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Article

Chronic Pulmonary Aspergillosis: Genomic Variant Analysis and Protein Dysfunction Susceptibility in a Brazilian Cohort

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Abstract: Chronic pulmonary aspergillosis (CPA) is a debilitating condition often affecting immunocompetent patients with underlying structural lung diseases, particularly pulmonary tuberculosis. This study investigates single nucleotide variants (SNVs) in immunogenetic-related genes among a Brazilian cohort with CPA. Twelve patients with confirmed CPA, based on ESCMID/ERS criteria, were sequenced using NGS technology. Variants were annotated, classified using ACMG guidelines, and analyzed for potential impact on protein interactions and immune pathways. A set of SNVs in *CX3CR1*, *IL12B*, *IL4R*, *PTX3*, *CCR5* and *IFNG* genes were classified as variants of uncertain significance (VUS), but protein–protein interaction analysis suggests a potential role in immune evasion and dysfunction. This is the first study to apply a custom multigenic panel for CPA susceptibility in a Brazilian cohort, contributing to future functional and clinical studies in fungal immunogenetics.

Keywords: CPA; immune genetic; *Aspergillus*

1. Introduction

Chronic pulmonary aspergillosis (CPA) is a progressive and potentially fatal fungal infection primarily caused by *Aspergillus spp.*, an opportunistic fungus commonly found in the environment. Its clinical manifestation is influenced by both host immune status and the presence of pre-existing pulmonary sequelae(1-3). Immunocompromised individuals—such as those undergoing solid organ or bone marrow transplantation—are particularly vulnerable to invasive aspergillosis(4). However, CPA predominantly affects immunocompetent or non-neutropenic patients, especially those with chronic lung conditions such as tuberculosis sequelae(5, 6).

CPA encompasses several clinical subtypes, including simple aspergilloma, chronic cavitary pulmonary aspergillosis, subacute invasive aspergillosis, *Aspergillus* nodules, and chronic fibrosing pulmonary aspergillosis. Diagnosis relies on a combination of radiological evidence, immunoprecipitation antibody titers, and clinical symptoms, which is especially critical in low-resource settings(7-10).

Despite treatment efforts, CPA is associated with significant morbidity and mortality, often progressing to pulmonary fibrosis and severely impacting quality of life (11, 12). Globally, an estimated 3 million individuals are affected by CPA annually, with Brazil reporting approximately 112,000–160,000 new cases per year and a 5-year mortality rate ranging from 38% to 85%(13).

Genetic predisposition may influence susceptibility to CPA. Single nucleotide polymorphisms (SNPs) are common in the general population and can result from evolutionary adaptation, spontaneous mutations, or environmental exposure(14, 15). While not all genetic mutations result in pathogenic protein alterations, certain SNPs may affect immune response pathways. Genome-wide association studies (GWAS) have identified variants in immune-related genes, such as PTX3, associated with impaired neutrophil function and increased susceptibility to fungal infections(5, 16, 17).

Given the limited literature on host–pathogen genetic interactions in fungal infections, this study aims to investigate genomic variants in Brazilian patients diagnosed with CPA. Using next-generation sequencing (NGS) and American College of Medical Genetics (ACMG) classification guidelines, we identified and evaluated genetic variants that may contribute to susceptibility to CPA.

2. Materials and Methods

This study enrolled Brazilian patients receiving outpatient follow-up in the Infectious Diseases and Pulmonology departments at the Hospital das Clínicas, University of São Paulo (HC-FMUSP) during the period of 2023, with a confirmed diagnosis of chronic pulmonary aspergillosis (CPA). All patients with CPA had the following characteristics, in accordance with the ESCMID/ERS criteria(18): (1) the computerized tomography scan of the chest findings suggestive of aspergillosis (one or more cavities with or without a fungal ball present or nodules); (2) microbiological evidence of *Aspergillus* infection [microscopy or culture from sputum, bronchoalveolar lavage (BAL) or, biopsy, histology, positive serum or BAL galactomannan] or serological evidence (serum immunodiffusion test and counterimmunoelectrophoresis); (3) All present for at least 3 months or at least a 1-month duration in SAIA; (4) Exclusion of alternative diagnoses, such as tuberculosis, malignancy, and other similar pathological conditions.

A total of 12 patients (n=12) were included, of whom 8 were male (66.7%) and 4 were female (33.3%). All patients presented post-tuberculosis pulmonary. The clinical subtypes observed in the cohort were: chronic cavitary pulmonary aspergillosis in 8 patients (66.7%), simple aspergilloma in 3 patients (25.0%), and chronic fibrosing pulmonary aspergillosis in 1 patient (8.3%). The exclusion criteria for the study were: immunosuppressive conditions (eg. HIV infection, malignancies, prolonged corticosteroid administration, diabetes mellitus, and cirrhosis) and confirmed genetic diseases.

DNA Extraction

Peripheral blood (4 mL) was collected in EDTA-containing tubes. All participants signed informed consent forms (CAAE: 76601023.1.0000.0068). DNA was extracted at the Cytogenomics Laboratory, Department of Pathology, Faculty of Medicine, University of São Paulo, using the *QIAamp DNA Blood Mini Kit* (Qiagen®, Hilden, Germany), following the manufacturer's protocol. A final volume of 30 µL was obtained for subsequent analyses, and DNA concentration and quality were measured using a *Qubit fluorometer* (Thermo Fisher Scientific®).

Next-Generation Sequencing (NGS)

Exome libraries were prepared using the *Illumina® DNA Prep with Exome 2.0 Plus Enrichment* kit, according to the manufacturer's instructions. Sequencing was performed using the *NovaSeq 6000 System* (Illumina®, 2025). Raw data were generated in Variant Call Format (VCF) and annotated by comparison with the reference genome GRCh38/hg38 using bioinformatics tools.

Variant Filtering and Classification

A custom-designed virtual multigenic panel was developed including genes previously reported to be associated with susceptibility to fungal infections. The panel included: *CCR5, CX3CR1, IFNG, IFNGR2, IL10, IL12A, IL12B, IL13, IL4, IL4R, CXCL8, CXCR1, CXCR2, MBL2, MIF, NOS3, PTX3,*

and ARNT2. VCF files from each participant were analyzed using the Franklin by Genoox® platform (<https://franklin.genoox.com>). Allele frequencies were obtained from the gnomAD v4.1.0 and ABraOM databases (Brazilian Online Archive of Mutations). Table 1.

Variants classified as single nucleotide variants (SNVs) were evaluated using the American College of Medical Genetics and Genomics (ACMG) guidelines, considering allele frequency, in silico prediction tools, and mutation type. The criteria PM2_SUPP and BP4 were applied, and most variants were categorized as variants of uncertain significance (VUS). Table 2

Additional analyses included computational modeling of amino acid substitutions and chemical dissimilarity to assess potential protein dysfunction. Protein–protein interaction networks and immune pathway enrichment were evaluated for implicated genes.

3. Results

3.1. Variant Analysis and Classification

Variant analysis was guided by the development of a virtual multigenic panel consisting of genes previously associated with susceptibility to fungal infections (Table 1). The selected genes were: CCR5, CX3CR1, IFNG, IFNGR2, IL10, IL12A, IL12B, IL13, IL4, IL4R, CXCL8, CXCR1, CXCR2, MBL2, MIF, NOS3, PTX3, and ARNT2. VCF files containing the sequencing data of each participant were uploaded to the Franklin by Genoox® platform for variant annotation and classification. Global allele frequencies were obtained from gnomAD v4.1.0, while Brazilian-specific frequencies were retrieved from the ABraOM database.

3.2. ACMG Guidelines

Variants were evaluated using the ACMG criteria, including absence in population databases, in silico pathogenicity predictions, and variant type. Most SNVs were classified under the criteria PM2_SUPP + BP4 as variants of uncertain significance (VUS) (Table 2). Due to inconsistencies in in silico predictors, we further assessed the potential for structural and functional disruption by analyzing the chemical dissimilarity of substituted amino acids. This analysis supported the hypothesis of potential protein dysfunction. Additional evaluation using PPI networks revealed possible impacts on antifungal immune mechanisms.

3.3. Protein–Protein Interaction and Pathway Enrichment

PPI network analysis was performed using STRING v12.0, incorporating genes that showed VUS with potential immunological relevance (PTX3, CX3CR1, IL4R, IL12B, IFNG, CCR5). Significant enrichment was observed (PPI enrichment p-value < 0.05), suggesting interactions relevant to antifungal defense. Figure 1.

Biological processes identified included: cytokine-mediated signaling pathway (GO:0019221), macrophage activation in immune response (GO:0002281), response to fungal pathogens (GO:0009620), positive regulation of phagocytosis (GO:0050766), host modulation of symbiont processes (GO:0051851), and regulation of cytokine production (GO:0001817).

Relevant KEGG pathways included: cytokine–cytokine receptor interaction (FDR: 1.26e-06), Th1 and Th2 cell differentiation (FDR: 0.00019), and the JAK-STAT signaling pathway (FDR: 0.00071).

Table 1. Virtual multigenic panel. Column 1: name of genes. Column 2: Reference of literature Column 3: SNP ID. Column 4: Related to infections.

Gene	Reference	SNP ID	Related to infections
<i>MBL</i>	PMID: 37558798	rs11003125	yes

<i>IL4</i>	PMID: 26667837	rs2243248	yes
<i>IL4R</i>	PMID: 27708669	rs3024656	yes
<i>IL8</i>	PMID: 26667837	rs2227307	yes
<i>CXCR1</i>	PMID: 26667837	rs2234671	yes
<i>CXCR2</i>	PMID: 26667837	rs1126580	yes
<i>PTX3</i>	PMID: 33240991	rs3816527	yes
<i>CX3CR1</i>	PMID: 31964743	rs9823718; rs7631529	yes
<i>MIF</i>	PMID: 36166743	NOT INFORMED	yes
<i>CCR5</i>	PMID: 26667837	rs1799987; rs2734648	yes
<i>IFNγR1</i>	PMID: 37327531	rs2234711	yes
<i>IFNγ</i>	PMID: 26667837	rs2069705	yes
<i>IL13</i>	PMID: 26667837	rs1800925	yes
<i>IL12B</i>	PMID: 26667837	rs3212227C	yes
<i>NOS3</i>	PMID: 38407762	rs1549758	yes
<i>ARNT2</i>	PMID: 31964743	rs1374213	yes

Table 2. Variant classification. Column 1: Name of gene. Column 2: SNP ID. Column 3: type of molecular alteration found. Column 4: Classification according ACMG. Column 5: Level of mutation damage. Column 6: rediction of the effect of substitutions between amino acids based on chemical properties..

Gene	SNP (ID number)	Alteration	Classification	prediction tools	Grantham distance
<i>CX3CR1</i>	rs555964469	missense	VUS	0.303	Conservative (29)
<i>IL12B</i>	rs1245834629	missense	VUS	0.0580	Moderately conservative (56)
<i>IL4R</i>	rs780006435	missense	VUS	0.0850	Moderately conservative (58)
<i>PTX3</i>	rs138818541	missense	VUS	0.237	Conservative (43)
<i>CCR5</i>	rs1800940	missense	VUS	0.154	Moderately radical (110)
<i>IFNG</i>	rs76012457	missense	VUS	0.0340	Moderately conservative (94)

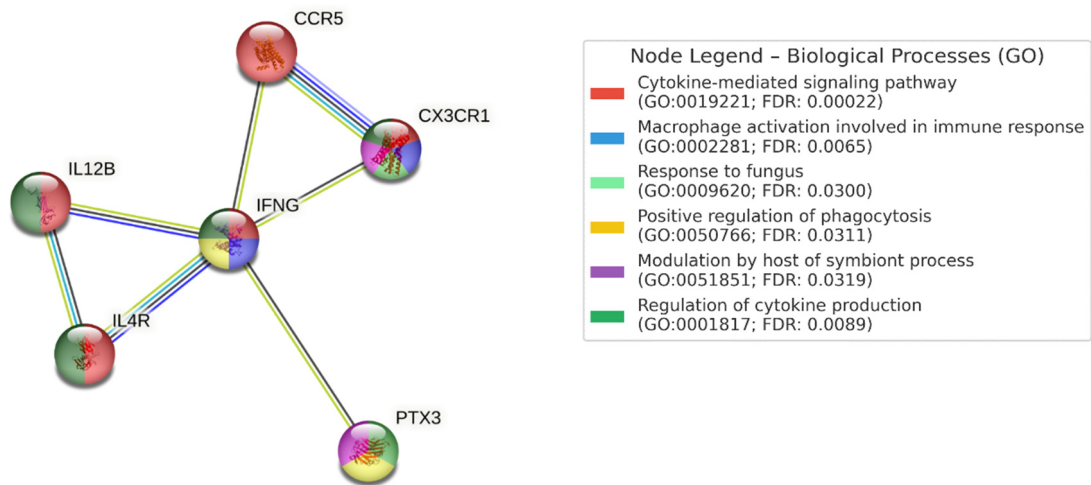


Figure 1. Number of nodes: 6 number of edges: 7 average node degree: 2.33 avg. local clustering coefficient: 0.867 expected number of edges: 1 PPI enrichment p-value: 6.27e-05 (0.0000627).

4. Discussion

The primary clinical feature associated with progressive chronic *Aspergillus spp.* infection is structural damage caused by previous pulmonary diseases, such as tuberculosis. Interestingly, most affected individuals do not present significant alterations in leukocyte or lymphocyte lineages, which raises questions about immune system functionality in immunocompetent hosts. Effective immune responses require adequate signaling pathways. Although humoral immunity is not yet fully understood in fungal infections, it is clear that the absence of humoral signaling compromises the recruitment of cellular immune mechanisms. Pathogen recognition is mediated by soluble molecules that participate in immune cascades, including opsonization, phagocytic activation, and direct pathogen neutralization. This immunological synergy—particularly against *Aspergillus conidia*—requires the coordinated action of pattern recognition molecules and cellular immunity.

PTX3 is a soluble pattern recognition molecule that plays a vital role in recruiting macrophages and neutrophils to sites of inflammation. Its multifunctional nature includes modulating inflammation, tissue remodeling, and complement activation. Previous studies have shown that polymorphisms in *PTX3* are associated with increased susceptibility to fungal infections in non-neutropenic patients due to impaired conidial opsonization. In our study, the variant rs138818541 in *PTX3* was identified in one patient (8.3%). Although classified as a variant of uncertain significance (VUS) by ACMG guidelines, emerging evidence suggests a deleterious impact of such variants. Similarly, the rs555964469 variant in *CX3CR1* was also found in one patient. This chemokine receptor is predominantly expressed in natural killer cells, cytotoxic CD8+ T cells, macrophages, and monocytes. As leukocytes are essential in fungal clearance, defects in receptor function may compromise host defense. Studies like that of Lupiañez et al. (2020) have shown that macrophages are key players in eliminating fungal pathogens, and alterations in their activation may contribute to CPA persistence. The pulmonary epithelium consists of alveolar epithelial cells type I and II (ATI and ATII). While ATII cells retain regenerative potential in adults, aging and chronic infections impair their functionality. The sustained recruitment of macrophages and inflammatory mediators, including profibrotic signals, can exacerbate CPA progression. Our cohort's average age of 48 years may reflect environmental and age-related impacts on epithelial repair. Interestingly, the most frequently observed variant in our study was located in the *IL4R* gene (75% of patients). Although this gene is widely studied in allergic bronchopulmonary aspergillosis (ABPA), its role in CPA is less established. IL-4 is essential for Th2 cell differentiation, and abnormal expression of IL4R is implicated in Th2-skewed diseases. However, given the high allele frequency of most *IL4R* variants

in the Brazilian population (ABraOM > 0.5), they were excluded from ACMG classification—except for the rare rs780006435 variant (allele frequency < 0.01), which met classification criteria. The presence of these variants suggests a molecular mechanism underlying increased vulnerability to CPA. While some in silico predictors did not classify them as deleterious, additional analyses of amino acid property changes suggested structural and functional alterations in the encoded proteins. Our study is limited by its small sample size and the scarcity of published functional studies on host genetics in chronic fungal infections. Moreover, the Brazilian population's genetic diversity complicates variant interpretation due to limited genome databases. Although we used ABraOM as a reference, it remains underpowered compared to European datasets. Nevertheless, this is the first study to apply a custom multigenic panel to a Brazilian CPA cohort. Our findings underscore the need for further functional studies and biomarker development, including quantification of PTX3 and MBL2 in clinical settings. Ultimately, integrating genetic insights into CPA management may improve diagnostic accuracy and therapeutic outcomes.

5. Conclusions

This exploratory study highlights the potential role of rare genomic variants in immune-related genes among patients with chronic pulmonary aspergillosis. Although the variants identified were classified as VUS, structural and network analyses indicate possible impacts on protein function and antifungal immune responses. This research represents the first application of a multigenic panel in a Brazilian cohort with CPA and emphasizes the importance of integrating genomic data into the investigation of host susceptibility to fungal diseases. Future studies with larger cohorts and functional validation are essential to better understand the molecular basis of CPA and to advance personalized approaches for its diagnosis and treatment.

6. Contributions

R. S. Mendes, V. F. Oliveira, A. N. Costa and M. M. C. Magri were involved in writing—review and editing, formal analysis, methodology; B.M. Wolff, L.L. Vieira, M.R. Costa Siemann, Karina Marinho Nascimento and L. S. Rolim helped in methodology—Next Generation Sequencing, review; Y. Gasparini, and G.F.S. Carvalho, review; E. A. Moura helped in NGS analysis; L.D. Kulikowski took charge of project administration and funding acquisition.

7. Ethics Declarations

Ethics approval and consent to participate. The ethics committee of the University of São Paulo approved this study (HC-FMUSP—CAPPesq 76601023.1.0000.0068. The patients or parents of the patients signed the consent form for participation in the study.

Consent for publication: All authors have agreed to have this article published.

Competing interests: The authors declare no competing interests.

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