**Name of Journal:** *World Journal of Orthopedics*

**Manuscript NO:** 79620

**Manuscript Type:** EDITORIAL

**Genetics of congenital anomalies of the hand**

Kyriazis Z *et al*. Genetics of congenital anomalies

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**Author contributions:** Kyriazis Z wrote the paper and participated in the analysis and drafting of the literature data; Kollia P participated in the conception and design of the study, analysis and interpretation of the literature data, and critical revision of the article; Grivea I and Sotiriou S participated in analysis and interpretation of the literature data and critical revision of the article; Dailiana ZH participated in the conception and design of the study, analysis and interpretation of the literature data, and drafting and critical revision of the article; all authors actively reviewed and revised the manuscript and approved the finally submitted manuscript.

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**Received:** August 29, 2022

**Revised:** October 9, 2022

**Accepted:** October 14, 2022

**Published online:**

**Abstract**

Congenital anomalies of the hand are malformations occurring during the development of the human limb, and present as isolated disorders or as a part of a syndrome. During the last years, molecular analysis techniques have offered increasing knowledge about the molecular basis of hand malformations. Disturbances in the signaling pathways during the development of the upper limb result in malformations of the upper extremity. At present, several genes have been identified as responsible for hand anomalies and other have been recognized as suspect genes related to them. Different and new high throughput methods have been introduced for the identification of the gene mutations. In the current editorial, we summarize concisely the current molecular status of isolated hand genetic disorders and the recent progress in molecular genetics, including the genes related to the disorder. This progress improves the knowledge of these disorders and has implications on genetic counselling and prenatal diagnosis.

**Key Words:** Hand; Gene; Mutation; Molecular; Diagnosis; Disorders

Kyriazis Z, Kollia P, Grivea I, Sotiriou S, Dailiana ZH. Genetics of congenital anomalies of the hand. *World J Orthop* 2022; In press

**Core Tip:** The genetic basis of hand disorders is elucidated by the expansion of knowledge and introduction of molecular analysis techniques which contribute to the identification of new genes responsible for them. New genes and mutations are being isolated and correlated with the disorder based on the advances in sequencing technology, such as next generation sequencing and genetic consultation, and future therapeutic developments are enhanced. There appears to be a gap in the literature concerning the knowledge about the genetic basis of all hand disorders. The current molecular status of them is discussed and a summary of different genes, already identified or suspected to be related with them, is presented.

**INTRODUCTION**

Congenital anomalies of the hand very often have an autosomal dominant pattern of inheritance and most of them have a monogenic genetic basis with variable penetrance[1]. They are related with a disturbance of the normal procedure of the development of the limb, with diverse aetiology and variable clinical features, and their proposed classification system has changed throughout the years, incorporating a rather molecular than anatomic scope of the phenotypes[2]. Initially, the Swanson Classification of congenital anomalies of the hand, including nine categories of malformations, was used by the International Federation of Societies for Surgery of the Hand Committee on Congenital Conditions, as it was considered effective according to the knowledge at the time. Later on, as the awareness of pathogenetic routes and molecular basis of limb formation expanded, the object modeling technique classification was presented, dividing hand anomalies into four groups: Malformations (which include the majority of the disorders), deformations, dysplasias, and syndromes[3].

To date, several loci and disease-causing genes, including all four categories of hand disorders, have been identified in humans, and correlated to specific phenotypes.

Since these phenotype manifestations are indicators that the fetus or the newborn may suffer from a syndrome, the ability to identify the potential syndromes associated with these anomalies, is important for the clinician. Additionally, it is important to distinguish between syndromic and non-syndromic cases for reasons of genetic counselling. Therefore, we present a concise summary of the main genes that are responsible for the disorders, whose etiology is mainly based on known genetic and not external factors, and lead to hand disorder phenotypes when mutated.

**Polydactyly**

***Preaxial/Radial polydactyly***

Preaxial polydactyly is a malformation described by an extra digit on the radial side of the hand with an incidence as high as 1 in 3000 births. It follows an autosomal dominant inheritance model with reduced penetrance[4]. Thumb polydactyly has been further subdivided into six subtypes by Wassel according to the level of (metacarpal, proximal, or distal phalange) and the extent of duplication (partial and complete)[5,6]. Next generation sequencing (NGS) analysis in a patient with Wassel III polydactyly identified three gene mutations as follows: (1) Substitution c.1639 G > T in *RPGRIP1* gene; (2) Insertion of adenine in *TMEM216* gene; and (3) A > G nucleotide substitution (c.490) in *FBN1* gene. In a patient with Wassel IV duplication, the following mutations were identified: (1) Adenine duplicated in exon 45 of *CEP290* gene; (2) Two substitutions in *RPGRIP1* gene, c.1639 G > T and c.685 G > A; (3) Adenine insertion in *TMEM216* gene, c.432-11 432-10 insA; (4) Substitution G > C c.8249 in *MEGF8* gene; and (5) Substitution T > A c.548 in *CEP164* gene. These mutations are suspected to be involved in the formation of thumb duplication phenotype[7]. Another suspected way of development of preaxial polydactyly is the overexpression of *HES1* gene. The produced protein is considered to intervene in SHH/GLI signaling axis and results in the manifestation of preaxial polydactyly[8]. The disease gene locus with triphalangeal thumb was identified in chromosome 7q36[9]. Point mutations (105 C > G, 305A > T, 323T > C, 404G > A, 295T > C, 4909 C > T, 297G > A, 334T > G, 402C > T, and 545G > A) have been identified, and a 739A > G transition near the 5-end of the zone of polarizing activity regulatory sequence (*ZRS*)and a 621C > G mutation in the *ZRS* of the *LMBR1* gene have also been mapped[10,11]. Two more novel mutations (ZRS131A > T and ZRS474C > G) correlated with preaxial polydactyly were identified in a recent study of a Chinese family[12]. No mutations have been identified for index finger polydactyly, which is inherited with an autosomal dominant trait[13].

***Postaxial polydactyly***

Postaxial polydactyly presents with extra digits on the ulnar side of the hands. Mutations in genes *ZNF141, GLI3, IQCE, GLI1, FAM92A1, KIAA0825*, and *DACH1* have been isolated and their involvement in this manifestation is identified[14].

Responsible gene loci have been mapped to 7pl5-q11.23, 13q21-q32, 19p13.2-p13.1, 7q21-q34, and 13q13.3- 13q21.2 regions using genome-wide scan[15]. Subsequently, two heterozygous mutations, p.A765PfsX14 and p.R539TfsX12 in *GLI3* gene,and P.T474I mutation in the *ZNF141* gene have been identified using exome sequencing[16-18]. Recently, a new suspected mutation in *GLI1* gene (c.1133C > T) was isolated in an Indian family with the disorderand a mutation in *KIAA0825* gene has been isolated and suspected, although the role of the protein encoded by this gene in limb formation is still unclear[19,20].

***Central polydactyly and complex types of polydactyly***

Central polydactyly phenotype is characterized by duplication of the 2nd, 3rd, or 4th digit[21]. No disease causing locus or gene responsible for central polydactyly has been identified. Mirror image polydactyly is characterized by mirror-image duplication of fingers and toes[22].A mutation of the *MIPOL1* gene at 14q13 and two heterozygous deletions including the *PITX1* gene were identified[23].

**Split-hand malformation**

Split-hand malformation may occur as an isolated trait or accompanied with other defects. It manifests as a clinically heterogeneous disorder characterized by absent central digital rays, which result in median clefts of the hand. Responsible mutations map to chromosome 7q21.3-q22.1, chromosome Xq26, and chromosome 10q25[24]. *LBX1, BTRC, POLL, FBXW4,* and *BTRC* gene mutations are reported as responsible for the disease[25]. Recent molecular studies have expanded the list of suspected gene mutations. A *TP63* gene translocation and *FGFR1, BHLHA9, LRP6, UBA2,* and *WNT10B* gene mutations have been recently identified[26-30].

**Radial ray defects**

Radial ray defects occur as an isolated malformation or syndromic. They are characterized by partial or complete absence of radial ray structures. Radial defects comprise a large group of diseases. They are associated with *TBX3* gene, coding for a T-box transcriptional factor. *TBX3* is widely expressed in a variety of tissues including forelimbs and hindlimbs, epithelium of the mammary gland, the genital tubercle, and the uterus[31].

**Defects in Dorso-Ventral Patterning**

This disorder category involves nail-patella syndrome, which is autosomal dominant and is expressed with defects affecting the nails, skeleton, kidneys, and eyes. Loss of function mutations in the *LMXlB* gene lead to the syndrome[32,33]. *LMXlB* is involved in determination of dorso-ventral patterning of the limb. A mutation of *WIF1* gene has been isolated as a potential novel cause of the phenotype[34].

**Brachydactyly**

Brachydactyly phenotype may present as an isolated defect or in association with other malformations and refers to disproportionately short fingers and toes. Isolated brachydactylies usually occur as autosomal dominant traits and show a high degree of phenotypic variability. A locus on chromosome 5p13.3-p13.2 and the Indian hedgehog gene on chromosome 2q35-36 are involved in Type A1 brachydacty[35]. A mutation in the human bone morphogenetic protein receptor 1B gene (*BMPR1B*) on chromosome 4q can cause Type A2 brachydactyly. Mutations in growth/differentiation factor-5 gene (*GDF5*) alter the receptor binding affinities and can also cause symphalangism. No gene or locus for Type A3 brachydactyly has been identified. Type B Brachydactyly phenotype involves isolated mutations in the receptor kinase-like orphan receptor 2 gene (*ROR2*) on 9q22[36]. Type C Brachydactyly phenotype is considered to be caused by mutations in *GDF5*[37].

**Syndactyly**

Syndactyly is characterized by the fusion of soft and/or bony tissue of the fingers of the hand and is the most common congenital malformation of the hand in North America and Europe. Syndactyly can occur as an isolated malformation or as part of a syndrome. *HOXD13, FBLN1, GJA1, LMBR1, LRP4, GREM, FGF16,* and *BHLHA9* genesare incriminated for the disorder, when mutated[38]. Syndactyly type I presents as fusion between the middle and ring fingers. It is an autosomal dominant malformation and the most common type of syndactyly. Mutations in human chromosome 2q34-q36 have been isolated. Syndactyly type II is a dominantly inherited malformation. Its phenotype contains soft tissue syndactyly between the middle and ring fingers and sometimes clinodactyly or camptodactyly of the little finger. *HOXDl3* gene mutations are involved in this disorder[39]. Syndactyly type III affects the ring and little fingers with the middle phalanx of the little finger being absent or rudimentary. Mutations in Connexin 43 are involved in this type of syndactyly. Syndactyly type IV manifests as syndactyly of all fingers, and syndactyly type V as metacarpal synostosis. The genetic background of the last two types is not yet investigated[40].

**CONCLUSION**

Genetic hand disorders and their genetic heterogeneity and allelic heterogeneity between families indicate more complex mechanisms besides simple Mendelian inheritance. These mechanisms include underlying genetic, epigenetic, and environmental factors[41]. With the advent of NGS technology, including exome sequencing and whole-genome sequencing, new mutations causing hand malformations are isolated and the molecular pathogenesis is exposed. Systematic bioinformatics analysis of the responsible genes, using high-throughput sequencing, is a valuable tool in establishing the precise genotype–phenotype correlations of hand genetic disorders. Diagnosis is still largely postbirth, although prenatal diagnosis *via* molecular and genetic methodologies exists. The expansion of our knowledge related to the mutations leading to different phenotypes, with the use of next generation sequencing analysis, will contribute to prenatal diagnosis, prediction of operative treatment strategy, and potential future applications in gene therapy.

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**Footnotes**

**Conflict-of-interest statement:** All the authors have no conflict of interest related to the manuscript.

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**Provenance and peer review:** Invited article; Externally peer reviewed.

**Peer-review model:** Single blind

**Peer-review started:** August 29, 2022

**First decision:** September 26, 2022

**Article in press:**

**Specialty type:** Orthopedics

**Country/Territory of origin:** Greece

**Peer-review report’s scientific quality classification**

Grade A (Excellent): 0

Grade B (Very good): B

Grade C (Good): C

Grade D (Fair): 0

Grade E (Poor): 0

**P-Reviewer:** Chisthi MM, India; Kotlyarov S, Russia **S-Editor:** Wang LL **L-Editor:** Wang TQ **P-Editor:** Wang LL