Research Article

The Identification of Microdeletion and Reciprocal Microduplication in 22q11.2 Using High-Resolution CMA Technology

Ana Julia Cunha Leite,^{1,2} Irene Plaza Pinto,^{2,3} Damiana Mirian da Cruz e Cunha,² Cristiano Luiz Ribeiro,^{2,3} Claudio Carlos da Silva,^{1,2,3,4} Aparecido Divino da Cruz,^{1,2,3,4} and Lysa Bernardes Minasi^{1,2}

¹ Programa de Pós-Graduação (Mestrado) em Genética, Pontifícia Universidade Católica de Goiás, Rua 235,

No. 40, Setor Leste Universitário, 74605-050 Goiânia, GO, Brazil

²Núcleo de Pesquisas Replicon, Departamento de Biologia, Pontifícia Universidade Católica de Goiás, Rua 235,

No. 40, Setor Leste Universitário, 74605-050 Goiânia, GO, Brazil

³*Programa de Pós-Graduação em Biotecnologia e Biodiversidade, Universidade Federal de Goiás, Goiânia, GO, Brazil*

⁴Laboratório de Citogenética Humana e Genética Molecular, Secretaria do Estado da Saúde de Goiás, Goiânia, GO, Brazil

Correspondence should be addressed to Lysa Bernardes Minasi; minasilb@gmail.com

Received 18 December 2015; Accepted 13 March 2016

Academic Editor: Jonathan Muraskas

Copyright © 2016 Ana Julia Cunha Leite et al. This is an open access article distributed under the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

The chromosome 22q11.2 region has long been implicated in genomic diseases. Some genomic regions exhibit numerous low copy repeats with high identity in which they provide increased genomic instability and mediate deletions and duplications in many disorders. DiGeorge Syndrome is the most common deletion syndrome and reciprocal duplications could be occurring in half of the frequency of microdeletions. We described five patients with phenotypic variability that carries deletions or reciprocal duplications at 22q11.2 detected by Chromosomal Microarray Analysis. The CytoScan HD technology was used to detect changes in the genome copy number variation of patients who had clinical indication to global developmental delay and a normal karyotype. We observed in our study three microdeletions and two microduplications in 22q11.2 region with variable intervals containing known genes and unstudied transcripts as well as the LCRs that are often flanking and within this genomic rearrangement. The identification of these variants is of particular interest because it may provide insight into genes or genomic regions that are crucial for specific phenotypic manifestations and are useful to assist in the quest for understanding the mechanisms subjacent to genomic deletions and duplications.

1. Introduction

The copy number variations (CNVs) changes result in meiotic nonallelic homologous recombination (NAHR) between low copy repeats (LCR) that are often flanking these genomics rearrangements. The crossover mediated by NAHR can be interchromosomal or intrachromosomal [1]. Some genomic regions exhibit numerous LCRs with high identity sequences (95–99%) which provide increased genomic instability and mediate deletions and duplications in many disorders [2]. The microarray technology has increased the detection of CNVs and the diagnosis of patients with multiple congenital anomalies and intellectual disability.

The chromosome 22q11.2 region contains eight LCRs, designated from A to H. Four centromeric LCR22s (A to D) were related to reciprocal microduplications and DiGeorge/Velocardiofacial Syndromes (DGS/VCFS) (OMIM 188400 and 192430). On the other hand, telomeric LCR22s, named

LCR22D–H, located in a distal portion of 22q11.2 region, were related to distal microdeletions and reciprocal microduplications [3, 4].

DGS is the most common deletion syndrome with an incidence of 1: 4000 newborns and has a spectrum of clinical abnormalities that affects multiple systems, including cardio-vascular, neurological, psychiatric, endocrine, and immune systems. Palatal abnormalities and characteristic facial features also can be present [5]. Around 85–90% of individuals with DGS have the deletions of 3 Mb spanning LCR22A–D, while 8–10% have a 1.5 Mb deletion in LCR22A-B [6].

Most of the individuals with 22q11.2 microduplications carry approximately 3 Mb in length, among LCR22A and LCR22D, which are the reciprocal of the common deleted region found in DGS/VCFS, while few patients have 1.5 Mb duplications among LCR22A and LCR22B [7]. Larger duplications of 4 Mb and 6 Mb were also reported and involve LCRA–LCRE and LCRA–LCRG, respectively [8].

According to Portnoï [9], the frequency of 22q11.2 microduplication is approximately half the frequency of microdeletions which can be explained for the highly variable and mild phenotype leading to a low investigation of individuals affected. In addition, the phenotype of individuals with 22q11.2 microduplications can share features with DGS/VCFS, including heart defects, velopharyngeal insufficiency with or without cleft palate or hypernasal speech, and urogenital abnormalities. Here, we described five patients with phenotypic variability that carries deletions or reciprocal duplications at 22q11.2 detected by Chromosomal Microarray Analysis (CMA).

2. Materials and Methods

2.1. Biological Samples. All five participants had global developmental delay (GDD) without etiological diagnosis after undergoing a thorough clinical evaluation. Assistant physicians from the Goiás state public health system referred each patient to the genetic service at both the Laboratory of Human Cytogenetic and Molecular Genetics and the Biology Department at Pontifical Catholic University in Central Brazil. Parents or guardians signed the informed consent forms approved by the Ethics Committee on Human Research at the Pontifical Catholic University of Goiás (CEP-PUC/GO), under the protocol number 1721/2011.

For each proband and their biological parents, a total of 5 mL of peripheral blood was collected using a standard vacuum extraction blood-collecting system containing EDTA and heparin. Genomic DNA was isolated from whole blood using QIAamp DNA Mini kit (Qiagen, Germany), following the manufacturer's instructions. Conventional cell cultures, harvesting, and G-banding at the level of 550 bands were performed in all patients, following standardized procedures [10]. Chromosome observations were performed using a Zeiss Axioscope (Göttingen, Germany) and analyses using the IKAROS software (Metasystems Corporation, Altlussheim, Germany).

2.2. Chromosomal Microarray Analysis. Genomic DNA was obtained from peripheral blood from the probands and their

parents. The analyses were carried out on the probands and their biological parents to establish whether the DNA rearrangements were *de novo* or inherited. Total DNA (250 ng) for each sample was digested with *NspI*, ligated, PCR amplified and purified, fragmented, biotin-labeled, and hybridized for use in a GeneChipTM HD CytoScan Array (Affymetrix, Santa Clara, CA, USA). The array was designed specifically for cytogenetic research, including ≈2,696,550 copy number variation markers, 743,304 are single-nucleotide polymorphism markers, and >1,953,246 are nonpolymorphic markers. CEL files obtained by scanning the arrays were analyzed using the Chromosome Analysis Suite software (Affymetrix). Gains and losses that affected a minimum of 50 and 25 markers, respectively, in a 100 kb length were initially considered.

2.3. 22q11.2 LCR Structure Analysis. Using the segmental duplication track of the http://genome.ucsc.edu/ browser (Human Genome Build 36.1), we performed an analysis of duplicated genomic sequences including known LCRs (segmental duplication >1 kb of nonrepeat masked sequence with over 90% similarity), comparing the CNV size surrounding the proximal 22q11.2 locus 3 times (chr22: 18,640,000–25,080,000) against itself.

2.4. Clinical Report

Patient 1. Patient 1 was a 12-year-old female patient born to nonconsanguineous parents at 36-week gestation to a 42-year-old mother and 45-year-old father, and her birth weight was 3020 g. Child delivery was carried out through a caesarean section procedure. At 3 months of age, she had epilepsy and reflux. Physical examination of the proband revealed thumb brachydactyly, long finger, retrognathia, short philtrum, and large ear. The family history revealed that her oldest sister has heart malformation (Figure 1).

Patient 2. Patient 2 was a 15-year-old male patient born to nonconsanguineous parents, at 38-week gestation to a 38year-old mother and 38-year-old father, and his birth weight was 3200 g and crown-heel length 45 cm. Child delivery was carried out through a caesarean section procedure. After birth, the child showed cyanotic, did not cry, and did not perform the act of sucking. At the age of 14 years, he presented with seizures, cried a lot, had a slow development, and talked nonsense. Physical examination of the proband revealed hypertelorism, ear protuberant with attached lobes, blunted nose, broad nose and narrow bridge, short philtrum, short neck, thumb brachydactyly, clinodactyly of the fifth finger, prognathism, and permanent microdontia. The family history revealed that his aunt's grandmother had intellectual disability (Figure 2).

Patient 3. Patient 3 was a 7-year-old male patient born to nonconsanguineous parents, at 39-week gestation to a 30-year-old mother and 29-year-old father, and his birth weight was 3840 g and crown-heel length 51 cm. Child delivery was carried out through a caesarean section procedure. Child sat at eight months of age with the physical therapist's help, did not crawl, and had hypotonia. He started walking at one

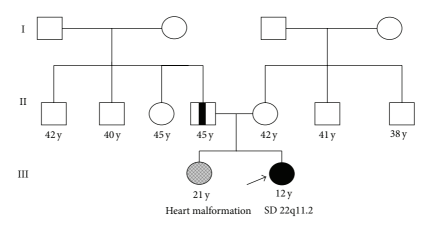


FIGURE 1: Pedigree of the patient 1 family.

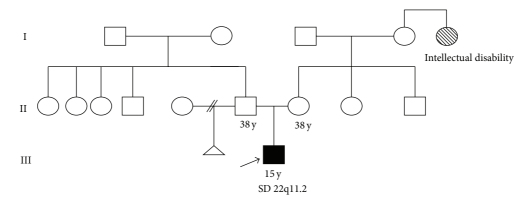


FIGURE 2: Pedigree of the patient 2 family.

year of age and only spoke at two years of age with the help of a speech therapist. Physical examination of the proband revealed retrognathia, low set ears, ear tags, ear protuberant with attached lobes, low nasal bridge, broad base of the nose, and epicanthal fold. He also had lowered orbital on the right side of the face and ptosis. The family history revealed that a maternal uncle was schizophrenic and paternal uncle had intellectual disability (Figure 3).

Patient 4. Patient 4 was a 14-year-old male patient born to nonconsanguineous parents, at 32-week gestation to a 35year-old mother and 39-year-old father, and his birth weight was 1600 g and crown-heel length 42 cm. Child delivery was carried out through a caesarean section procedure. Child was born with esophageal atresia and had no suction act and growth delay. His bone age is two years less than the biological age. Physical examination of the proband revealed hypertelorism, micrognathia, thumb brachydactyly, and asymmetric face. The family history revealed that a maternal third cousin was diagnosed with Torre Syndrome and a paternal cousin was born with a cleft lip (Figure 4).

Patient 5. Patient 5 was a 4-year-old female patient born to nonconsanguineous parents, at 39-week gestation to a 30-year-old mother and 34-year-old father, and her birth weight was 3140 g and crown-heel length 49 cm. She had jaundice. Child delivery was carried out through a caesarean

section procedure. Child sat at eight months and spoke at three years of age. She presents difficulty in walking and has low immunity. Physical examination of the proband revealed hypertelorism, epicanthic fold, low nasal bridge, micrognathia, and thin upper lip. The family history revealed that a father's niece has intellectual disability and difficulty in walking (Figure 5).

3. Results and Discussion

In the current study, we describe molecular findings in 5 probands with clinical diagnosis of global developmental delay, presenting microdeletion or microduplication in the apparently identical genomic region within 22q11.2 region and the LCRs which surround these regions. Samples were initially screened using G-banding karyotypes and showed no visible numerical and structural alterations (46,XX or 46,XY). After this analysis, we performed Chromosomal Microarray Analysis that demonstrated abnormalities of 22q11.2 region. All individuals in our study had deletion and duplication breakpoints that flanked or fell within previously characterized LCR22s. For the duplications breakpoints we only identified LCR-H within the 22q11.2 region in contrast with deletions breakpoints that presented LCR22s flanking the 22q11.2 region. The results from CMA and LCRs are shown in Table 1.

	LCR identity	%66 %66	%66 %66	96%	96%	96%	
TABLE 1: Clinical and molecular features of five probands screened by high-resolution CMA technology.	LCRs start/stop	B-D B-D	A-D A-D	After H	After H	After H	
	Origin**	Inherited pat	De почо	De поvо	Inherited mat	Inherited mat	
	Genes	ZNF74, SCARF2, MEDI5, SNAP29, CRKL, Inherited LZTRI, SLC7A4 pat	PRODH, DGCR5, DGCR9, DGCR10, DGCR2, DGCR11, DGCR14, GSC2, SLC25A1, CLTCL1, HIRA, MRPL40, UFD11, CLDN5, SEPT5, GP1BB, TBX1, GNB11, COMT, ARVCF, DGCR8, DGCR61, ZNF74, SCARF2, MED15, SNAP29, CRKL, LZTR1, SLC7A4, HIC2	IGLL3P, LRP5L, CRYBB2P1	MIR650	POMI2ILIP, GGTLC2, IGLL5, RTDR1, GNAZ, RAB36, BCR, FBXW4P1	
	Microarray nomenclature	22q11.21(20,716,902–21,465,662) × 1	22q11.21(18,916,842-21,800,797) × 1	22q11.23q12.1(25,656,237–25,930,479) × 1	22q11.23q12.1(25,656,237–25,994,326) × 3	22q11.22q11.23(22,962,196–23,652,512) × 3	
	Size (Mb)	0.75	2.88	0.27	0.34	0.69	ernal.
	Sex CNV Cytoband	Loss 22q11.21	Loss 22q11.21	Loss 22q11.21	M Gain 22q11.21 0.34	Gain 22q11.21 0.69	rnal; Pat: pat
	CNV	Loss	Loss	Loss	Gain	Gain	Mat: mate
	Sex	щ	W	Μ	Μ	щ)elay; ** l
	Age (yo)	12	15	9	14	4	omental I
	Clinical features*	GDD	GDD	GDD	GDD	GDD	*GDD: Global Developmental Delay; **Mat: maternal; Pat: paternal.
	Case		7	3	4	2J	*GDD: (

olo	
ц	
ech	
7	
CM/	
5	
n	
īti	
olt	
res	
h-r	
/ hig	
y l	
dþ	
ne	
ee	
SCL	
Is	
n.	
ba	
of five probands screened by hi	
ve	
f five	
ofo	
res	
ature	
22	
ur fe	
ula	
Sec	
Ы	
lπ	
nc	
linical and mol	
ica	
Clini	
Ö	
3LE 1: C	
3LE	

BioMed Research International

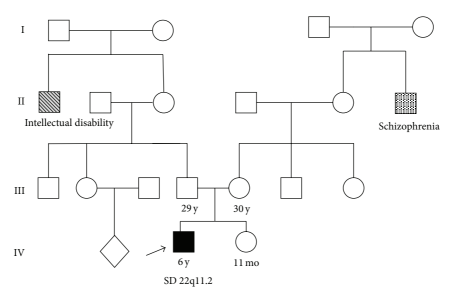


FIGURE 3: Pedigree of the patient 3 family.

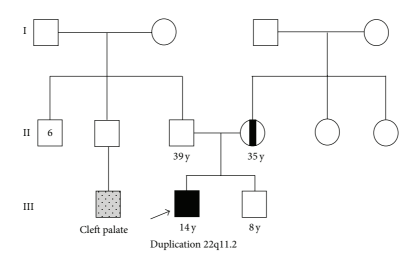
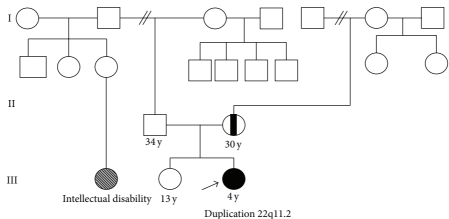


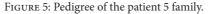
FIGURE 4: Pedigree of the patient 4 family.

We identify three microdeletions at 22q11.2 region from patients 1, 2, and 3 (Figure 6). In patient 2, the CMA showed a de novo 22q11.2 microdeletion with 2.88 Mb, flanked by LCR-A and LCR-D, which involves more than 70 genes, including TBX1 that is suppressed. The majority of individuals have a 3 Mb deletion whose proximal region contains the presumed disease associated with T-box transcription factor 1 (TBX1) gene [11-16]. TBX1 gene plays a vital role during development and deletion of this gene leads to a variety of craniofacial and cardiac structures [17]. Rump et al., in 2014, [18] highlighted that loss of CRKL combined with loss of either TBX1 or MAPKI in individuals with A–D or larger deletions results in higher rates of cardiac defects than in those individuals with central deletions that involve only CRKL gene. Another gene present in this region is the catechol-O-methyltransferase (COMT) gene that catalyzes one of the major degradative pathways of the catecholamine transmitters, and one copy of this gene leads to abnormal regulation of catechol-Omethyltransferase levels in the brain. Researchers believe that

changes involving this enzyme in the prefrontal cortex may help explain the increased risk of behavioral problems and mental illness associated with 22q11.2 deletion Syndrome [19, 20].

The CMA from patient 1 showed a paternal origin 0.75 Mb deletion at 22q11.2, between LCR-B and LCR-D, classified by Rump et al., 2014 [18] in a Central Group of LCRs. Atypical or distal microdeletions between LCR-B and LCR-D were identified in a limited number of studies [21–26]. Patient 3 demonstrated a *de novo* 0.27 Mb microdeletion at 22q11.2 and we identified only LCRs within this CNV. There are three genes deleted at 0.27 Mb microdeletion: *IGLL3P*, *LRP5L*, and *CRYBB2P*. Haploinsufficiency in seven genes was observed in 0.75 Mb microdeletion (Table 1) that were associated with DGS [15, 27–29]. The *CRKL* gene has also been implicated in the underlying molecular mechanism of 22q11 deletion syndrome. Interestingly, CRKL was mapped in the distal half of the typical deleted region, suggesting that it may be responsible for those cases with the most distal





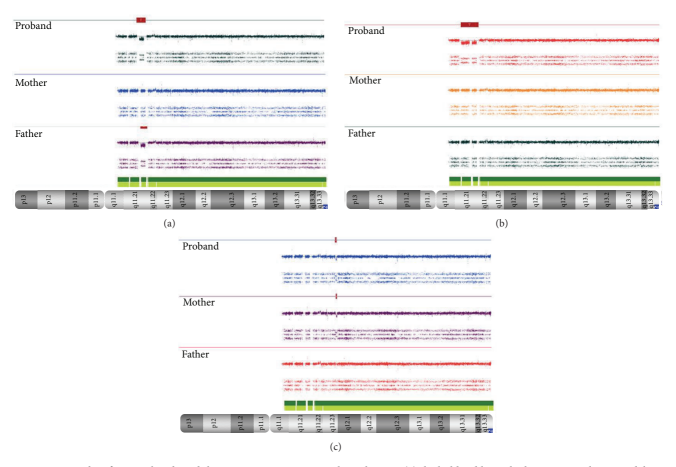


FIGURE 6: CMA data from probands and their parents at 22q11.2. Analysis showing (a) the bold red line which represents the microdeletion from proband 1 and her parents, (b) 2.88 Mb of de novo microdeletion from proband 2 and his parents, and (c) inherited maternal microdeletion from proband 3 and his parents.

deletions that do not involve TBX1 gene [30, 31]. TBX1 is the important gene related to the 22q11.2 deletion syndrome and the haploinsufficiency of this gene is presented in most of the phenotype characteristics. However, this gene is not always included in all 22q11.2 deletions, like patients 1 and 3, and many authors suggest that there are regulatory elements or

modifier genes mapped far away from this region which may affect the TBX1 expression [30].

According to Carelle-Calmels et al., 2009, [32] deletion at 22q11.2 was usually sporadic and was reported to be inherited in 6 to 28% of patients with these syndromes. Different from what was observed in patient 1, inherited microdeletions were

BioMed Research International

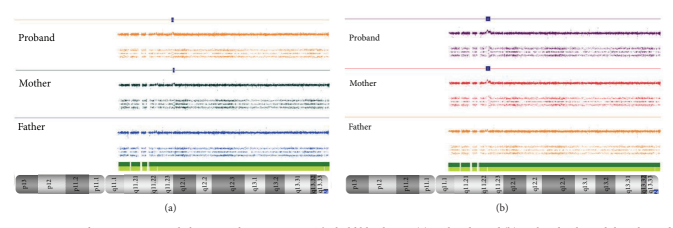


FIGURE 7: CMA depicts genomic imbalances in chromosome 22. The bold blue line in (a) proband 4 and (b) proband 5 showed the inherited maternal microduplication at 22q11.2.

commonly of maternal origin, and these findings occur due to the recombination rate at 22ql1.2 being 1.6 to 1.7 times greater in females than in males [33]. Furthermore, when inherited, these microdeletions had parent-of-origin effect, and phenotypic variability was observed [9].

Patients 1 and 3 presented global developmental delay and ears anomalies; developmental delay or cognitive impairment is reported in over 80% of patients but tends to be relatively mild, with no severe affect and the typical facial gestalt of the proximal microdeletion syndrome was absent. Moreover, ear anomalies are observed in most distal microdeletion and they are nonspecific [11–14].

Additionally, we observed two patients, 4 and 5, who harbor microduplication in 22q11.2 region. Patients 4 and 5 showed maternally inherited duplication of 0.34 Mb and 0.69 Mb, respectively (Figure 7). Both microduplications are located within the block of LCR22-F. *MIR650* gene is located at 0.34 Mb duplication and the potential role of miRNAs in the pathogenesis of duplication phenotypes has to be better investigated. In a study by Merico et al., 2014, [34], the authors showed that the deleted miRNAs in 22q11.2 are involved with the regulation of expression of genes in several developmental pathways which could be affected by miRNAs reduced levels, and the dysregulation of miRNA processing could be due to the haploinsufficiency of *DCGR8* gene [6, 35].

In general, duplications confer milder phenotypes and are more likely to be inherited than their reciprocal deletions, with approximately 93% of small duplications [36]. Although genomic imbalances inherited from a clinically normal parent are usually considered benign CNVs, parents of a child with an inherited chromosome abnormality may sometimes show mild variations of the child's phenotype, such as what have been reported for the 22q11.2 deletion syndrome [2].

Careful clinical assessment of both child and parents is crucial to understand the causative role of duplications at 22q11.2. Considering the unaffected phenotypes of parents, the presence of a genetic modifier has been proposed for other syndromes with variable expressivity. In this context, a genetic modifier is characterized by a combination of CNVs at the same or different loci inherited from parents in whom the single variation was insufficient to cause the disease [37]. Genomic imbalances inherited from phenotypically unaffected parents may contribute to the progeny phenotype through variable penetrance or expressivity, or both, through epigenetic effects, response to environmental factor challenge, modulated epistasis, or stochastic cellular events during fetal development. Moreover, genomic imbalances may uncover a recessive mutation on the nondeleted allele [5, 8, 9].

We need to understand more regarding the reasons for similarities features and widely variable phenotype for individuals with 22q11.2 deletions and duplications. According to Zhang and Lupski, 2015, [38] these variable phenotypes suggested the possible existence of some modifying genetic factors in addition to the large genomic CNVs and recent progress has shown that the SNPs and CNVs in noncoding regions can be the genetic modifiers.

We identified two groups of LCR22 that are flanking the genomic rearrangements from patients 1 and 2. These findings suggest that the occurrence of genomic rearrangements might be mediated by NAHR between the LCRs, increasing the susceptibility to the generation of CNVs. On the order hand, patients 3, 4, and 5 showed LCRs within the reported deletion and duplications breakpoints; thus, it is not possible to speculate the detailed mechanism in relation to how this deletion and duplication initially appeared in these patients. Although high density of long and short interspersed nuclear elements (LINEs and SINEs) around the proximal breakpoint may have been contributory, detailed sequence data characterizing the breakpoints are necessary to recognize these elements and to propose the mechanisms of the genomic rearrangements.

4. Conclusions

We observed in our study variable intervals containing known genes and unstudied transcripts as well as the LCRs that are within and often flanking this genomic rearrangement. The recognition of LCRs provides important insights related to the role of genomic architecture in chromosomal rearrangements, chromosome evolution, and human disease. This report further illustrates the potential for determination of small microduplications and microdeletions through of the use of Chromosomal Microarray Analysis. These array-based methods would allow more sensitive and rapid breakpoint localization without the need for multiple FISH experiments. Besides, the applications of microarray allow the identification of distal deletions and also facilitate breakpoint identification in the proximal 22q11.2 deletions and duplications. The identification of these variants is of particular interest because it may provide insight into genes or genomic regions that are crucial for specific phenotypic manifestations and are useful to assist in the quest for understanding the mechanisms subjacent to genomic deletions and duplications.

Competing Interests

The authors declare that they have no competing interests.

Authors' Contributions

Aparecido Divino da Cruz and Lysa Bernardes Minasi conceived and designed the experiments. Ana Julia Cunha Leite, Irene Plaza Pinto, Cristiano Luiz Ribeiro, and Damiana Mirian da Cruz e Cunha performed the experiments. Irene Plaza Pinto, Lysa Bernardes Minasi, Ana Julia Cunha Leite, and Aparecido Divino da Cruz analyzed the data. Ana Julia Cunha Leite, Aparecido Divino da Cruz, Irene Plaza Pinto, and Lysa Bernardes Minasi wrote the paper.

Acknowledgments

The authors thank Dr. Rinaldo Wellerson Pereira from Catholic University of Brasília for his assistance and for coordinating the ExeGenS, Rede de Excelência em Genética e Genômica Molecular Aplicada a Saúde Humana. Their gratitude goes to each patient and his/her family for taking part in this study. This work was supported by the CAPES and FAPEG.

References

- E. Blennow, K. Lagerstedt, H. Malmgren, S. Sahlén, J. Schoumans, and B. M. Anderlid, "Concurrent microdeletion and duplication of 22q11.2," *Clinical Genetics*, vol. 74, no. 1, pp. 61– 67, 2008.
- [2] J. Coppinger, D. McDonald-Mcginn, E. Zackai et al., "Identification of familial and de novo microduplications of 22q11.21q11.23 distal to the 22q11.21 microdeletion syndrome region," *Human Molecular Genetics*, vol. 18, no. 8, pp. 1377–1383, 2009.
- [3] T. H. Shaikh, H. Kurahashi, and B. S. Emanuel, "Evolutionarily conserved low copy repeats (LCRs) in 22q11 mediate deletions, duplications, translocations, and genomic instability: an update and literature review," *Genetics in Medicine*, vol. 3, no. 1, pp. 6– 13, 2001.
- [4] F. M. Mikhail, R. D. Burnside, B. Rush et al., "The recurrent distal 22q11.2 microdeletions are often de novo and do not represent a single clinical entity: a proposed categorization system," *Genetics in Medicine*, vol. 16, no. 1, pp. 92–100, 2014.
- [5] R. E. Ensenauer, A. Adeyinka, H. C. Flynn et al., "Microduplication 22q11.2, an emerging syndrome: clinical, cytogenetic, and

molecular analysis of thirteen patients," *The American Journal of Human Genetics*, vol. 73, no. 5, pp. 1027–1040, 2003.

- [6] R. D. Burnside, "22q11.21 deletion syndromes: a review of proximal, central, and distal deletions and their associated features," *Cytogenetic and Genome Research*, vol. 146, pp. 89–99, 2015.
- [7] A. Alberti, C. Romano, M. Falco et al., "1.5 Mb de novo 22q11.21 microduplication in a patient with cognitive deficits and dysmorphic facial features," *Clinical Genetics*, vol. 71, no. 2, pp. 177–182, 2007.
- [8] Z. Ou, J. S. Berg, H. Yonath et al., "Microduplications of 22q11.2 are frequently inherited and are associated with variable phenotypes," *Genetics in Medicine*, vol. 10, no. 4, pp. 267–277, 2008.
- M.-F. Portnoï, "Microduplication 22q11.2: a new chromosomal syndrome," *European Journal of Medical Genetics*, vol. 52, no. 2-3, pp. 88–93, 2009.
- [10] R. Verma and A. Babu, Human Chromosomes: Principles and Techniques, McGraw-Hill, 2nd edition, 1995.
- [11] A. Rauch, R. A. Pfeiffer, G. Leipold, H. Singer, M. Tigges, and M. Hofbeck, "A novel 22q11.2 microdeletion in DiGeorge syndrome," *The American Journal of Human Genetics*, vol. 64, no. 2, pp. 659–666, 1999.
- [12] F. M. Mikhail, M. Descartes, A. Piotrowski et al., "A previously unrecognized microdeletion syndrome on chromosome 22 Band q11.2 encompassing the BCR Gene," *The American Journal* of Medical Genetics A, vol. 143, no. 18, pp. 2178–2184, 2007.
- [13] S. Ben-Shachar, Z. Ou, C. A. Shaw et al., "22q11.2 distal deletion: a recurrent genomic disorder distinct from DiGeorge syndrome and velocardiofacial syndrome," *The American Journal* of Human Genetics, vol. 82, no. 1, pp. 214–221, 2008.
- [14] O. K. Rødningen, T. Prescott, A.-S. Eriksson, and O. Røsby, "1.4 Mb recurrent 22q11.2 distal deletion syndrome, two new cases expand the phenotype," *European Journal of Medical Genetics*, vol. 51, no. 6, pp. 646–650, 2008.
- [15] S. E. Racedo, D. M. McDonald-Mcginn, J. H. Chung et al., "Mouse and human CRKL is dosage sensitive for cardiac outflow tract formation," *The American Journal of Human Genetics*, vol. 96, no. 2, pp. 235–244, 2015.
- [16] C. R. Fagerberg, J. Graakjaer, U. D. Heinl et al., "Heart defects and other features of the 22q11 distal deletion syndrome," *European Journal of Medical Genetics*, vol. 56, no. 2, pp. 98–107, 2013.
- [17] H. Parker, E. Conway, J. Goldsberry, S. Jeffries, E. Price, and J. T. Oxford, "Genetic and molecular aspects of DiGeorge syndrome," *Bio One, The Biologists' Forum*, vol. 86, no. 2, pp. 109–117, 2015.
- [18] P. Rump, N. de Leeuw, A. J. van Essen et al., "Central 22q11.2 deletions," *American Journal of Medical Genetics—Part A*, vol. 164, no. 11, pp. 2707–2723, 2014.
- [19] M. J. Zeitz, P. P. Lerner, F. Ay et al., "Implications of COMT long-range interactions on the phenotypic variability of 22q11.2 deletion syndrome," *Nucleus*, vol. 4, no. 6, pp. 487–493, 2013.
- [20] S. Yu, W. D. Graf, and R. J. Shprintzen, "Genomic disorders on chromosome 22," *Current Opinion in Pediatrics*, vol. 24, no. 6, pp. 665–671, 2012.
- [21] L. Fernández, J. Nevado, F. Santos et al., "A deletion and a duplication in distal 22q11.2 deletion syndrome region. Clinical implications and review," *BMC Medical Genetics*, vol. 10, article 48, 2009.

- [22] L. Garavelli, S. Rosato, A. Wischmeijer et al., "22q11.2 Distal deletion syndrome: description of a new case with truncus arteriosus type 2 and review," *Molecular Syndromology*, vol. 2, no. 1, pp. 35–44, 2011.
- [23] S. Ledig, C. Schippert, R. Strick, M. W. Beckmann, P. G. Oppelt, and P. Wieacker, "Recurrent aberrations identified by array-CGH in patients with Mayer-Rokitansky-Küster-Hauser syndrome," *Fertility and Sterility*, vol. 95, no. 5, pp. 1589–1594, 2011.
- [24] S. Yu, W. D. Graf, A. Ramalingam et al., "Identification of copy number variants on human chromosome 22 in patients with a variety of clinical findings," *Cytogenetic and Genome Research*, vol. 134, no. 4, pp. 260–268, 2011.
- [25] J. Breckpot, B. Thienpont, M. Bauters et al., "Congenital heart defects in a novel recurrent 22q11.2 deletion harboring the genes CRKL and MAPKI," *American Journal of Medical Genetics A*, vol. 158, no. 3, pp. 574–580, 2012.
- [26] S. Sanna-Cherchi, K. Kiryluk, K. E. Burgess et al., "Copynumber disorders are a common cause of congenital kidney malformations," *The American Journal of Human Genetics*, vol. 91, no. 6, pp. 987–997, 2012.
- [27] P. Ravassard, F. Côté, B. Grondin, M. Bazinet, J. Mallet, and M. Aubry, "ZNF74, a gene deleted in DiGeorge syndrome, is expressed in human neural crest-derived tissues and foregut endoderm epithelia," *Genomics*, vol. 62, no. 1, pp. 82–85, 1999.
- [28] D. M. McDonald-McGinn, S. Fahiminiya, T. Revil et al., "Hemizygous mutations in SNAP29 unmask autosomal recessive conditions and contribute to atypical findings in patients with 22q11.12Ds," *Journal of Medical Genetics*, vol. 50, no. 2, pp. 80– 90, 2013.
- [29] H. Kurahashi, K. Akagi, J. Inazawa et al., "Isolation and characterization of a novel gene deleted in DiGeorge syndrome," *Human Molecular Genetics*, vol. 4, no. 4, pp. 541–549, 1995.
- [30] S. Garcia-Miñaur, J. Fantes, R. S. Murray et al., "A novel atypical 22q11.2 distal deletion in father and son," *Journal of Medical Genetics*, vol. 39, no. 10, p. E62, 2002.
- [31] D. L. Guris, J. Fantes, D. Tara, B. J. Druker, and A. Imamoto, "Mice lacking the homologue of the human 22q11.2 gene CRKL phenocopy neurocristopathies of DiGeorge syndrome," *Nature Genetics*, vol. 27, no. 3, pp. 293–298, 2001.
- [32] N. Carelle-Calmels, P. Saugier-Veber, F. Girard-Lemaire et al., "Genetic compensation in a human genomic disorder," *The New England Journal of Medicine*, vol. 360, no. 12, pp. 1211–1216, 2009.
- [33] M. Delio, T. Guo, D. M. McDonald-McGinn et al., "Enhanced maternal origin of the 22q11.2 deletion in velocardiofacial and DiGeorge syndromes," *The American Journal of Human Genetics*, vol. 92, no. 3, pp. 439–447, 2013.
- [34] D. Merico, G. Costain, N. J. Butcher et al., "MicroRNA dysregulation, gene networks, and risk for schizophrenia in 22q11.2 deletion syndrome," *Frontiers in Neurology*, vol. 5, article 238, 2014.
- [35] L. M. Brzustowicz and A. S. Bassett, "miRNA-mediated risk for schizophrenia in 22q11.2 deletion syndrome," *Frontiers in Genetics*, vol. 3, article 291, 2012.
- [36] J. D. Weisfeld-Adams, L. Edelmann, I. K. Gadi, and L. Mehta, "Phenotypic heterogeneity in a family with a small atypical microduplication of chromosome 22q11.2 involving TBX1," *European Journal of Medical Genetics*, vol. 55, no. 12, pp. 732– 736, 2012.
- [37] J. Liao, L. Kochilas, S. Nowotschin et al., "Full spectrum of malformations in velo-cardio-facial syndrome/DiGeorge

syndrome mouse models by altering Tbx1 dosage," *Human Molecular Genetics*, vol. 13, no. 15, pp. 1577–1585, 2004.

[38] F. Zhang and J. R. Lupski, "Non-coding genetic variants in human disease," *Human Molecular Genetics*, vol. 24, no. 1, pp. R102–R110, 2015.