

1. Record Nr.	UNINA9910349295903321
Autore	Chen Harold
Titolo	Atlas of genetic diagnosis and counseling // Harold Chen
Pubbl/distr/stampa	New York : , : Springer New York : , : Imprint : Springer, , 2020
ISBN	1-4614-6430-7
Descrizione fisica	1 online resource (LX, 2224 p. 2472 illus., 2018 illus. in color.)
Disciplina	611.01816 599.935
Soggetti	Genetic counseling Genetic disorders - Diagnosis Scientific atlases
Lingua di pubblicazione	Inglese
Formato	Materiale a stampa
Livello bibliografico	Monografia
Nota di contenuto	Acardia -- Achondrogenesis -- Achondroplasia -- Adams-Oliver syndrome -- Agnathia -- Aicardi syndrome -- Alagille syndrome -- Albinism -- Alpha thalassemia-mental retardation (ATR-X) syndrome -- Ambiguous genitalia -- Amniotic deformity, adhesions, mutilations (ADAM) syndrome -- Androgen insensitivity syndrome -- Angelman syndrome -- Apert syndrome -- Aplasia cutis congenita -- Arthrogryposis multiplex congenita -- Asphyxiating thoracic dystrophy -- Ataxia telangiectasia -- Atelosteogenesis -- Autism -- Bannayan-Riley-Ruvalcaba syndrome -- Beckwith-Wiedemann syndrome -- Behcet disease -- Biotinidase deficiency -- Bladder exstrophy -- Blepharophimosis, ptosis, and epicanthus inversus syndrome (BPES) -- Body stalk anomaly -- Brachydactyly -- Branchial cleft cyst -- Calcinosis cutis -- Campomelic dysplasia -- Carpenter syndrome -- Cat-eye syndrome -- Celiac disease -- Cerebral palsy -- Cerebrocosto-mandibular syndrome -- Charcot-Marie-Tooth disease -- CHARGE association -- Cherubism -- Chiari malformation -- Chondrodysplasia punctata -- Chromosome abnormalities in pediatric solid tumors -- Cleft lip/palate -- Cleidocranial dysplasia -- Cloacal exstrophy -- Clubfoot (talipes equinovarus) -- Collodion baby -- Congenital adrenal hyperplasia -- Congenital cutis laxa -- Congenital cytomegalovirus infection -- Congenital generalized lipodystrophy --

Congenital hemihyperplasia (congenital hemihypertrophy) --
 Congenital hydrocephalus -- Congenital hypothyroidism -- Congenital
 muscular dystrophy -- Congenital toxoplasmosis -- Conjoined twins
 -- Corpus-callosum agenesis/dysgenesis -- Craniometaphyseal
 dysplasia -- Cri-du-chat syndrome -- Crouzon syndrome -- Cutis
 marmorata telangiectatica congenita -- Cystic fibrosis -- Dandy-
 Walker malformation -- De Lange syndrome -- Del(18p) syndrome --
 Del(22q11.2) syndrome -- Del(Yq) syndrome -- Diabetic embryopathy
 -- Down syndrome -- Duncan syndrome (X-linked lymphoproliferative
 disease) -- Dyschondrosteosis/Langer mesomelic dysplasia --
 Dysmelia (limb deficiency/reduction) -- Dysplasia epiphysealis
 hemimelica -- Dystonia -- Dystrophinopathies -- EEC syndrome --
 Ehlers-Danlos syndrome -- Ellis-van Creveld syndrome --
 Enchondromatosis -- Epidermolysis bullosa -- Epidermolytic
 palmoplantar keratoderma -- Faciogenital (Aarskog) syndrome --
 Facioscapulohumeral muscular dystrophy (FSH) -- Familial
 adenomatous polyposis syndrome -- Familial hyperlysineuria --
 Familial Mediterranean fever -- Fanconi anemia -- Femoral hypoplasia-
 unusual facies syndrome -- Fetal akinesia syndrome -- Fetal alcohol
 syndrome -- Fetal hydantoin syndrome -- Fibrodysplasia ossificans
 progressiva -- Finlay-Marks syndrome -- Floppy infant -- Fragile X
 syndrome -- Fraser syndrome -- Freeman-Sheldon (whistling face)
 syndrome -- Friedreich ataxia -- Frontonasal dysplasia --
 Galactosemia -- Gastroschisis -- Gaucher disease -- Generalized
 arterial calcification -- Genitopatellar syndrome -- Giant congenital
 melanocytic nevi (giant congenital nevi) -- Glucose-6-phosphate
 dehydrogenase deficiency -- Glycogen storage disease, type II (Pompe)
 -- Goldenhar syndrome -- Gorlin (nevoid basal cell carcinoma)
 syndrome -- Greig cephalopolysyndactyly syndrome -- Hallermann-
 Streiff syndrome -- Harlequin fetus -- Hemophilia A -- Hereditary
 hearing loss -- Hereditary hemochromatosis -- Hereditary multiple
 exostosis -- Herlyn-Werner-Wunderlich syndrome --
 Holoprosencephaly -- Holt-Oram syndrome -- Huntington disease --
 Hydrolethrus syndrome -- Hydrops fetalis -- Hyper-IgE syndrome --
 Hypochondroplasia -- Hypoglossia-hypodactyly (oromandibular limb
 hypogenesis) syndrome -- Hypohidrotic ectodermal dysplasia --
 Hypomelanosis of Ito -- Hypophosphatasia -- Hypopituitarism -- I(1p),
 I(1q) syndrome -- Isodic(Yq) syndrome -- Incontinentia pigmenti --
 Infantile myofibromatosis -- Ivemark syndrome -- Jarcho-Levin
 syndrome -- Joubert syndrome -- Kabuki syndrome -- Kassback-
 Merritt syndrome -- KID syndrome -- Klinefelter syndrome -- Klippel-
 Feil syndrome -- Klippel-Trenaunay syndrome -- Kniest dysplasia --
 Larsen syndrome -- LEOPARD syndrome -- Lesch-Nyhan syndrome --
 Lethal multiple pterygium syndrome -- Loeys-Dietz syndrome -- Lowe
 syndrome -- Marfan syndrome -- McCune-Albright syndrome --
 Meckel-Gruber syndrome -- Megalencephalic leukoencephalopathy
 with subcortical cysts (van der Knaap disease) -- Menkes disease --
 Metachromatic leukodystrophy -- Miller-Dieker syndrome --
 Mitochondrial Leber hereditary optic neuropathy -- Mobius syndrome
 -- Mowat-Wilson Disease -- Mucopolysaccharidosis II (Hurler syndrome)
 -- Mucopolysaccharidosis III (pseudo-Hurler Polydystrophy) -- MPS I (Hurler
 syndrome) -- MPS II (Hunter syndrome) -- MPS III (Sanfilippo syndrome)
 -- MPS IV (Morquio syndrome) -- MPS VI (Maroteaux-Lamy syndrome)
 -- Multiple endocrine neoplasia Syndrome -- Multiple epiphyseal
 dysplasia -- Multiple pterygium syndrome -- Myotonic dystrophy --
 Nail-Patella Syndrome (hereditary Osteo-onychodysplasia) -- Neonatal
 Herpes simplex infection -- Nephrogenic diabetes insipidus --
 Netherton syndrome -- Neu-Laxova syndrome -- Neural tube defects

-- Neurofibromatosis 1 -- Neurofibromatosis 2 -- Noonan syndrome -- Oblique facial cleft syndrome -- Oligohydramnios sequence -- Omphalocele -- Oro-Facial-Digital Syndrome -- Osteogenesis imperfecta -- Osteopetrosis -- Osteopoikilosis -- Otopalatodigital spectrum disorders -- Pachyonychia congenita -- Pallister-Killian syndrome -- Phenylketonuria (PKU) -- Pierre Robin sequence -- Polycystic kidney disease, AD form -- Polycystic kidney disease, AR form -- Popliteal pterygium syndrome -- Prader-Willi syndrome -- Progeria -- Prune belly syndrome -- Pseudoachondroplasia -- R(18) syndrome -- Retinoid embryopathy -- Rett syndrome -- Rickets -- Rigid spine syndrome -- Roberts syndrome -- Robinow syndrome -- Rubinstein-Taybi syndrome -- Saethre-Chotzen syndrome -- Sagittal synostosis associated with chromosome abnormalities -- Schizencephaly -- Schmid metaphyseal chondrodystrophy -- Seckel syndrome -- Severe combined immune deficiency -- Short rib polydactyly syndromes (SRPS) -- Sickle cell disease -- Silver-Russell syndrome -- Sirenomelia -- Smith-Lemli-Optiz syndrome -- Smith-Magenis syndrome -- Sotos syndrome -- Spinal muscular atrophy -- Spondyloepiphyseal dysplasia -- Stickler syndrome -- Sturge-Weber syndrome -- Tay-Sachs disease -- Tetrasomy 9p syndrome -- Thalassemia -- Thanatophoric dysplasia -- Thrombocytopenia-absent radius (TAR) syndrome -- Treacher-Collins syndrome -- Trimethylaminuria -- Triploidy -- Trismus pseudocamptodactyly (Hecht syndrome) -- Trisomy 8 mosaicism (Warkany) syndrome -- Trisomy 13 syndrome -- Trisomy 18 syndrome -- Tuberous sclerosis -- Turner syndrome -- Twin-twin transfusion syndrome -- Ulnar-mammary syndrome -- Urofacial (Ochoa) Syndrome -- VATER (VACTERL) association -- Von Hippel-Lindau disease -- Waardenburg syndrome -- Weill-Marchesani syndrome -- Williams syndrome -- Wolf-Hirschhorn syndrome -- X-linked agammaglobulinemia (Bruton type) -- X-linked ichthyosis -- XX Male -- XXX syndrome -- XXXXX syndrome -- XXXXY syndrome -- XY female.-XYY syndrome.

Sommario/riassunto

Dr. Chen shares his almost 40 years of clinical genetics practice in a comprehensive pictorial atlas of almost 250 genetic disorders, malformations, and malformation syndromes. The author provides a detailed outline for each disorder, describing its genetics, basic defects, clinical features, diagnostic tests, and counseling issues, including recurrence risk, prenatal diagnosis, and management. Numerous color photographs of prenatal ultrasounds, imagings, cytogenetics, and postmortem findings illustrate the clinical features of patients at different ages, patients with varying degrees of severity, and the optimal diagnostic strategies. The disorders cited are supplemented by case histories and diagnostic confirmation by cytogenetics, biochemical, and molecular techniques, when available. The Atlas of Genetic Diagnosis and Counseling will help all physicians to understand and recognize genetic diseases and malformation syndromes and better evaluate, counsel, and manage affected patients. In this new edition, 47 additional genetic disorders are added, as well as extensive updates made to the previous disorders. New illustrations, as previous edition, will be supplemented by case and family history, clinical features, and laboratory data, especially molecular confirmation.

2. Record Nr.	UNINA9910786712903321
Autore	Wolf Diane L
Titolo	Beyond Anne Frank [[electronic resource]] : hidden children and postwar families in Holland / / Diane L. Wolf
Pubbl/distr/stampa	Berkeley, Calif. ; ; London, : University of California Press, c2007
ISBN	1-282-35846-4 1-4337-0131-6 1-4294-5576-4 9786612358463 0-520-93970-0
Descrizione fisica	1 online resource (419 p.)
Collana	S. Mark Taper Foundation imprint in Jewish studies Beyond Anne Frank
Disciplina	940.531808209492
Soggetti	Jews - Persecutions - Netherlands Hidden children (Holocaust) - Netherlands Holocaust, Jewish (1939-1945) - Netherlands Holocaust survivors - Netherlands Netherlands Ethnic relations
Lingua di pubblicazione	Inglese
Formato	Materiale a stampa
Livello bibliografico	Monografia
Note generali	Description based upon print version of record.
Nota di bibliografia	Includes bibliographical references and index.
Nota di contenuto	Front matter -- Contents -- Acknowledgments -- Introduction -- 1. The History and Memory of Hidden Children -- 2. Before and During the War: The Netherlands and the Jews -- 3. After the War: The Jews and the Netherlands -- 4. "My Mother Screamed and Screamed": Memories of Occupation, War, and Hiding -- 5. "I Came Home, but I Was Homesick": When Both Parents Returned -- 6. "They Were Out of Their Minds": When One Parent Returned -- 7. "Who Am I?": Orphans Living with Families -- 8. "There Was Never a Kind Word": Life in Jewish Orphanages -- 9. Creating Postwar Lives, Creating Collective Memory: From the Personal to the Political -- Conclusion -- Notes -- Glossary -- References -- Index
Sommario/riassunto	The image of the Jewish child hiding from the Nazis was shaped by Anne Frank, whose house-the most visited site in the Netherlands- has become a shrine to the Holocaust. Yet while Anne Frank's story

continues to be discussed and analyzed, her experience as a hidden child in wartime Holland is anomalous-as this book brilliantly demonstrates. Drawing on interviews with seventy Jewish men and women who, as children, were placed in non-Jewish families during the Nazi occupation of Holland, Diane L. Wolf paints a compelling portrait of Holocaust survivors whose experiences were often diametrically opposed to the experiences of those who suffered in concentration camps. Although the war years were tolerable for most of these children, it was the end of the war that marked the beginning of a traumatic time, leading many of those interviewed here to remark, "My war began after the war." This first in-depth examination of hidden children vividly brings to life their experiences before, during, and after hiding and analyzes the shifting identities, memories, and family dynamics that marked their lives from childhood through advanced age. Wolf also uncovers anti-Semitism in the policies and practices of the Dutch state and the general population, which historically have been portrayed as relatively benevolent toward Jewish residents. The poignant family histories in *Beyond Anne Frank* demonstrate that we can understand the Holocaust more deeply by focusing on postwar lives.

3. Record Nr.	UNINA9910780332103321
Autore	Rouse Tina I
Titolo	Exploring a vision [[electronic resource]] : integrating knowledge for food and health : a workshop summary / / by Tina I. Rouse and Debra P. Davis ; planning group for a workshop on Exploring a vision--integrating knowledge for food and health ; Board on Agriculture and Natural Resources, Division on Earth and Life Studies, National Research Council of the National Academies
Pubbl/distr/stampa	Washington, D.C., : National Academies Press, c2004
ISBN	0-309-16636-5 0-309-52705-8
Descrizione fisica	1 online resource (92 p.)
Altri autori (Persone)	DavisDebra P
Disciplina	363.8/0973
Soggetti	Nutrition - Research - United States Public health - Research - United States Food consumption - Health aspects - United States Health planning - United States Obesity - Research - United States
Lingua di pubblicazione	Inglese
Formato	Materiale a stampa
Livello bibliografico	Monografia
Note generali	Bibliographic Level Mode of Issuance: Monograph
Nota di bibliografia	Includes bibliographical references.