	Myeloid hemopathy Acute lymphoblastic leukemia (adult)
Erasmus MC	Multiple myeloma treatment center	Multiple myeloma Aggressive B-cell non-Hodgkin lymphoma
Erasmus MC	Paediatric Brain Tumour Center	Rare nervous system tumor
Erasmus MC	Brain Tumor Center	Glial tumor Tumor of the meninges Primary central nervous system lymphoma Primary germ cell tumor of the central nervous system Embryonal tumor of the neuroepithelial tissue Hemangioblastoma Craniopharyngioma Tumor of cranial and spinal nerves
Erasmus MC	Academic Breast Cancer Center	Hereditary Breast Cancer
Erasmus MC	Nasal, Paranasal, Ear and Skull base tumors workgroup	Tumors arising from the epithelium of the nasal, paranasal and skull base regions Esthesioneuroblastoma
Erasmus MC	Rotterdam Head and Neck Tumor work group	Oral and laryngyal squamous carcinoma
Erasmus MC	Soft tissue sarcoma center	Rare soft tissue tumor
Erasmus MC	Pediatric Surgical Centre for Anatomical Congenital Malformations	Congenital and syndromic diaphragmatic hernia Esophageal atresia Hirschsprung disease Anorectal malformation Omphalocele Gastroschisis Intestinal malformation Chronic intestinal failure
Erasmus MC	Erasmus MC Center for Pancreatic Diseases	Hereditary chronic pancreatitis Recurrent acute pancreatitis Autoimmune pancreatitis type 1 Autoimmune pancreatitis type 2 Familial pancreatic carcinoma Congenital pancreatic cyst
Erasmus MC	Rotterdam Oesophageal and Gastric Cancer Working Group	Esophageal carcinoma; incl. Barrett's oesophagus
Erasmus MC	Erasmus MC Liver Center	Hepatocellular adenoma Adult hepatocellular carcinoma Klatskin tumor <i>Rare hepatic and biliary tract tumor</i>
Erasmus MC	Immunodeficiency center	Primary immunodeficiency
Erasmus MC	Center for systemic allergic diseases	Systemic mastocytosis

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Erasmus MC	Center of rare skin diseases	Netherton syndrome Suppurative hidradenitis Vascular anomaly or angioma
Erasmus MC	Centre of expertise for Children with Autoimmune Diseases	
Erasmus MC	Centre of expertise for Children with Autoimmune Diseases	Juvenile idiopathic arthritis
Erasmus MC	Center for Rare Systemic Immune Disease	Rare systemic disease; Uveitis, Morbus Behcet, Morbus Sjogren, Systemic sclerosis
Erasmus MC	Turner Syndrome Center	Turner syndrome
Erasmus MC	Craniofacial Center	Isolated craniosynostosis Syndromic craniosynostosis; craniofrontonasal syndrome Pierre Robin syndrome associated with branchial arches anomalies <i>Cleft lip with or without cleft palate</i> <i>Cleft palate</i>
Erasmus MC	ErasmusMC Center for Congenital Hand and Upper Limb Malformations	Syndrome with limb malformations as a major feature Non-syndromic limb malformation
Erasmus MC	Expertise center DSD	Disorder of sex development
Erasmus MC	Expert Center spinal disraphism Rotterdam	Spina bifida aperta
Erasmus MC	Center for inherited cardiovascular diseases	Hypertrophic cardiomyopathy Unclassified cardiomyopathy Rare cardiac disease; rare familial occurrence of thoracic aortic abnormalities incl. dissection with (un)known genetic cause, e.g. Aneurysm-osteoarthritis syndrome
Erasmus MC	Center for congenital heart diseases Erasmus MC Rotterdam	Congenital heart malformation
Erasmus MC	Expert Center Prader Willi syndrome	Chromosomal anomaly; Prader Willi syndrome
Erasmus MC	Expert Center Rare Growth Disorders	Silver-Russell syndrome due to maternal uniparental disomy of chromosome 7; also H19 hypomethylation chromosome 11 or unknown Non-acquired pituitary hormone deficiency; not yet known whether or which genes Non-acquired pituitary hormone deficiency; due to known GH gene and yet unknown genetic variants Growth hormone insensitivity syndrome
Erasmus MC	Expertise center Erasmus MC Vascular Genetics	Homozygous familial hypercholesterolemia Familial lipoprotein lipase deficiency
Erasmus MC	Erasmus MC Cystic Fibrosis Center	Cystic fibrosis
Erasmus MC	Pulmonary hypertension center	Rare respiratory disease
Erasmus MC	Interstitial Lung Disease Centre	Interstitial lung disease; adult
Erasmus MC	Sarcoidosis Centre ErasmusMC	Sarcoidosis
Erasmus MC	Mesothelioma centre	Mesothelioma <i>Malignant peritoneal mesothelioma</i> <i>Thymoma</i> <i>Small cell lung cancer</i>

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Erasmus MC	Center for Bronchopulmonary Dysplasia	Bronchopulmonary dysplasia	
Erasmus MC	Erasmus MC Bone Center	Primary bone dysplasia	
Erasmus MC	Center for Perinatal Psychiatry	Postpartum psychosis	
Erasmus MC	Erasmus MC Leprosy Centre	Leprosy	
Erasmus MC	Academic Center Kidney & Hypertension	rare renal tubular disease familial cystic renal disease	
Erasmus MC	Center for Familial and Hereditary Tumors	Lynch Syndrome	
Erasmus MC	Center for pregnancy induced diseases	Pre-eclampsia (< 34 wks) HELLP-syndrome	
Erasmus MC	Centre of Expertise for Uveitis	Uveitis Fuchs hetrochromic iridocyclitis Birdshot chorioretinopathy Paraneoplastic uveitis (ipv uveitis) Cancer-associated retinopathy	* * * * *
Erasmus MC	Center for Genetic Eye Diseases Rotterdam	Retinal dystrophy Retinitis pigmentosa Developmental defect of the eye Color-vision disease Achromatopsia	
Erasmus MC	Center for Congenital bone marrow failure and leukemia predisposition syndromes	Myelodysplastic syndrome Inherited acute myeloid leukemia Constitutional neutropenia	
	Rotterdam Ocular Melanoma Center (ROMC)	Uveal melanoma	
LUMC	Center for Bone Quality	Sclerosteosis Primary bone dysplasia with decreased bone density Primary bone dysplasia with defective bone mineralization Fibrous dysplasia of bone Sternocostoclavicular Hyperostosis Rare parathyroid disease and phosphocalcic metabolism anomaly Primary bone dysplasia with increased bone density Primary bone dysplasia with disorganized development of skeletal components Chronic recurrent multifocal osteomyelitis	
LUMC	Nerve Centre	Nerve lesion	
LUMC	Neuromuscular Center LUMC	Neuromuscular disease Duchenne en Becker muscular dystrophy Acquired neuromuscular junction disease Facioscapulohumeral dystrophy Oculopharyngeal muscular dystrophy Inclusion body myositis	
LUMC	Huntington Disease Center Leiden	Huntington disease	
LUMC	Cerebral Hereditary Angiopathy Center	CADASIL, and RVCL - HCHWA-D	
LUMC	Headache Center LUMC	Rare headache; Sporadic/Familial Hemiplegic Migraine, Visual Snow, SUNCT, Cluster Headache, Trigeminus Neuralgia, Hemicrania Continua, Paroxysmal Hemicrania, TAC nao, Hypnic Headache.	
LUMC	Autonomic Disease Center	Pure autonomic failure, PAF	

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LUMC	Gender Clinic Leiden, WAKZ-Curium LUMC	Gender dysphoria
LUMC	Expertise center for monogenic diabetes mellitus	MODY syndrome
LUMC	Expertise center for lipodystrophy	Primary lipodystrophy
LUMC	Center for Endocrine Tumors Leiden (CETL)	Rare hypothalamic and pituitary disease (behalve MEN 1 en 2) Hereditary pheochromocytoma-paranglioma Thyroid tumor Adrenocortical carcinoma Parathyroid carcinoma <i>Acquired chronic primary adrenal insufficiency</i>
LUMC	Expertise Center Genetics of growth	Growth disorders with a height <-3 SDS Growth disorders with a height >3 SDS Leri-Weill dyschondrosteosis/syndrome Disorders in the GH-IGF1 axis and signaling pathways IGSF1 deficiency syndrome
LUMC	Hemophilia treatment centre LUMC-Haga	Hemophilia A Hemophilia B Von Willebrand disease Acquired hemophilia Acquired von Willebrand syndrome
LUMC	Expert center for aplastic anemia	Idiopathic aplastic anemia
LUMC	Expert center for hemoglobinopathies	Hemoglobinopathy, incl alpha- beta-thalassemia, Sickle Cell Disease, HbS and Hb variants
LUMC	Expert center for pediatric stem cell transplantation	Combined T and B cell immunodeficiency, mainly SCID and ICF syndr Primary immunodeficiency due to a defect in adaptive immunity Acute graft versus host disease
LUMC	Bone and soft tissue tumour clinic	Soft tissue sarcoma; Gastrointestinal stromal (cell) tumour
LUMC	Bone and soft tissue tumour clinic	Rare bone tumor Multiple osteochondromas Adamantinoma Chondromyxoid fibroma Osteosarcoma Ewing sarcoma Chondrosarcoma Giant cell tumor of bone Rare soft tissue tumor
LUMC	Leiden Ocular Oncology Center	Uveal melanoma
LUMC	Female Cancer Center - Leiden (FCC-L)	Rare cancer of the cervix uteri Rare vulvovaginal tumor; Vulvar cancer Rare cancer of the corpus uteri Rare ovarian cancer
LUMC	Expertise Center Pediatric ophthalmology	Retinopathy of prematurity
LUMC	Expertise Center Fetal medicine	Hemolytic disease due to fetomaternal alloimmunization Twin to twin transfusion syndrome Fetal and neonatal alloimmune thrombocytopenia Fetal parvovirus syndrome Hydrops fetalis Congenital heart malformation; fetal cardiac interventions Posterior urethral valve; Lower Urinary Tract Obstruction Non-syndromic respiratory or mediastinal malformation

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LUMC	Prenatal and congenital infections by cytomegalovirus and parvovirus B19	Infectious embryofetopathy; Congenital CMV inf. and fetal and congenital parvovirus B19 inf.	
LUMC	Expert center for cutaneous lymphomas	Primary cutaneous lymphoma	
LUMC	Expertise Center Rare autoinflammatory diseases	Systemic sclerosis Juvenile rheumatoid factor-negative polyarthritis with anti-nuclear antibodies	
LUMC	Congenital Heart malformations (CAHAL pediatric)	Congenital heart malformation; pediatric Rare cardiac rhythm disease; non-genetic	
LUMC	Marfan-FTAAD Clinic	Marfan syndrome; including neonatal Marfan syndrome, FTAAD	
LUMC	Hereditary bowel cancer centre	Hereditary nonpolyposis colon cancer; Lynch caused by MLH1 or MSH2 mutation Hereditary nonpolyposis colon cancer; Lynch caused by MSH6 or PMS2 mutation MUTYH-related attenuated familial adenomatous polyposis Familial adenomatous polyposis; APC associated polyposis	
LUMC	Clinic for Lupus-, Vasculitis- and Complement-mediated systemic diseases	C3 glomerulonephritis <i>Pauci-immune glomerulonephritis</i> <i>Anti-neutrophil cytoplasmic antibody-associated vasculitis</i> <i>Immunoglobulin-mediated membranoproliferative glomerulonephritis</i>	
LUMC	Center for Inherited kidney disease	Autosomal recessive polycystic kidney disease Autosomal Dominant Polycystic Kidney Disease, PKD1 mutation Autosomal Dominant Polycystic Kidney Disease PKD2 mutation Autosomal Dominant Medullary Cystic Kidney Disease	
LUMC	Alpha1 International Registry (AIR)	Alpha-1-antitrypsin deficiency	
LUMC	Center for Narcolepsia	Narcolepsy-cataplexy	
LUMC	Expert center for familial cutaneous melanoma	Familial melanoma; incl. FAMMM syndrome and FAMMMPC syndrome	
LUMC	Center of expertise Coffin-Siris syndrome	Coffin-Siris syndroom	
LUMC	Familial and hereditary breast cancer center	Hereditary breast and ovarian cancer syndrome Hereditary breast cancer	
LUMC	Center for vestibular schwannoma and NF2	Vestibularis schwannoom	*
LUMC & AMC & Vumc	Center for Hereditary Retinal Diseases Leiden Amsterdam	Retinal dystrophy Centrale sereuze chorioretinopathie	
LUMC & Erasmus MC	Center for Congenital bone marrow failure and leukemia	Myelodysplastic syndrome Inherited acute myeloid leukemia Constitutional neutropenia	
LUMC	Center for Polyomavirus-associated skin infections and cancer (Acronym PASIC)	Virus-associated trichodysplasia spinulosa, usually known as trichodysplasia spinulosa	*
LUMC	Expert Center for hematopoietic stem cell transplantation and T-cell immunotherapy	Acute myeloid leukemia Acute lymphoblastic leukemia Plasma cell tumor	
LUMC	Center for primary and secondary immunodeficiencies	Mendelian susceptibility to mycobacterial diseases Combined T and B cell immunodeficiency	
MUMC+	Expertise Center Galactosemia	Galactosemia	

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MUMC+	Expertise Center Hyperostosis of the skull	Osteopetrosis
MUMC+	Expertise Center Huntington's disease	Huntington disease
MUMC+	Neuromuscular Centre MUMC+	Neuromuscular disease Myotonic dystrophy Sodium channelopathy-related small fiber neuropathy Duchenne and Becker muscular dystrophy
MUMC+	Expertise Center Neural tube defects	Neural tube defect
MUMC+	Academic Center for Epilepsy	Epilepsy syndrome Continuous spikes and waves during sleep, epileptic encephalopathy Early infantile epileptic encephalopathy
MUMC+	Expertise Center Cerebral Palsy	Spastic diplegia - infantile type. 1. spastic unilateral cerebral palsy 2. dyskinetic cerebral palsy
MUMC+	Expertise Center Hereditary Tumors	Hereditary breast and ovarian cancer
MUMC+	Center for Genodermatoses	Inherited ichthyosis Erythrokeratoderma Lymphedema Birt-Hogg-Du syndrome Rare genetic skin disease <i>Hereditary palmoplantar keratoderma</i> <i>Basal cell nevus syndrome</i>
MUMC+	Expertise center Rare syndromes and cognitive disorders	Rare developmental defect during embryogenesis Kabuki syndrome Rett Syndrome
MUMC+	Expertise center Cardiogenetics MUMC+	Rare familial disorder with hypertrophic cardiomyopathy Dilated cardiomyopathy Arrhythmogenic right ventricular dysplasia Familial long QT syndrome Idiopathic ventricular fibrillation - not Brugada type Brugada syndrome
MUMC+	Expertise center Pulmonary hypertension	Pulmonary hypertension with unclear multifactorial mechanism; auto immune mechanisms in PH, and right ventricular failure
MUMC+ & Radboudumc	Marfan and related disorders policlinic	Marfan syndrome Loeys-Dietz syndrome Familial thoracic aortic aneurysm and aortic dissection
MUMC+	Limburg renal registry	EGPA, GPA and MPA Cryoglobulinemic vasculitis
MUMC+	Gastro-intestinal center Maastricht	Biliary tract carcinoma en Hepathocellular carcinoma
MUMC+	Maastricht Soft Tissue Tumor Center	Rare soft tissue tumor; long list, together all soft tissue sarcomas in adults
MUMC+	Maastricht Gynaecological Oncology Center	Rare ovarian cancer; epithelial and non- epithelial, also tumor of Fallopian tubes Rare cancer of the corpus uteri
MUMC+	Center for Endocrine tumors	Thyroid carcinoma
MUMC+	Neuroendocrine tumours Center	Bronchial NET en Merkelcell carcinoma
MUMC+	Neuro-oncologie centrum Maastricht	Glial tumor Tumor of cranial and spinal nerves

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MUMC+	Maastricht Head & Neck Cancer Center	Rare otorhinolaryngologic tumor; as a group	
MUMC+	Lung cancer center maastricht	Small cell lung cancer	
MUMC+	<i>Division of Balance Disorders</i>	<i>Idiopathic Bilateral Vestibulopathy</i>	
MUMC+	<i>MUMC Mediastinal tumors</i>	<i>Thymic tumor</i>	*
Radboudumc	Nijmegen centre for mitochondrial disorders	Mitochondrial disease	
Radboudumc	Nijmegen Center for Disorders of Glycosylation (NCDG)	Congenital disorder of glycosylation	
Radboudumc	Centre for genetic movement disorders	Rare hereditary ataxia, mainly autosomal dominant and recessive cerebellar ataxias Hereditary spastic paraplegia Mainly Sjogren-Larsson syndrome, GLUT1 deficiency syndrome, and disorders of dopamine metabolism	
Radboudumc	Neuromuscular Centre	Neuromuscular disease Facioscapulohumeral dystrophy Oculopharyngeal muscular dystrophy Duchenne and Becker muscular dystrophy Myotonic dystrophy Neuralgic amyotrophy Idiopathic inflammatory myopathy Congenital myotonia and paramyotonia congenita Non-dystrophic myopathy	
Radboudumc	Center for rare CNS and retinal vascular disease	Acquired aneurysmal subarachnoid hemorrhage	
Radboudumc	Hearing & Genes Centre	Rare genetic deafness Usher syndrome	
Radboudumc	Radboud Adrenal Centre	Cushing syndrome Adrenogenital syndrome Rare primary hyperaldosteronism Primary adrenal insufficiency Adrenal/paraganglial tumor, incl Von Hippel Lindau and MEN-2 syndrome Adrenal/paraganglial tumor; except catecholamines, aldosterone or cortisol producing . Incl. incidentalomas and carcinomas	
Radboudumc	Centre of Paroxysmal Nocturnal hemoglobinuria	Paroxysmal nocturnal hemoglobinuria	
Radboudumc	Haemophilia treatment centre	Rare coagulation disorder; hemophilia Rare coagulation disorder; von Willebrand Disease Rare coagulation disorder; other	
Radboudumc	Radboud Center for iron disorders	Disorder of iron metabolism and transport, focus on hereditary hemochromatosis, FTH1-related iron overload, congenital atransferrinemia, microcytic anemia with iron overload and aceruloplasminemia Sideroblastic anemia Constitutional anemia due to iron metabolism disorder Constitutional dyserythropoietic anemia, mainly type I-IV	
Radboudumc	Hereditary cancer centre	Hereditary breast and ovarian cancer syndrome; BRCA mutation carriers Hereditary nonpolyposis colon cancer; Lynch Syndrome Familial gastric cancer, incl her. diffuse GC APC-related attenuated familial adenomatous polyposis	
Radboudumc	Centre for thyroid carcinomas	Thyroid tumor	
Radboudumc	Radboud Skull base centre	Rare tumor; different very rare skull base tumors Rare nervous system tumor; acoustic neuroma; cerebello pontine angle tumor; excl. NF2 patients	

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		Tumor of endocrine glands; Hereditary pheochromocytoma-paraganglioma	
Radboudumc	Radboudumc pediatric center for congenital malformations of the intestinal tract	High anorectal malformation	
		Intermediate anorectal malformation	
		Low anorectal malformation	
		Anal fistula	
		VACTERL/VATER association	
		Caudal regression sequence	
		Hirschsprung disease	
		Cloacal exstrophy	
		Intestinal malformation	*
		Esophageal malformation	*
		Gastroschisis	*
		Omphalocele	*
Radboudumc	Ophthalmogenetic center	Genetic vitreous-retinal disease	
		Choroideremia	
		Stargardt disease and other ABCA4-related diseases	
		Retinal dystrophy; central serous retinopathy	
		Leber congenital amaurosis	
Radboudumc	Liver cyst center	Isolated polycystic liver disease	
Radboudumc	Radboudumc Expertise Center for immunodeficiency and autoinflammation	Primary immunodeficiency	
		Autoinflammatory syndrome with immune deficiency	
		Immunodeficiency due to a complement cascade protein anomaly	
		Schnitzler syndrome	
		Mevalonate kinase deficiency = Hyper IgD Syndrome (HIDS)	
		Autoinflammatory syndrome	*
Radboudumc	Mycology reference center	Aspergillosis; chronic aspergillosis and ABPA	
		Chronic mucocutaneous candidiasis	
		Rare mycosis; in patients with hyper IgE syndrome	
		Rare mycosis; in patients with chronic granulomatous dis.	
		Aspergillosis	
Radboudumc	Radboud Center Renal Disorders	Glomerular disease	
		Rare renal tubular disease	
		Familial cystic renal disease; all cystic kidney dis. in children, incl. ciliopathies/nephronophthoses	
		Renal or urinary tract malformation	
		Cystinosis	
		Thrombotic microangiopathy	
Radboudumc	Centre for genetic neurodevelopmental disorders	Kleefstra syndrome due to a point mutation	
		Koolen-de Vries syndrome	
		Cowden syndrome	
		KBG syndrome	
		Noonan syndrome	
		Prader Willi Syndrome	
Radboudumc	Craniofacial team Nijmegen	Cranial malformation	
		Craniosynostosis	
Radboudumc	Radboud DSD centre	46 -XX disorder of sex development induced by fetal androgens excess	
		46 -XY disorder of sex development	
		Turner syndrome	
Radboudumc & MUMC+	Marfan and related disorders policlinic	Marfan syndrome	
		Loeys-Dietz syndrome	
		Familial thoracic aortic aneurysm and aortic dissection	

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Radboudumc	Centre for Hemangiomas and Congenital Vascular Anomalies Nijmegen (HECOVAN)	Vascular tumor; incl complicated hemangiomas	
		Venous malformation	
		Lymphatic system malformation	
		Arteriovenous malformation	
		Complex - combined vascular malformation; incl Klippel-Trenaunay-syndrome	
Radboudumc	(Pediatric) urology center	Bladder exstrophy; incl. cloacal exstrophy and epispadias	
		Agensis and aplasia of uterine body; Cloacal anomalies and anal atresia in combination with urological tract anomalies	
		Non-syndromic urogenital tract malformation of male and female; e.g. in spina bifida	
		Posterior urethral valve	
		Posterior hypospadias	
Radboudumc	Radboud Intestinal failure Unit	Chronic intestinal failure	
Radboudumc	Cleft (lip and) palate center Nijmegen	Cleft palate; cleft hard and / or soft palate	
		Cleft palate; cleft lip/palate	
		Cleft palate; Submucosal cleft palate	
		Oculo-auriculo-vertebral spectrum (=hemifaciale microsomia)	
Radboudumc	Radboudumc Center for Pulmonary Hypertension	Idiopathic pulmonary arterial hypertension; all kinds of PH	
Radboudumc/UCCZ Dekkerswald	Center for Mycobacterial diseases	Tuberculosis; also incl. nontuberculous mycobacterial inf.	
Radboudumc	Radboud Center for Congenital Diaphragmatic Hernia and neonatal pulmonary hypertension	Congenital diaphragmatic hernia; non- syndromic and syndromic	
		Pulmonary hypertension owing to lung disease and/or hypoxia; Persistent Pulmonary Hypertension of the newborn	
		<i>Congenital alveolo-capillary dysplasia</i>	
Radboudumc	Radboud Pituitary Center	Cushing disease	
		Somatotropic adenoma	
		Prolactinoma; also incl. other rare types of functioning pituitary tumor	
		Pituitary deficiency; also incl. all hypothalamic and pituitary diseases resulting in pituitary def.	
Radboudumc	Radboud Sarcoma Center	Bone sarcoma; incl. bone and soft tissue tumors and GIST (gastrointestinal stromal tumors)	
Radboudumc	Radboud Centre for vulvar and cervix cancer and Dutch Mole Registry	Gestational trophoblastic neoplasm; all different subgroups	
		Vulvar intraepithelial neoplasia; rare vulvar cancers and VIN	
Radboudumc	Radboudumc Neuro-oncological center	Tumor of the neuroepithelial tissue	
		Primary melanocytic tumor of the CNS	
Radboudumc	Center for Head and Neck Oncology	Squamous cell carcinoma of head and neck	
		Malignant epithelial tumor of the salivary glands	
Radboudumc	Radboudumc center for congenital disorders of dental development	Oligodontia	
		<i>Schöpf-Schulz-Passarge syndrome</i>	*
Radboudumc	Radboudumc center for facial palsy	Paralytic facial malformation	*
Radboudumc	Radboud University Medical Centre CF centre	Cystic fibrosis	
Radboudumc	Radboudumc Center for male infertility	Male infertility due to obstructive azoospermia	*
		Rare idiopathic male infertility	*
		Male infertility due to gonadal dysgenesis or sperm disorder	*
Radboudumc	Radboud Center for Infectious diseases	Lyme disease	*
		Q fever	*

UMC	Naam van Expertisecentrum	Cluster van / Specifieke aandoening(en)	
		<i>Arbovirus fever</i>	*
		<i>Malaria</i>	*
		<i>Idiopathic recurrent and disabling cutaneous herpes</i>	*
		<i>Whooping cough (kinkhoest)</i>	*
		<i>Viral hemorrhagic fevers</i>	*
		<i>Fulminant virale hepatitis</i>	*
Radboudumc	Radboudumc center for systemic autoimmune diseases	Systemic sclerosis	
		Localized scleroderma	
		Eosinophilic fasciitis	
UMCG	Expert centre for Phenylketonuria (PKU) and Tyrosinemia type I	Disorder of phenylalanin or tyrosine metabolism	
UMCG	Expert centre for hepatic Glycogen Storage Diseases	Glycogen storage disease due to glucose-6-phosphatase deficiency type a	
		Glycogen storage disease due to glucose-6-phosphatase deficiency type b	
		Glycogen storage disease due to glycogen debranching enzyme deficiency	
		Glycogen storage disease	
		Glycogen storage disease due to liver glycogen phosphorylase deficiency	
UMCG	Expert centre for M(C)ADD	Medium chain acyl-CoA dehydrogenase deficiency	
		Multiple acyl-CoA dehydrogenation deficiency - severe neonatal type	
		Multiple acyl-CoA dehydrogenation deficiency - mild type	
UMCG	Expert centre for serine deficiencies	Neurometabolic disorder due to serine deficiency	
UMCG	Neurovascular Team UMCG	Cerebral malformation, intracranial dural AV-fistula	
		Spinal arteriovenous shunts	
		Cerebral malformation, brainstem cavernomas	
		Cerebral malformation, proliferative angiopathy	
UMCG	Expert centre for movement disorders in adults and children	Rare dystonia: myoclonus, focal, generalised, dopa responsive	
		Hyperekplexia	
		Neurodegeneration with brain iron accumulation; a.o. PKAN	
		Rare myoclonus; myoclonus dystonia GOSR2 and FCMTE	
		Psychogenic movement disorders	
		Autosomal dominant cerebellar ataxia, incl recessive ataxias	
		Paroxysmal dyskinesia	
		Inherited congenital spastic tetraplegia	
		Rare choreic movement disorder, huntington's disease	
		Neurometabolic disease, related tot movement disorders	
UMCG	Expertise Center Groningen Papilloma studies	Recurrent respiratory papillomatosis	
UMCG	Adrenal centre UMCG	Catecholamine-producing tumor, incl. pheochromocytoma/ paraganglioma and non-secreting head and neck paragangliomas	
		Rare primary hyperaldosteronism	
UMCG	Hemophilia Treatment Centre UMCG	Rare hemorrhagic disorder; hemophilia	
		Rare hemorrhagic disorder; von Willebrand Disease	
		Rare hemorrhagic disorder: other allied bleeding disorders	
UMCG	Expert centre mastocytosis Netherlands (ECMN)	Mastocytosis	
UMCG	Expert Center head- and neck oncology	Squamous cell carcinoma of head and neck	
UMCG	Expert Center neuro-oncology in adults	Glial tumor	
UMCG	Expert centre for carcinoid / neuroendocrine carcinoma's (NEC)	Carcinoid tumor and carcinoid syndrome	
UMCG	Centre of familial tumors	Inherited cancer-predisposing syndrome; incl VHL, MEN1, MEN2 and familial paraganglioma/PCC.	
UMCG	Thyroid cancer centre	Thyroid tumor	
UMCG	Familial Breast Ovarian Cancer Clinic	Hereditary breast and ovarian cancer syndrome; BRCA1 BRCA2	

UMC	Naam van Expertisecentrum	Cluster van / Specifieke aandoening(en)
UMCG	Familial Colorectal Cancer Clinic	Hereditary nonpolyposis colon cancer Familial adenomatous polyposis; incl FAP and MUTYH
UMCG	Expertise Center Germcell tumors	Testicular germ cell tumor
UMCG	Expertise Center Soft tissue and bone tumors	Osteosarcoma Soft tissue sarcomas
UMCG	Pediatric Neuro-oncology Team	Medulloblastoma; and PNET Rare nervous system tumor
UMCG	Expertise Center Gyneco-oncology UMCG	Rare vulvovaginal tumor; squamous cell carcinoma of vulva Rare cancer of the cervix uteri; incl. squamous cell carcinoma Malignant epithelial tumor of ovary; different types adenocarcinoma
UMCG	UMCG/ oesophageal/ gastric cancer tumorgroup	Esophageal adenocarcinoma
UMCG	Small bowel rehabilitation and transplant centre	Chronic intestinal failure
UMCG	Expertise Center pediatric liver disease, pediatric liver surgery and pediatric liver transplantation	Biliary atresia
UMCG	Paediatric centre for Rheumatologic and immunologic diseases	Juvenile idiopathic arthritis
UMCG	Tuberculosis centre Beatrixoord	Tuberculosis
UMCG	Expertise Center for Polycystic Kidney Diseases	Familial cystic renal disease
UMCG	Center for Blistering Diseases	Inherited epidermolysis bullosa Autoimmune bullous skin disease; all forms of pemphigus and pemfigoid
UMCG	Groningen Unit for Amyloidosis Research & Development (GUARD)	Primary systemic amyloidosis Primary localized amyloidosis Secondary amyloidosis Familial amyloid polyneuropathy Transthyretin-related familial amyloid cardiomyopathy Senile systemic amyloidosis
UMCG	Expert centre for systemic vasculitis	Vasculitis
UMCG	Expert centre Sjögren syndrome	Systemic autoimmune disease; Sjögren (including MALT lymphoma) Tumor of hematopoietic and lymphoid tissues; MALT lymphoma associated with Sjögren's disease
UMCG	Clinic for Connective tissue disorders	Marfan syndrome
UMCG	The multidisciplinary CHARGE clinic	CHARGE syndrome
UMCG	Clinic for rare chromosome disorders	Autosomal anomaly; wide diversity of chromosomal deletions and duplications (and not the more common trisomies)
UMCG	Expert Centre for Cardiogenetics	Familial isolated arrhythmogenic ventricular dysplasia - biventricular form Cardiomyopathy Genetic cardiac rhythm disease
UMCG	Expert Center for Children and Adults with rare Congenital Heart Diseases	Congenital heart malformation
UMCG	Dutch expertise centre for lympho-vascular medicine	Primary lymphedema Syndromic lymphedema
UMCG	Expert Center for Children and Adults with Pulmonary Hypertension	Pulmonary arterial hypertension

UMC	Naam van Expertisecentrum	Cluster van / Specifieke aandoening(en)	
UMCG	Cystic Fibrosis centre Groningen	Cystic fibrosis	
UMCG	UMCG Pituitary Center	Rare pituitary disease	
UMCG	<i>Pediatric Oncology group UMCG</i>	<i>Tumor of hematopoietic and lymphoid tissues</i>	
UMCG	<i>Center for rare inherited inborn errors of metabolism</i>	<i>Molybdenum cofactor deficientie type A</i>	
UMCG	<i>Paediatric Colorectal Expertise Center Groningen</i>	<i>Hirschsprung disease</i> <i>Anorectal malformation</i>	* *
UMCG	<i>Centre of expertise for choledochal malformations</i>	<i>Choledochal cyst</i>	
UMCG	<i>Necrotizing Enterocolitis</i>	<i>Necrotiserende Enterocolitis</i>	
UMCU	Center Inherited Metabolic Diseases	Disorder of fatty acid oxidation and ketone body metabolism Disorder of pyridoxine metabolism	
UMCU	Brain Centre Rudolf Magnus, Neuromuscular Diseases	Neuromuscular disease Spinal muscular atrophy	
UMCU	Center for Refractory Pediatric Epilepsy	Rare epilepsy Tuberous sclerosis Continuous spikes and waves during sleep and ESES Epilepsy syndrome	
UMCU	Center of Excellence in Congenital Orofacial and Dental Anomalies	Rare odontologic disease; oligodontia 22q11.2 deletion syndrome Isolated Pierre Robin syndrome Hemifaciale microsomie & Microtie Cleft/lip palate	
UMCU	Expertise centre for benign hematology, thrombosis and hemostasis, Van Creveld clinic	Rare hemorrhagic disorder due to a coagulation factors defect; hemophilia Rare hemorrhagic disorder due to a coagulation factors defect; von Willebrand Disease Rare hemorrhagic disorder due to a coagulation factors defect; deficiency of factor II/V/ VII/X/XI Rare hemorrhagic disorder due to a coagulation factors defect; antiplasmin deficiency Rare anemia Rare constitutional medullary aplasia; Fanconi, Diamond-Blackfan anemia and congenital neutropenia Rare hemorrhagic disorder; Congenital and acquired platelet disorders	
UMCU	Expertise centre for malignant hematology	Multiple myeloma Non Hodgkin lymphoma Acute lymphoblastic and myeloid leukemia	
UMCU	Centre for rare tumors	Inherited cancer-predisposing syndrome; Men 1 Multiple endocrine neoplasia type 2A; incl. fam medullary thyroid carcinoma, MEN2B and sporadic medullary thyroid carcinoma Von Hippel-Lindau disease Thyroid tumor	
UMCU	Expertise Center Rare GI and hepatic diseases	Progressive familial intrahepatic cholestasis Wilson disease Intractable diarrhea of infancy; due to genetic defects	
UMCU	UMCU ophthalmology uveitisgroup	Anterior uveitis; (non) infectious Posterior uveitis; (non) infectious and in syst dis.	

UMC	Naam van Expertisecentrum	Cluster van / Specifieke aandoening(en)	
		Systemic diseases with panuveitis	
		Rare inflammatory eye disease	
		Intermediate uveitis; (non) infectious	
UMCU	Expertise centre for primary immunodeficiencies	Immunodeficiency predominantly affecting antibody production; incl. CVID, XLA, other types of complete agammaglobulinemia	
		Immunodeficiency predominantly affecting antibody production; incl. SADNI, IgG subclass- and IgA def.	
		Primary immunodeficiency due to a defect in adaptive immunity; B and T cell immunodeficiencies: SCID, CID	
		Primary immunodeficiency due to a defect in innate immunity	
		Primary hemophagocytic lymphohistiocytosis	
		Graft versus host disease	
UMCU	Expertise centre systemic autoimmune diseases	Juvenile idiopathic arthritis	
		Periodic fever syndrome, incl. CAPS, FMF, behcet, Traps, Pfapa	
		Juvenile dermatomyositis	
		Pediatric systemic lupus erythematosus	
		Vasculitis	
		Systemic sclerosis	
		Rare coagulation disorder; Antiphospholipid syndrome	
		Systemic autoimmune disease; extraglandular manifestations in Sjogren	
UMCU	WKZ center for congenital malformations	Esophageal atresia	
		Anorectal malformation	
		Hirschsprung Disease	
UMCU	Multidisciplinary Center for Limb Reduction Defects	Rare bone disease; m.n. primaire skeletdysplasieën en dysostoses, ook osteogenesis imperfecta en achondroplasia	
		Non-syndromic limb reduction defects	
UMCU	Children's Heartcenter WKZ	Congenital heart malformation	
UMCU	Clinic for Tuberous Sclerosis Complex	Rare genetic neurological disorder; Tuberous Sclerosis Complex	
UMCU	Cystic Fibrosis Clinic	Cystic fibrosis	
UMCU	Expert Centre Hereditary and congenital nephrologic and urologic disorders	Rare renal disease; congenital or inherited renal or urinary tract disease	
		Familial cystic renal disease; nephronophthisis, as feature of i.e. Joubert and Meckel or isolated or part of other ciliopathies.	
		Non-syndromic renal or urinary tract malformation; CAKUT	
UMCU	Center for Rare Ear and Hearing Diseases	Middle ear anomaly	
UMCU & NKI-AvL	Expert Center of Neuroendocrine carcinomas	Gastroenteropancreatic endocrine tumor	
UMCU	Center of vascular anomalies Utrecht	Vascular anomaly	
UMCU	Sylvia Toth Center for Multi-disciplinary follow up of Lysosomal Storage Disorders, University Medical Center Utrecht	Hurler disease	
UMCU	Mobility Clinic	Osteochondritis Dissecans	
UMCU	Center for inherited cardiovascular disease	Cardiomyopathy	*
		Genetic cardiac rhythm disease	*
		Rare genetic vascular disease	*
UMCU	Centre of expertise for extracranial carotid artery aneurysms (ECAA)	Extracraniale carotis aneurysma	*

UMC	Naam van Expertisecentrum	Cluster van / Specifieke aandoening(en)	
UMCU	Dutch National Expertise Center for Pseudoxanthoma elasticum (DNECP)	Pseudoxanthoma elasticum-	
UMCU	Utrecht Center for Keratoconus and Corneal Dystrophies	Keratoconus	*
UMCU	Center for Neonatal Neurology	Hypoxic ischemic brain injury Pediatric arterial ischemic stroke Periventricular leukomalacia	
UMCU	Center for Intracranial Vasculopathies	Sneddon syndrome Primary central nervous system vasculitis Moya Moya angiopathy:	* * *
UMCU	Head and Neck Working Group Utrecht	Squamous cell carcinoma of head and neck Squamous cell carcinoma of the oral tongue Rare tumors of salivary glands Nasopharyngeal carcinoma	
UMCU	UMCU Neuro-Oncology Center	Glial tumor	
VUmc	Centre for Genetic Metabolic Diseases Amsterdam (CGMA) - VUmc specific	Disorder of creatine biosynthesis	
VUmc	Center for Childhood White Matter Disorders	Rare neurologic disease	
VUmc	Expertise Center for Osteogenesis Imperfecta	Primary bone dysplasia with decreased bone density	
VUmc	Center for obstetric brachial plexus lesion	Rare neurologic disease; obstetric brachial plexus lesions	
VUmc	Center for pediatric rehabilitation medicine	Spastic diplegia - infantile type	
VUmc	Center for pediatric oral and maxillofacial surgery	Robin sequence	
VUmc	Expertise Center Head and Neck tumors	Nasopharyngeal carcinoma Squamous cell carcinoma of the oral cavity and lip Rare head and neck tumor Hereditary pheochromocytoma-paranglioma	
VUmc	Celiac disease center	Refractory celiac disease Autoimmune hepatitis Short Bowel Syndrome Enteropathy Associated T-cell Lymphoma	
VUmc	Center for rare haematologic cancers	Multiple myeloma Non Hodgkin lymphoma Acute lymphoblastic and myeloid leukemia Myelodysplastic syndromes (al erkend onder cluster NHL) Chronic Myeloid Leukemia (al erkend onder cluster NHL) Primary Myelofibrosis Hodgkin Lymphoma (al erkend onder cluster AL&ML) Myeloid hemopathy Lymphoid hemopathy	
VUmc	Dutch Center for Oral Medicine and Oral Pathology	Squamous cell carcinoma of head and neck; Ameloblastomas and Keratocystic odontogenic tumors, Gorlin-Goltz syndrome	
VUmc	Brain Tumor Center Amsterdam	Rare tumor; Brain Tumors	
VUmc	Dutch Retinoblastoma Center	Retinoblastoma	

UMC	Naam van Expertisecentrum	Cluster van / Specifieke aandoening(en)	
VUmc	Expertise Center Neurophthalmology	Neuromyelitis optica: autoimmune optic neuropathies incl. NMO, CRION, RION, ION, MSON	
		Paraneoplastic neurologic syndrome; optic neuropathies relevant to DD of CRION, RION and NMO-SD	
		Adult-onset myasthenia gravis; ocular MG	
		Idiopathic intracranial hypertension; loss of vision due to IIH	
		Acute zonal occult outer retinopathy	
VUmc	Expertise Center Preeclampsia from origin to healthy aging	Preeclampsia	
VUmc	Centre for Systemic Sclerosis and Systemic Lupus Erythematosus, embedded in Amsterdam Rheumatology and immunology Centre	Systemic sclerosis	
VUmc	Pediatric nephrology centre	Idiopathic nephrotic syndrome	
		Non-syndromic renal or urinary tract malformation; CAKUT	
VUmc	Cleft Lip and Palate Team	Rare developmental defect during embryogenesis; cleft lip and palate	
VUmc	Birt-Hogg-Dubé task force	Inherited renal cell cancer-predisposing syndrome; Birt-Hogg-Dubé syndrome	
VUmc	Expertcenter for Fibrodysplasia Ossificans Progressiva	Fibrodysplasia ossificans progressiva	
		X linked osteoporosis with fractures	*
		Fibrous dysplasia of bone (Head)	*
VUmc	VUmc multidisciplinary (genetic) breast cancer team	Hereditary breast cancer	
VUmc	Down Center the Netherlands, location West	Down syndrome	
VUmc	VUMC PH Centre	Rare pulmonary hypertension; many subforms of PH; associated: collagen vascular disease, Hereditary PH, IPAH, type III WHO associated with emfysema.	
		Chronic thromboembolic pulmonary hypertension	
Vumc & AMC	Cystic Fibrosis Centre Amsterdam	Cystic Fibrosis	
VUmc	PCD-center Vumc	Primary ciliary dyskinesia	
VUmc	Fetal Akinesia Deformation Sequence Centre	Fetal Akinesia Deformation Sequence	
VUmc & AMC	Amsterdam Centre of congenital malformations	Hirschsprung Disease	
		Esophageal atresia	
		Anorectal malformation	
		Sacroccocygeal teratoma	
		Chronic intestinal pseudoobstruction	
VUmc & AMC	Amsterdam Center for ILD and sarcoïdosis	Sarcoïdosis	*
VUmc	Center on Atypical Sex or Gender development	Gender dysphoria	
		Disorder of sex development	
VUmc & LUMC & AMC	Center for Hereditary Retinal Diseases Leiden Amsterdam	Retinal dystrophy	
		Centrale sereuze chorioretinopathie	
NKI-AVL	Sarcoma Expertise Centre Amsterdam	Soft tissue sarcoma	
		Rare soft tissue tumor; Gastrointestinal stromal tumor (GIST)	
NKI-AVL	Expert Center of familial GI tumours	Genetic digestive tract tumor; 1. Hereditary nonpolyposis colon cancer 2. Familial adenomatous polyposis 3. Attenuated Familial adenomatous polyposis 4. Hereditary mixed poliposis syndrome	

UMC	Naam van Expertisecentrum	Cluster van / Specifieke aandoening(en)	
		Familial gastric cancer	
NKI-AvL& UMCU	Expert Center of Neuroendocriene carcinomas	Gastroenteropancreatic endocrine tumor	
NKI-AvL	Expert Centre of rare thoracic tumours	Mesotheloom	
NKI-AvL	Expert centre for rare urological diseases	Testicular cancer	
		Penile cancer	
NKI-AvL	Center of rare head and neck tumours	Rare otorhinolaryngologic disease; Tumours of the head and neck Rare otorhinolaryngologic disease; Tumours of the salivary glands	
NKI-AvL	Expert Center for Hereditary Cancer	Li-Fraumeni syndrome Hereditary breast and ovarian cancer syndrome	
NKI-AvL	<i>Rare Skin Cancer Center</i>	<i>Merkel Cell carcinoma</i>	
NKI-AvL	<i>Centre for rare nervous system tumor</i>	<i>Glial tumor</i>	
NKI-AvL	<i>Center of rare GI tumours</i>	<i>Epithelial tumor of anal cancer</i> <i>Rare gastroesophageal tumor</i>	*
Bartiméus	Bartiméus Diagnostisch Centrum	Genetic vitreous-retinal disease Congenital stationary night blindness Oculocutaneous or ocular albinism Cerebral visual impairment in children	
Kempenaeghe	Center of Sleep Medicine Kempenaeghe	Rare Sleep disorders	
The Rotterdam Eye Hospital	Rare Eye Disease Center Rotterdam	Uveal melanoma Central serous chorioretinopathy Rare acquired eye disease; Herpes simplex virus keratitis, stromal, neutrophic and endotheliitis Uveitis Rare genetic eye disease; Retinal Dystrophies	
STZ-Albert Schweitzer hospital, Dordrecht	Centre of expertise Retroperitoneal Fibrosis	Retroperitoneal fibrosis	
SZT-Medisch Spectrum Twente	<i>Gastrointestinal ischemia Centre</i>	<i>Celiac trunk compression syndrome</i>	
STZ-OLVG	Center for HME-MO (Hereditaire Multipele exostosen-Multipele Osteochondromen)	Multiple osteochondromas; Hereditary Multiple Exostoses - Multiple Osteochondromas	
STZ-OLVG	EC for interstitial lungdiseases OLVG	Idiopathic pulmonary fibrosis en Idiopathic interstitial pneumonia Exposure-related interstitial lung disease; in its broadest sense	
STZ-St. Antonius Ziekenhuis	Interstitial Lung Diseases Center of Excellence	Interstitial lung disease idiopathic pulmonary fibrosis (IPF) hypersensitivity pneumonitis (PH) sarcoidosis	
STZ-St. Antonius Ziekenhuis	Center for Pulmonary vascular diseases	Hereditary hemorrhagic telangiectasia Chronic thromboembolic pulmonary hypertension Idiopathic pulmonary arterial hypertension	

UMC	Naam van Expertisecentrum	Cluster van / Specifieke aandoening(en)	
STZ - St. Antonius Ziekenhuis	St. Antonius Oesofagus Centrum	Esophageal carcinoma	
STZ-CWZ Nijmegen	Malignant Hyperthermia investigation unit Nijmegen	Malignant hyperthermia	
STZ-CWZ Nijmegen	Center for Cerebrotendinous xanthomatosis	Cerebrotendinous xanthomatosis	
STZ-Maxima Medisch Centrum	SolviMáx, Center of Excellence for Abdominal Wall and Groin Pain	Anterior cutaneous nerve syndrome (ACNES)	
STZ-Maxima Medisch Centrum	Center for Adrenal Tumors	Adrenocortical carcinoma	
SZT-St. Elisabeth Ziekenhuis	Neurovascular Center Tilburg	Neurovascular malformation	
SZT-St. Elisabeth Ziekenhuis	Neurovascular Center Tilburg	Acquired aneurysmal subarachnoid hemorrhage	
STZ- St Elisabeth Hospital Tilburg	Neuro-oncology Center Tilburg	Glial tumor	
STZ-Jeroen Bosch Hospital	Center for Primary immunodeficiencies	Immuno-deficiency predominantly affecting antibody production, mainly the various types of unclassified antibody deficiency (voorlopig erkenning t/m sep 2017)	
SZT-Maasstad Hospital	Burn Centre Maasstad Hospital, in cooperation with Burn Centre Red Cross Hospital and Martini Hospital (ADBC: Association of Dutch Burn Care Centers)	Toxic epidermal necrolysis	
STZ-Medisch Centrum Haaglanden-Bronovo-Nebo	Center for Neuro-oncology The Hague	Gliomen	
PMC	Princess Máxima Center for pediatric solid tumors	(Pediatric) germ cell tumor	*
		(Pediatric) rare soft tissue tumor	*
		(Pediatric) rare renal tumor	*
		(Pediatric) rare digestive tumor	*
		Neuroblastoma	*
PMC	Princess Máxima Center for pediatric hematological malignancies and stem cell transplants	(Pediatric) tumor of hematopoietic and lymphoid tissues	*
		(Pediatric) myeloid hemopathy	*
		(Pediatric) lymphoid hemopathy	*
		(Pediatric) acute myeloid leukemia	*
		(Pediatric) acute lymphoblastic leukemia	*
		(Pediatric) lymphoma	*
		(Pediatric) myelodysplastic syndrome	*
PMC	Princess Máxima Center for pediatric brain tumors	(Pediatric) rare nervous system tumor	*
		(Pediatric) glial tumor	*
		(Pediatric) astrocytoma	*
		(Pediatric) medulloblastoma	*