The Role of Interleukin 10 Genetic Variations in Pulmonary Tuberculosis: Perspectives of Genetics, Pathogenesis and Immunology

By Debie Anggraini

181

HEME: Health and Medical Journal

pISSN: 2685 – 2772 eISSN: 2685 – 404x

Available Online at: https://jurnal.unbrah.ac.id/index.php/heme/issue/view/50

The Role of Interleukin 10 Genetic Variations in Pulmonary Tuberculosis: Perspectives of Genetics, Pathogenesis and Immunology

Anggraini, D.1,*

35

Clinical Pathology Departement, Faculty of Medicine, Baiturrahmah University, Padang *debieanggraini@fk.unbrah.ac.id

Abstract

Pulmonary tuberculosis remains a significant global public health challenge. In 674 ts to overcome this disease, a deeper understanding of the role of individual genetics, such as IL-10 genetic variation, in the re 56 nse to M. tuberculosis infection is critical. Research that has been conducted shows that IL-10, which has an im 20 ant role in regulating the immune response, can also influence the development of TB. Genetic variations in the IL-10 gene play a role in determining the extent of the immune response to TB infection and an individual's risk of this disease. The interaction between Treg cells, IL-10, and TB is also an important aspect in the pathogenesis and management of TB. Although Treg cells and IL-10 have a role in controlling excessive inflammation, too much of either can dampen the immune response needed to overcome infections. The implication of this research is that the development of more targeted and personalized therapy is an important step in overcoming TB. The use of individual genetic knowledge, such as IL-10 genetic variations, can help design more effective therapies and improve patient prognosis. However, challenges such as drug resistance and the complexity of 30 etic-immunological interactions remain challenges that need to be overcome in TB management. Overall, this study shows the importance of involving the fields of genetics and immunolog 4n global efforts to address pulmonary tuberculosis. With a deeper understanding of the 5 ctors that influence the immune response to TB infection, we can hope to develop more effective strategies in the prevention, diagnosis and treatment of this disease and reduce the burden of TB worldwide.

Keywords: Interleukin 10; Pulmonary Tuberculosis; Genetic Variation, Pathogenesis; Immunology

Abstrak

Tuberkulosis paru tetap menjadi tantangan global yang signifikan dalam bidang kesehatan masyarakat. Dalam upaya untuk mengatasi penyakit ini, pemahaman lebih dalam tentang peran genetika individu, seperti variasi genetik IL-10, dalam respon terhadap infeksi M. tuberculosis sangat penting. Penelitian yang telah dilakukan menunjukkan bahwa IL-10, yang memiliki peran penting dalam mengatur respon imun, juga dapat memengaruhi perkembangan TB. Variasi genetik dalam gen IL-10 memainkan peran dalam menentukan sejauh mana respons imun terhadap infeksi TB dan risiko individu terhadap penyakit ini. Interaksi antara sel Treg, IL-10, dan TB juga merupakan aspek penting dalam patogenesis dan pengelolaan TB. Meskipun sel Treg dan IL-10 memiliki peran dalam mengendalikan peradangan yang berlebihan, terlalu banyak dari keduanya dapat meredakan respons imun yang diperlukan untuk mengatasi infeksi. Implikasi penelitian ini adalah bahwa pengembangan terapi yang lebih terarah dan personalisasi adalah langkah penting dalam mengatasi TB. Penggunaan pengetahuan genetik individu, seperti variasi genetik IL-10, dapat membantu merancang terapi yang lebih efektif dan meningkatkan prognosis pasien. Namun, tantangan-tantangan seperti resistensi obat dan kompleksitas interaksi genetik-imunologi tetap menjadi tantangan yang perlu diatasi dalam pengelolaan TB. Dalam keseluruhan, penelitian ini menunjukkan pentingny 67 elibatkan bidang genetika dan imunologi dalam upaya global untuk mengatasi tuberkulosis paru. Dengan pemahaman yang lebih dalam tentang faktor-faktor yang memengaruhi respons imun terhadap infeksi TB, kita dapat berharap untuk mengembangkan strategi yang lebih efektif dalam pencegahan, diagnosis, dan pengobatan penyakit ini serta mengurangi beban TB di seluruh dunia.

Kata Kunci: Interleukin 10; Tuberkulosis Paru; Variasi Genetika, Patogenesis; Imunologi

Email: heme@unbrah.ac.id

I. Introduction

40

Pulmonary tuberculosis is one of the most deadly infectious diseases in the world, caused by Mycobacterium tuberculosis. Despite great efforts to control it, pulmonary tuberculosis remains a significant global health challenge. This disease 4 affects millions of people every year and is the main cause of death due to bacterial infections. Therefore, a deeper understanding of the mechanisms of pathogenesis, individual susceptibility to this disease, and the development of more effective therapies are essential. Amid scientific efforts to understand the factors that influence pulmonary tuberculosis, the role of genetics in susceptibility to infection and disease progression is a topic of increasing interest. Variations 43 human genes have long been known to play a key role in determining the response to infection and the development of disease, including pulmonary tuberculosis. One of the genes that is the center of attention in this research is the interleukin 10 (IL-10) gene.¹

IL-10 is a cytokine that has an important role in regulating the body's immune regionse. This is a key factor in maintaining a balance between pro-inflammatory (fighting infection) and anti-inflammatory (preventing excessive tissue damage) immune responses. Therefore, IL-10 is particularly relevant in the context of pulmonary tuberculosis, where an appropriate immune response is key in controlling M. tuberculosis infection, but at the same time, excessive inflammation can cause serious lung damage. Genetic polymorphisms in IL-10 have been an important area of research in understanding the role of genetics in pulmonary tuberculosis. Genetic polymorphisms are variations in DNA sequences that can affect the expression or function of certain genes. These variations may influence how efficiently IL-10 regulates the immune response and its impact on pulmonary tuberculosis infection. Therefore, in this

literature review, we will investigate the role of genetic variation in IL-10 in the context of pulmonary tuberculosis, focusing on understanding it from the viewpoint of genetics, pathogenesis, and immunology.^{2,3}

31

It is important to note that the role of IL-10 in pulmonary tuberculosis is complex. On the one hand, IL-10 can help in controlling excessive inflammation that can damage lung tissue, but on the other hand, too much IL-10 can inhibit an effective immune response against M. tuberculosis. Thus fore, genetic variations in IL-10 can have a significant impact on the development and outcome of pulmonary tuberculosis in different individuals. In this literature review, v24 will explore various aspects related to the role of IL-10 genetic variations in pulmonary tuberculosis. We will begin by providing background on pulmonary tuberculosis as a significant global disease, as well as the important role of interleukin 10 in the immune system. We will also provide an explanation of the genetics of IL-10, including the structure of the gene and the types of genetic polymorphisms that can influence the expression and function of this gene.4,5

In the next section, we will discuss current efforts in the development of therapeutics immunotherapy for pulmonary tuberculosis. We will discuss therapeutic approaches being developed and how knowledge of IL-10 genetics can be used to design more effective therapies Thus, this literature review will provide comprehensive understanding of the role of IL-10 genetic variation in pulmonary tuberculosis, with a focus on genetics, pathogenesis, and immunology perspectives. It is hoped that this knowledge will provide a strong foundation for further research and development of more effective therapies for this disease.

II. INTERLEUKIN 10 (IL-10) GENETICS

A. IL-10 GENE STRUCTURE

The interleukin 10 (IL-10) gene is an important component of the human immune system and has a significant role in regulating immune responses inflammation. To understand the genetic role of IL-10 in pulmonary tuberculosis, we need 55 understand the basic structure of this gene. The IL-10 gene is located 75 chromosome 1 in humans, more precisely on the long arm of chromosome 1 (1q31-q32). This gene consists of five exons separated by four introns. This basic structure is the basis for IL-10 protein synthesis, which in turn influences various aspects of the immune response.1,6

IL-10 protein is an anti-inflammatory cytokine produced by various types of immune cells, including T cells, B cells, macrophages, and dendritic cells. The main function of IL-10 is to reduce inflammation and calm excessive immune responses. It achieves this by inhibiting the activation of cells that participate in pro-inflammatory immune responses and by inhibiting the production of pro-inflammatory cytokines such as interleukin 1 (IL-1), interleukin 6 (IL-6), and nuclear factor kappa B (NF- κB). The IL-10 gene structure also contains a control region that regulates gene expression. Polymorphisms within these regions may influence how efficiently the IL-10 gene is expressed and, consequently, how efficiently IL-10 functions in regulating immune and inflammatory responses.^{7,8}

B. IL-10 GENETIC POLYMORPHISM

Zenetic polymorphisms are variations in DNA sequence that can be found between individuals in different populations. These variations can affect gene expression and the function of the proteins produced by those genes. In the context of IL-10, genetic polymorphisms are important factors that can

individual's response influence an infection and the development of pulmanary tuberculosis. There are several polymorphisms that have been identified in the IL-10 gene, and some of the most important are the polymorphisms at positions -1082 (referred to as -1082 G/A), -819 (referred to 63-819 C/T), and -592 (referred to as as -592 C/A) in the IL-10 gene promoter control region. This polymorphism may influence IL-10 expression levels.

The -1082 G/A polymorphism has been associated with differences in IL-10 production, where in iduals with the -1082 A allele tend to have highers L-10 production than individuals with the -1082 G allele. C/A has also been associates with varying levels of IL-10 production. Genetic variations in IL-10 maganfluence the body's ability to regulate the immune response to M. tuberculosis. Some studies have shown that psividuals with certain polymorphisms in the IL-10 gene may have a higher risk of developing pulmoury tuberculosis or may have a different immune response 152 M. tuberculosis infection. In addition, IL-10 genetic polymorphisms can also influence the effectiveness of therapy and response to treatment. Therefore, further understanding of IL-10 genetic polym48 hisms is critical in efforts to develop more targeted and personalized therapeutic approaches in the treatment of pulmonary tuberculosis. 7,8,10

C. TYPES OF GENETIC POLYMORPHISMS IN IL-10

Genetic polymorphisms in the interleukin 10 (IL-10) gene include variations in the DNA sequence that can be found between individuals in different populations. In the context of IL-10, there are several types of genetic polymorphisms that have been identified and studied in depth.

First, there is a polymorphism at position - 1082 in the IL-10 gene promoter control region, which is often referred to as -1082

G/A. This polymorphism can influence the level of IL-10 expression, where individuals ith the -1082 A allele tend to have higher IL-10 production than individuals with the -1082 G allele. Second, there is a polymorphism at position -819, called -19 C/T, as well as at position -592, called -592 C/A, also in the promoter region of the IL-10 gene. 11,12

Variations in these two ositions have also been associated with differences in IL-10 production. For example, individuals with the -819 T allele and -592 A allele tend to have lower levels of IL-10 production. Additionally, there are other genetic polymorphisms that can occur in different regions of the IL-10 gene, which can also influence IL-10 expression and function. The combination of these polymorphisms can create complex genetic diversity populations, which can have a significant impact on an individual's immune response to infection and the development of disease, including pulmonary tuberculosis.8,10

D. Geographic and Ethnic Distribution of Il-10 Polymorphisms

Genetic polymorphisms in IL-10 not only vary between individuals, but can also vary in geographic and ethnic distribution. Various human populations around the world different 146 erns of genetic polymorphisms in the IL-10 gene. The geographic distribution of IL-10 polymorphisms may be reflected in the allelic variations of the gene in different ethnic groups in different regions of 42 world. Studies have shown that the frequencies of different alleles in the IL-10 gene can differ between different populations. This could be the result of evolution and environmental differences in different regions, which can influence patterns of genetic variation populations. 13,14

Additionally, the geographic distribution of IL-10 polymorphisms may also have implications in susceptibility to certain diseases. Populations with certain genetic polymorphisms may have different risks of developing diseases such as pulmonary tuberculosis. Therefore, understanding the geographic and ethnic distribution of IL-10 polymorphisms may provide important insights into disease epidemiology and differences in response to infection based on genetic and geographic factors. 12,15,16

E. ASSOCIATION OF IL-10 GENETIC POLYMORPHISMS WITH SUSCEPTIBILITY TO PULMONARY TUBERCULOSIS

One of the most important aspects of research on IL-10 genetic polymorphisms is their relationship to individual susceptibility pulmonary tuberculosis. Genetic polymorphisms in IL-10 have been a focus of research because of their potential role in regulating the immune response against Mycobacterium tuberculosis, the bacterium causes pulmonary tuberculosis. Epidemiological and genetic studies have shown that in iduals with certain polymorphisms in the IL-10 gene may have a higher risk of developing pulmonary tuberculosis. These genetic variations may influence how the body responds to M. tuberculosis infection, and in some cases, individuals with polymorphism 26 that affect lower IL-10 production may have a better ability to control the infection.

However, the association between IL-10 polymorphisms and susceptibility to pulmonary tuberculo 18 remains the subject of active research. The role of IL-10 in regulating complex immune responses makes it an interesting focal point in understanding the pathogenesis of this disease. Therefore, further research is needed to better understand how genetic variation in IL-10 may influence an individual's risk of pulmonary tuberculosis and its potential

implications in the development of more targeted therapies. 12,15

III. PATHOGENESIS OF PULMONARY TUBERCULOSIS

A. MECHANISM OF MYCOBACTERIUM TUBERCULOSIS INFECTION

understand pathogenesis the pulmonary tuberculosis, it is important to look at how Mycobacterium tuberculosis (M. tuberculosis), the bacteria that causes this disease, infects and persists in the human body. The infection process begins when a person is exposed to M. tuberculosis through the air, for example through coughing or sneezing from an infected individual. After entering the body, M. tuberculosis reaches the lungs and spreads into the alveoli, the small units where gas exchange occurs. This is where these bacteria begin to multiply and infectious lesions form known tuberculomas. M. tuberculosis has thick cell walls and a lipid content that makes it resistant to the acidic environment inside macrophages, the first defense cells in the immune system.5

Macrophages are immune cells that are phagocytic, meaning they can engulf and digest pathogens such as bacteria. When macrophages engulf M. tuberculosis, the bacteria are not always completely destroyed. In contrast, some M. tuberculosis can live in macrophages and inhibit damaging immune responses. This is one way M. tuberculosis avoids detection and destruction by the immune system. During initial infection, pulmonary tuberculosis may not show significant symptoms. This is a form of infection known as latent tuberculosis, in which the bacteria remain in the body but do not cause active disease. Part of the reason why M. tuberculosis can persist in the body is its ability to modulate the immune response.16

B. IMMUNE RESPONSE TO MYCOBACTERIUM TUBERCULOSIS

The immune response to M. tuberculosis involves various components of the immune system, including immune cells that play a role in fighting infection. When M. tuberculosis enters the lungs, macrophages are one of the first defense cells that try to destroy the bacteria. Macrophages recognize M. tuberculosis through molecular patterns known as pattern recognition. They recognize bacterial cell components, such as lipopolysaccharide (LPS), peptidoglycan, and other substances, which are danger signals to the immune system. When macrophages recognize M. tuberculosis, they ivate an immune response by producing pro-inflammatory cytokines such interleukin-12 (IL-12) and interferon-gamma $(IFN-\gamma)^{-15-17}$

IL-12 is a cytokine that triggers the activation of T helper type 1 (Th1) cd 41. Th1 cells, which also produce IFN-γ, play an important role in coordinating the intersponse against M. tuberculosis. IFN-γ plays a central role in activating macrophages to destroy M. tuberculosis. When active, macrophages try to bind and digest bacteria more effectively. Apart from Th1 cells, cytotoxic T cells (CTL) also have a role in destroying cells infected by M. tuberculosis. CTL can recognize infected cells and damage them directly. All of these components work together to try to control the infection.

C. THE ROLE OF IL-10 IN THE REGULATION OF THE IMMUNE RESPONSE TO M. TUBERCULOSIS

Although this immune response 44 an important part of the body's defense against M. tuberculosis, the bacterium has strategies to evade the damaging immune response. One way M. tuberculosis inhibits immune 22 ponses is through the production of interleukin 10 (IL-10). IL-10 is an anti-

inflammatory cytokine produced by various types of immune cells, including macrophages, T cells, and B cells. The main function of IL-10 is to inhibit excessive inflammation and maintain balance in the immune response. When M. tuberculosis infects natural production of IL-10 by these cells.

The main role of IL-10 in the regulation of immune responses is to inhibit production of pro-inflammatory cytokines, such as interleukin 1 (IL-1), interleukin 6 (IL-6), and nuclear factor kappa B (NF-κB). This reduces inflammation that can damage lung tissue but also inhibits the body's ability to quick to eliminate M. tuberculosis. In other words, IL-10 plays a dual role in the context of pulmonary tuberculosis. On the one hand, it helps control excessive inflammation that can cause serious lung damage, but on the other hand, too much IL-10 can inhibit an effective immune response against M. tuberculosis. Genetic polymorphisms in the IL-10 gene may influence how efficiently IL-10 regulates the immune response, and therefore, may influence disease progression at the individual level. In addition, IL-10 can also influence the response to pulmonary tuberculosis therapy. A number of studies have attempted to understand how IL-10 regulation may influence treatment sugass. In some cases, high levels of IL-10 production have been associated with a slower response to therapy. 13,14

IV. INTERACTION BETWEEN IL-10 GENETIC VARIATIONS AND PULMONARY TUBERCULOSIS

In efforts to understand the role of genetic variation in the interleukin 10 (IL-10) gene in the context of pulmonary tuberculosis, epidemiological research has become an important component. Epidemiological evidence includes observational studies involving different human populations to identify correlations between IL-10 genetic

polymorphisms and risk for pulmonary tuberculosis. Early epidemiological studies suggest that indivitials with certain genetic polymorphisms in IL-10 may have a higher risk of developing pulmanary tuberculosis. For example, the -1082 G/A polymorphism in the promoter region of the IL-10 gene has been associated with an increased risk of pulmonary tuberculosis in some populations. However, this epidemiological evidence is not always consistent across studies and pulations. Some studies show a strong association between IL-10 genetic polymorphisms and pulmonary tuberculosis, while other studies find a weaker association or no association at all. This variability in results may be due to factors such as differences in the geographic and ethnic distribution of genetic polymorphisms, as well as differences in research methodology.

A. GENETIC STUDY OF IL-10 POLYMORPHISMS AND PULMONARY TUBERCULOSIS

In addition to epidemiological evidence, further genetic studies have been conducted to better understand how genetic variations in IL-10 may influence the risk and progression of pulmonary tuberculosis. These studies often involve in-depth molecular analysis of IL-10 genetic polymorphisms, with a focus on their relationship to the body's response to M. tuberculosis infection. Genetic studies have identified various genetic polymorphisms in IL-10 that may influence IL-10 production and activity. Some studies suggest that individuals with certain alleles, such as the -1082 A allerg and the -819 T allele, may have higher IL-10 production. This may lead to changes in the body's immune response to M. tuberculosis. Genetic studies are at a 170 trying to reveal the relationship between IL-10 genetic polymorphisms and susceptibility to pulmonary tuberculosis. In several studies, individuals with polymorphisms that affect lower IL-10 production have demonstrated better ability to control infection and disease progression. Conversely, individuals with high IL-10 production may have a higher risk of developing severe disease.

However, the results from this genetic study also demonstrate the complexity in the interaction between IL-10 genetic variation and pulmonary tuberculosis. There is no universal consistency in findings, and many other factors, including environmental and genetic factors, also play a role in disease risk and progression. Therefore, genetic research continues to advance to provide a deeper understanding of the role of IL-10 in pulmonary tuberculosis and how individual genetic variability may influence the response to this infection. In the context of developing personalized therapies and better understanding individual vulnerabilities, this research has great potential to advance our understanding of pulmonary tuberculosis.

B. MOLECULAR MECHANISMS INVOLVED

In an effort of understand how genetic variations in the interleukin 10 (IL-10) gene interact with pulmonary tuberculosis at the molecular level, research has attempted to identify the mechanisms involved. This research involves an in-depth analysis of how IL-10 genetic polymorphisms influence molecular responses cellular and Mycobacterium tuberculosis tuberculosis 54 One of the mechanisms involved is the impact of genetic variations in IL-10 on the production of IL-10 itself. Certain poly 36 rphisms, such as the -1082 A allele, have been associated with has er IL-10 production. This may result in increased concentrations of IL-10 microenvironment surrounding tuberculous lesions in the lung. These high concentrations of IL-10 can inhibit an effective immune respose against M. tuberculosis by inhibiting the production of pro-inflammatory cytokines such interleukin 12 (IL-12) and interferon-gamma $(IFN-\gamma).^5$

Additionally, genetic polymorphisms in IL-10 can also influence IL-10 receptor pression on immune cells. Cells that have lower levels of 151-10 receptors may be more susceptible to the inhibitory effects of IL-10. This may impact the extent to which the immune response can be modulated by IL-10 in response to M. tuberculosis infection. Additionally, studies have revealed that genetic polymorphisms in IL-10 can [53] uence the regulation of the production of other pro-inflammatory and inflammatory cyto 34 es. These genetic variations may alter the balance between the pro-inflammatory immune response needed control infection and inflammatory response that protects lung tissue from excessive damage. In some cases, certain genetic polymorphisms can trigger a shift in this balance in one direction or another, with direct implications for the development and outcome of pulmonary tuberculosis.1,5

C. IMPLICATIONS IN DIAGNOSIS, PROGNOSIS, AND THERAPY

Understanding the interactions between ILgenetic var₅₁ions and pulmonary tuberculosis has important implications in the diagnosis, prognosis, and therapy of this disease. Individual genetic variability may influence how a person responds to M. tuberculosis infection and the extent to which the disease may progress 62 n the context of diagnosis, knowledge of IL-10 genetic polymorphisms may aid in the identification of individuals who may have a greater susceptibility to pulmonary tuberculosis. This could be useful in monitoring high-risk individuals or in the development of more sensitive diagnostic tests for early detection of the disease. Additionally, in disease prognosis, knowledge of IL-10 genetic variations can help predict the extent to which the disease will progress and potentially become severe. Individuals with genetic polymorphisms that result in high IL-10 production may be at greater risk for serious disease progression and associated complications. This may allow more accurate assessment of prognosis and more intensive treatment planning. ^{11,12,18}

In therapeutic management, knowledge of IL-10 genetic polymorphisms can help design more targeted and personalized therapeutic approaches. Several studies have attempted to understand how IL-10 regulation may influence response to pulmonary tuberculosis therapy. Individuals with high IL-10 production may require a different therapeutic approach than those with lower IL-10 production. As research continues to develop, a deeper understanding of the interactions between IL-10 genetic variations and pulmonary tuberculosis has great potential to improve the management of this disease. With more targeted approaches, more sensitive diagnostics, and therapies tailored to individual genetic characteristics, we can hope to reduce the burden of pulmonary tuberculosis and improve clinical outcomes for those infected.9

V. THE ROLE OF IL-10 IN THE IMMUNE SYSTEM

A. FUNCTION OF IL-10 IN THE REGULATION OF IMMUNE RESPONSES

Interleukin 10 (IL-10) is an important cyto to e in the human immune system that has a major role in regulating the immune response. The main function of IL-10 is to control excessive inflammation and 68 intain balance in the immune response. It plays an important role in keeping the immune system balanced, avoiding immune responses that damage healthy tissue, and preventing various autoimmune diseas and excessive inflammation. One way IL-10 regulates the immune response is by in 72 biting the activation of pro-inflammatory immune cells such as nacrophages and T cells. IL-10 can inhibit the production of pro-inflammatory cytokines such as interleukin 1 (IL-1), interleukin 6 (IL- 6), and tumor necrosis factor alpha (TNF- α). This reduces inflammation and calms excessive immune responses.

In addition, [37-10 can also inhibit the activation of T helper type 1 (Th1) cells, which produce cytokines such as interferongamma (IFN-y). Cellular Th1 is important in fighting intracellular infections such as pulmonary tuberculosis. By inhibiting Th1 activation, IL-10 can limit excessive immune responses to infections and prevent tissue damage caused by excessive inflammation. However, although IL-10 has an important role in regulating the immune response, too much IL-10 or inappropriate regulation of its production and activity can have a negative impact on the body's ability to fight infections. Therefore, a proper balance in IL-10 regulation is key to maintaining a healthy immune system.

B. IMPACT OF IL-10 GENETIC POLYMORPHISMS ON IL-10 FUNCTION

Genetic polymorphisms are variations in DNA sequence that can be found between individuals in different populations. In the context of IL-10, genetic polymorphisms may influence the function and regulation of IL-10 in the immune response. There are several genetic polymorphisms in the IL-10 gene that have been identified, the most important of which is the polymorphism at positions -1082, -819, and -592 in the promoter region of the IL-10 gene. This polymorphism may influence the legal of IL-10 production. For example, the -1082 G/A polymorphism has been associated with differences in IL-10 production. Indiviously with the -1082 A allele tend to have higher IL-10 production than individuals with the -1082 G allele. This may result in differences in the body's ability to regulate the immune response.19

Genetic polymorphisms in IL-10 may also influence responses to infection and

C. CONTRIBUTION TO THE PATHOGENESIS OF PULMONARY TUBERCULOSIS

The role of IL-10 in the pathogenesis of pulmonary tuberculosis has been the focus of intensive resear 65. M. tuberculosis infection can stimulate the production of IL-10 in response to the inflammation produced by the bact 66. This IL-10 production can then inhibit the immune response needed to eliminate the infection. M. tuberculosis is an intracellular bacterium that infects macrophage cells in the lungs. Macrophages are the first defense cells that try to destroy these bacteria. However, by stimulating IL-10 production, M. tuberculosis can inhibit macrophage activation and inhibit the ability of these cells to destroy bacteria.

In addition, IL-10 can also influence the activation of T cells, including T helper type I (Th1) cells which are important in fighting M. tuber ulosis infection. Th1 activation triggers the production of pro-inflammatory cytokines such as interferon-gamma (IFN-γ), which has a central role in activating macrophages to fight M. tuberculosis. By inhibiting Th1 activation, IL-10 can inhibit IFN-γ production and dampen the immune response necessary to control infection. In the context of IL-10 genetic polymorphisms, genetic variations in this gene may influence IL-10 production and regulation. As a result, individuals with certain polymorphisms may

have different immune responses to M. tuberculosis infection. Some studies suggest that individuals with higher IL-10 production, which may be associated with certain polymorphisms, may have a higher risk of developing pulmonary tuberculosis or may have the progression of more serious disease.

The role of Treg cells (T-cell regulatory) and interleukin 10 (IL-10) in the pathogenesis of erculosis (TB) is an important aspect related to the regulation of the immune response to infection with Mycobacterium tuberculosis (M. tube 19 losis), the bacteria that causes TB. Treg cells are a group of 9 cells that have a major role in regulating the immune response to prevent excessive immune restions and damage to body tissue. IL-10 is an anti-inflammatory cytokine produced by various types of cells, including Treg cells, and plays a role in reducing inflammation 14 Below is a further explanation regarding the role of Treg cells and IL-10 in TB pathogenesis:

VI. TREG CELLS IN TB PATHOGENESIS

Treg cell 57 re a type of T cell that have the ability to inhibit the activation and 27 liferation of T cells and other immune cells. The main function of Treg cells is to maintain immune tolerance to the body's own antigens, thereby preventing damaging autoimmune reactions.

In the context of TB, the role of Treg cells is to inhibit excessive immune responses against M. tuberculosis. Uncontrolled TB infection can cause inflammation that damages lung tissue, and Treg cells help prevent this by dampening excessive immune responses.

A. INTERACTION BETWEEN TREG CELLS AND IL-10 IN TB PATHOGENESIS

Treg cells have the ability to produce IL-10 as a mechanism to control inflammation. IL-

10 is an anti-inflammatory cytokine that plays a role in inhibiting the activation of pro-inflammatory immune cels, such as macrophages and T cells, as well as inhibiting the production of proinflammatory cytokines. In the context of TB, Treg cells that produce IL-10 may help control excessive imigane responses against M. tuberculosis. IL-10 produced by Treg cells can reduce the activation of Th1 cells, which play a role in fighting intracellular infections such as TB. Thus, Treg cells and IL-10 contribute to maintaining the balance between the immune response that fights infection and prevents excessive tissue damage.

B. IMPACT ON ACTIVE AND LATENT TUBERCULOSIS

The role of Treg cells and IL-10 in TB pathogenesis may have consequences for the type of TB disease that develops. In active tuberculosis, high Treg cell activity and excessive IL-10 production may help M. tuberculosis evade detection and dampen an effective immune response, allowing the infection to progress to active dease. On the other hand, Treg cells and IL-10 may also play a role in tuberculoma formation and the development of latent tuberculosis. They help prevent an excessive immune response to the bacteria, allowing M. tuberculosis to persist in 13 e body without causing active disease. Understanding the role of Treg cells and L-10 in TB pathogenesis is important in the development of more effective therapeutic strategies and vaccines. Several studies have attempted to ident ways to regulate Treg cell activity or alter the balance between pro-inflammatory and anti-inflammatory immune responses in an effort to control TB infection. This is an important area of research in efforts to address the global TB burden and better understand the complex interactions between the human body and M. tuberculosis.

Interaction between Treg Cel 11 IL-10, and TB: Treg cells, 45 nich have the ability to produce IL-10, play a role in controlling excessive immune responses against M. tuberculosis. However, too many Treg cells and IL-10 can affect the body's ability to eliminate TB infection. Knowledge of IL-10 genetic variations may help in designing more personalized therapeutic approaches. This allows doctors to identify individuals who may have a different immune response to TB infection and prescribe more aprize therapy. A deeper understanding of the role of IL-10 and Treg cells in TB pathogenesis opens the door to the developmen 77 f new therapies focused on regulating the balance of the immune response. This includes the development of drugs that can influence IL-10 activity or alter Treg cell activity. IL-10 genetic variations can also be used in disease monitoring and prognosis assessment. with Individuals certain genetic polymorphisms may be at higher risk for the development of serious disease, and closer monitoring or different therapeutic approaches may be implemented.

VII. IMMUNOTHERAPY AND TARGETED THERAPY IN PULMONARY TUBERCULOSIS

A. POTENTIAL FOR UTILIZING IL-10 GENETIC KNOWLEDGE IN THERAPY

32 Knowledge of the role of IL-10 and genetic variations of IL-10 may provide the basis for more targeted therapeutic approaches. Genetic polymorphisms in IL-10 may influence an individual's response to M. tuberculosis infection and therapy. For example, individuals with polymorphisms that result in higher IL-10 production may have a more inhibited immune response to infection. In such cases, a more aggressive therapeutic approach may be necessary, including administration of stronger anti-TB drugs or adjuvant therapy aimed at overcoming immune barriers. On the other hand. individuals with lower IL-10

production may have a stronger immune response to infection. They may benefit from milder therapies or strategies that focus on enhancing the body's natural immune response.

Additionally, understanding the molecular mechanisms involved in IL-10 regulation may open the door to the development of drugs that can influence IL-10 activ 6. These drugs can be used to change the balance between pro-inflammatory and anti-inflammatory immune responses in the body, with the aim of increasing the body's ability to overcome TB infection

B. CHALLENGES AND OPPORTUNITIES

Although immune-based and targeted therapies offer great opportunities in treating TB, there are a number of challenges that need to be overcome. One of them is the complexity of the interactions between individual genetics, immune response, and TB infection. In some cases, a stronger immune response does not necessarily mean better protectan, and vice versa. Another challenge is the development of effective drugs and vaccines. The process of developing new drugs and vaccines requires time, rigorous clinical trials, and large financial investments. However, with evergrowing knowledge in the fields of genetics and immunology, opportunities to develop better therapies are increasing.

In addition, target-based therapy must also consider the issue of drug resistance. As a slow-growing bacterium, M. tuberculosis can develop resistance to drugs quickly. Therefore, there needs to be continued efforts to develop more robust and diverse therapies. Overall, the development of immunotherapy and target-based therapy in the treatment of pulmonary tuberculosis is an important step in the global effort to overcome this disease. With a deeper understanding of individual genetics and the immune response to M. tuberculosis, we can

hope to develop more personalized and effective therapies, and minimize the disease burden of pulmonary tuberculosis worldwide.

VIII. CONCLUSION



The role of interleukin 10 (IL-10) and its genetic variations in the pathogenesis and management of pulmonary tuberculosis (TB) has been discussed. The following is a summary of the main findings that can be conclided from this study: Role of IL-10 in TB: IL-10 is an important cytokine in the immune system that has a primary function regulating the 69 immune response. However, excessive production of IL-10 can inhibit the immune response required to overcome M. tuberculosis infection in pulmonary tubercul sis. IL-10 Genetic Variations: Genetic polymorphisms in the IL-10 gene can influence the production and regulation of IL-10 11 the body. Some polymorphisms have been associated with higher or lower risk of TB, and this raises the question of how these genetic variations influence the response to infection.

REFERENCES

- Anggraini D, Nasrul E, Susanti R, Suharti N. Polymorphysm of tumor necrosis factor-A interleukin-10 gene with pulmonary tuberculosis susceptibility. J Popul Ther Clin Pharmacol. 2023;30(2):50-8.
- [2] Commentary S, Review S, Review S, Review S, Review S, Review S, Redford PS, et al. The role of IL-10 in immune regulation during M . tuberculosis infection. Soc Mucosal Immunol [Internet]. 2011;4(3):261–70. Available from: http://dx.doi.org/10.1038/mi.2011.7
- [3] DiNardo AR, Gandhi T, Heyckendorf J, Grimm SL, Rajapakshe K, Nishiguchi T, et al. Gene expression signatures identify biologically and clinically distinct tuberculosis endotypes. Eur Respir J [Internet]. 2022;60(3). Available from: http://dx.doi.org/10.1183/13993003.02263-2021
- [4] Abel L, El-Baghdadi J, Bousfiha AA, Casanova JL, Schurr E. Human genetics of tuberculosis: A long and winding road. Philos Trans R Soc B Biol Sci. 2014;369(1645).
- [5] Ahmed A. Emerging patterns of regulatory T cell function in tuberculosis. 2020;273–87.

- [6] Bellerose MM. Genetic Identification of Novel Mycobacterium tuberculosis Susceptibility and Survival Mechanisms During Antibiotic Treatment Let us know how access to this document benefits you . 2020;
- [7] Sawitri NE, Prasetyo AA, Moewardi R. Polimorfisme Gen IL-10 -1082 G / A Sebagai Faktor Kerentanan Pejamu pada Pasien Tuberkulosis Multidrug Resistant Host Susceptibility Factor In Patients With Multidrug Resistant Tuberculosis. 2014;36(1):1–10.
- [8] Seviyelerinin S, Polimorfizminin I--AC, İlaç T. IL-10-592 A / C Gene Polymorphism and Cytokine Levels are Associated with Susceptibility to Drug Resistance in Tuberculosis. 2020;8(3):103-12.
- [9] Grigorov B, Trenova AG, Miteva LD, Stanilova SA. Interleukin-10 (IL-10) promoter polymorphism at position – 1082 in Bulgarian patients with multiple sclerosis. 2019;10(April).
- [10] Silva CA, Fernandes DCRO, Braga ACO, Cavalcante GC, Sortica VA, Hutz MH, et al. Investigation of genetic susceptibility to Mycobacterium tuberculosis (VDR and IL10 genes) in a population with a high level of substructure in the Brazilian Amazon region. Int J Infect Dis [Internet]. 2020;98:447–53. Available from: https://doi.org/10.1016/j.ijid.2020.06.090
- [11] McHenry ML, Bartlett J, Igo RP, Wampande EM, Benchek P, Mayanja-Kizza H, et al. Interaction between host genes and Mycobacterium tuberculosis lineage can affect tuberculosis severity: Evidence for coevolution? PLoS Genet [Internet]. 2020;16(4):1–18. Available from: http://dx.doi.org/10.1371/journal.pgen.1008728
- [12] Mchenry ML, Williams SM, Stein CM, Sciences QH, Sciences G, Western C. perspectives and approaches, 2021;2020:1–25.
- [13] Walker TM, Ip CLC, Harrell RH, Evans JT, Kapatai G, Dedicoat MJ, et al. Whole-genome sequencing to delineate Mycobacterium tuberculosis outbreaks: A retrospective observational study. Lancet Infect Dis [Internet]. 2013;13(2):137–46. Available from: http://dx.doi.org/10.1016/S1473-3099(12)70277-3
- [14] Wampande EM, Naniima P, Mupere E, Kateete DP, Malone LSL, Stein CM, et al. Genetic variability and consequence of Mycobacterium tuberculosis lineage 3 in Kampala-Uganda. PLoS One. 2019;14(9):1–14.
- [15] Ndhlovu V, Kiran A, Sloan D, Mandala W, Kontogianni K, Kamdolozi M, et al. Genetic diversity of Mycobacterium tuberculosis clinical isolates in Blantyre, Malawi. Heliyon [Internet]. 2019;5(10):e02638. Available from: https://doi.org/10.1016/j.heliyon.2019.e02638

- [16] Peresi E, Ragozo L, Oliveira C, Laurentino W, Alessandra É, Nunes P, et al. Cytokine Polymorphisms, Their Influence and Levels in Brazilian Patients with Pulmonary Tuberculosis during Antituberculosis Treatment, 2013;2013.
- [17] Raja A. Immunology of tuberculosis. Indian J Med Res. 2004;120(4):213–32.
- [18] Ismail N, Rivière E, Limberis J, Huo S, Metcalfe JZ, Warren RM, et al. Genetic variants and their association with phenotypic resistance to bedaquiline in Mycobacterium tuberculosis: a systematic review and individual isolate data analysis. The Lancet Microbe. 2021;2(11):e604–16.
- [19] Anggraini, D., & Oktora, M. Z. (2021). Hematology Profile of Tuberculosis Lymphadenitis Patients at Siti Rahmah Hospital, Padang, Indonesia. Indonesian Journal of Clinical Pathology and Medical Laboratory (IJCPML), 27(3).

The Role of Interleukin 10 Genetic Variations in Pulmonary Tuberculosis: Perspectives of Genetics, Pathogenesis and Immunology

ORIGI	NALITY REPORT				
	23% SIMILARITY INDEX				
PRIMARY SOURCES					
1	www.science.gov Internet	182 words — 3%			
2	encyclopedia.pub Internet	54 words — 1 %			
3	impactfactor.org Internet	41 words — 1 %			
4	"Tuberculosis", Springer Science and Business Med LLC, 2023 Crossref	ia 40 words — 1 %			
5	worldwidescience.org Internet	37 words — 1 %			
6	iv.iiarjournals.org Internet	32 words — 1 %			
7	link.springer.com Internet	32 words — 1 %			
8	Petr Nemec. "Association of Polymorphisms in Interleukin-10 Gene Promoter with Autoantibody Production in Patients with Rheumatoid Arthritis", A New York Academy of Sciences, 09/2009	31 words -1%			

- Shih, C.M.. "The involvement of genetic polymorphism of IL-10 promoter in non-small cell 29 words <1% lung cancer", Lung Cancer, 200512
- Anne O'Garra. "IL-10-producing and naturally occurring CD4+ Tregs: limiting collateral damage", Journal of Clinical Investigation, 11/15/2004
- Lela Kania Rahsa Puji, Tri Okta Ratnaningtyas, Frida Kasumawati, Nurwulan Adi Ismaya, Nur Hasanah. "The Correlation of Individual and External Factors to Work Fatigue in Employee at PT. Hutama Karya Building Division Project, Integrated Building Soekarno Hatta Airport Train Station", Health and Medical Journal, 2022
- www.openarchives.org $_{\text{Internet}}$ 24 words -<1%
- "INTERLEUKIN-10 AND THE INTERLEUKIN-10 RECEPTOR", Annual Review of Immunology, 04/2001

 Crossref
- David Sacks. "Re-examination of the immunosuppressive mechanisms mediating noncure of Leishmania infection in mice", Immunological Reviews, 10/2004Crossref
- Hai-Ling Qiao, Qiang Wen, Na Gao, Xin Tian, Lin-Jing Jia. "Association of IL-10 level and IL-10 promoter SNPs with specific antibodies in penicillin-allergic patients", European Journal of Clinical Pharmacology, 2007

- Luigi Boiardi, Bruno Casali, Enrico Farnetti, Nicolò $_{22\,words}$ < 1 9 0 Pipitone et al. "Interleukin-10 promoter polymorphisms in giant cell arteritis", Arthritis & Rheumatism, 2006
- Opdal, S.H.. "Il-10 gene polymorphisms are associated with infectious cause of sudden infant death", Human Immunology, 200312
- biosignaling.biomedcentral.com

 Internet

 19 words -<1%
- 19 www.frontiersin.org
 Internet 19 words < 1 %
- Catherine J. Edwards-Smith, Julie R. Jonsson, David M. Purdie, Amolak Bansal, Claudia Shorthouse, Elizabeth E. Powell. "Interleukin-10 promoter polymorphism predicts initial response of chronic hepatitis C to interferon alfa", Hepatology, 1999 Crossref
- Dewi Safnita, Hera Novianti, Yessy Setiawati, Pamelia Mayorita. "Adenokarsinoma Paru yang Didiagnosis dari Histopatologi Metastasis di Otak dan Perikardium", Health and Medical Journal, 2022 Crossref
- scholarworks.gsu.edu
 _{Internet}

 18 words < 1 %
- Mangia, A.. "IL-10 haplotypes as possible predictors of spontaneous clearance of HCV infection", Cytokine, 20040207

- Zhong, Qianfu, Cheng Ding, Meilin Wang, Ying Sun, and Yan Xu. "Interleukin-10 gene polymorphisms and chronic/aggressive periodontitis susceptibility: A meta-analysis based on 14 case-control studies", Cytokine, 2012.

 Crossref
- ELISABETH C. McGOWAN. "Placental IL-10 Dysregulation and Association With Bronchopulmonary Dysplasia Risk", Pediatric Research, 10/2009 Crossref
- Ikram Sghaier, Leila Mouelhi, Noor A. Rabia, Ezzedine Ghazoueni, Wassim Y. Almawi, Besma Yacoubi Loueslati. "IL-10 and IL-28B gene variants as predictors of sustained response to peginterferon and ribavirin therapy in chronic HCV infection", Cytokine, 2022 $_{\text{Crossref}}$
- adoc.pub 16 words < 1 %
- bjo.bmj.com
 Internet

 16 words -<1%
- discovery.dundee.ac.uk $_{\text{Internet}}$ 16 words -<1%
- www.mdpi.com
 Internet

 16 words < 1 %
- Alain P. Vicari. "Interleukin-10 in viral diseases and cancer: exiting the labyrinth?", Immunological Reviews, 12/2004

32	X. Jiang. "Mechanism of NKT Cell-Mediated Transplant Tolerance", American Journal of Transplantation, 6/2007 Crossref	15 words — < 1%
33	daneshyari.com Internet	14 words — < 1%
34	A. Tagore. "Interleukin-10 (IL-10) genotypes in inflammatory bowel disease", Tissue Antigens, 10/1999 Crossref	13 words — < 1%
35	Debie Anggraini. "Laboratory Examination in Hepatocelullar Carcinoma", Health & Medical Journal, 2019 Crossref	13 words — < 1%
36	L. Beretta. "Proximal interleukin-10 gene polymorphisms in Italian patients with systemic sclerosis", Tissue Antigens, 4/2007 Crossref	13 words — < 1 %
37	www.medsci.org Internet	13 words — < 1%
38	M Schrappe. "Association of initial response to prednisone treatment in childhood acute lymphoblastic leukaemia and polymorphisms wit tumour necrosis factor and the interleukin-10 ge Leukemia, 10/18/2002 Crossref	
39	api-ir.unilag.edu.ng Internet	12 words — < 1%

10 words -<1%P. Nahid, L. G. Jarlsberg, M. Kato-Maeda, M. R. 41 Segal et al. "Interplay of strain and race/ethnicity in the innate immune response to M. tuberculosis", PLOS ONE, 2018

Crossref

www.minervamedica.it

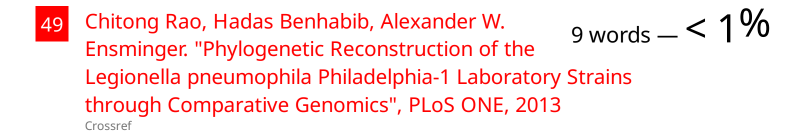
46

Internet

10 words -<1%Seiji Miyazoe. "Influence of interleukin-10 gene promoter polymorphisms on disease progression in patients chronically infected with hepatitis B virus", The American Journal of Gastroenterology, 8/2002 Crossref

10 words -<1%academic.oup.com Internet 10 words -<1%drrajivdesaimd.com Internet $_{10 \text{ words}} = < 1\%$ www.jimmunol.org Internet

- $_{10 \text{ words}} = < 1\%$ Internet 10 words -<1%www.researchgate.net
- Asner, S. A., S.A. Morré, Pierre-Yves Bochud, and G. $_{9 \text{ words}}$ < 1%48 Greub. "Host factors and genetic susceptibility for infections due to intracellular bacteria and fastidious organisms", Clinical Microbiology and Infection, 2014. Crossref



- F Farzaneh. "The IL-10 –1082G polymorphism is associated with clearance of HPV infection", BJOG

 An International Journal of Obstetrics and Gynaecology, 8/2006

 Crossref
- J. A. Whitsett. "Genetic Basis of Familial Interstitial Lung Disease: Misfolding or Function of Surfactant 9 words < 1% Protein C?", American Journal of Respiratory and Critical Care Medicine, 05/01/2002
- Omaima M. Abbas. "Interleukin-10 promoter polymorphisms in hepatitis C patients with and without <i>Schistosoma mansoni</i> co-infection", Liver International, 10/2009 Crossref
- W. Alpízar-Alpízar, G. I. Pérez-Pérez, C. Une, P. Cuenca, R. Sierra. "Association of interleukin-1B and interleukin-1RN polymorphisms with gastric cancer in a high-risk population of Costa Rica", Clinical and Experimental Medicine, 2005 Crossref
- djm.uodiyala.edu.iq
 Internet

 9 words < 1%
- e-sciencecentral.org
 Internet

 9 words -<1%
- garuda.kemdikbud.go.id

Crossref

9 words — <	1	%
-----------------------	---	---

molecular-cancer.biomedcentral.com

- 9 words -<1%
- "C5 Cardiovascular morbidity and mortality (T485 1 % words 1 % T497)", Nephrology Dialysis Transplantation, 2003.
- Ito, T.. "Two Functional Subsets of FOXP3 $^+$ Regulatory T Cells in Human Thymus and Periphery", Immunity, 20080613
- Kallas, Eveli, Kristi Huik, Merit Pauskar, Ene-Ly Jõgeda, Tõnis Karki, Don Des Jarlais, Anneli Uusküla, Radko Avi, and Irja Lutsar. "Influence of interleukin 10 polymorphisms -592 and -1082 to the HIV, HBV and HCV serostatus among intravenous drug users", Infection Genetics and Evolution, 2015.
- Paulo Vieira. "Regulatory T cells and mechanisms of immune system control", Nature Medicine, 08/2004Crossref
- S.-H. Kim. "Combined effect of IL-10 and TGF- β 1 promoter polymorphisms as a risk factor for aspirin-intolerant asthma and rhinosinusitis", Allergy, 08/2009 Crossref
- Spanish Group for the Study of Drug-Induced Liver $_{8 \text{ words}} < 1\%$ Disease (Grupo de Estudio para las Hepatopatias Asociadas a Medicamentos (GEHAM)). "Analysis of IL-10, IL-4

and TNF-@a polymorphisms in drug-induced liver injury (DILI) and its outcome", Journal of Hepatology, 200807

64	Tushar Patil, Ravindra Kumar Garg, Amita Jain, Madhu Mati Goel et al. "Serum and CSF cytokines and matrix metalloproteinases in spinal tuberculos Inflammation Research, 2014 Crossref	8 words — <	1%
65	d.docksci.com Internet	8 words — <	1%
66	docksci.com Internet	8 words — <	1%
67	jurnal.umj.ac.id Internet	8 words — <	1%
68	mol-biol4masters.masters.grkraj.org	8 words — <	1%
69	www.journalofbabylon.com Internet	8 words — <	1%
70	www.jptcp.com Internet	8 words — <	1%
71	www.wjgnet.com Internet	8 words — <	1%
72	www.zhb.uni-luebeck.de Internet	8 words — <	1%
73	Chi Chiu Mok. "Interleukin-10 promoter polymorphisms in Southern Chinese patients with	7 words — <	1%

systemic lupus erythematosus", Arthritis & Rheumatism, 06/1998

- Craig Kinnear, Eileen G. Hoal, Haiko Schurz, Paul D. 7 words <1% van Helden, Marlo Möller. "The role of human host genetics in tuberculosis resistance", Expert Review of Respiratory Medicine, 2017
- E G de la Concha. "Interleukin-10 polymorphisms in Spanish type 1 diabetes patients", Genes and Immunity, 06/2004Crossref
- Jasenka Trifunović, Larisa Miller, Željko Debeljak, Vesna Horvat. "Pathologic patterns of interleukin 10 expression A review", Biochemia Medica, 2015
- Li, M.O.. "Contextual Regulation of Inflammation: A words <1% Duet by Transforming Growth Factor-@b and Interleukin-10", Immunity, 20080411
- Nina Koldzic-Zivanovic, Huolin Tu, Terry L. Juelich, Peter L. Rady et al. "Regulation of adrenal glucocorticoid synthesis by interleukin-10: A preponderance of IL-10 receptor in the adrenal zona fasciculata", Brain, Behavior, and Immunity, 2006

 Crossref
- Steinke, J.W.. "Differential interleukin-10 production $_7$ words <1% stratified by -571 promoter polymorphism in purified human immune cells", Cellular Immunology, 200710

- "Posters", Journal of the European Academy of Dermatology and Venereology, 11/2004
- $_{6 \text{ words}}$ < 1%

- D Kube, T-D Hua, M Klöss, B Kulle, J Brockmöller, L Wojnowski, M Löffler, M Pfreundschuh, L Trümper. 6 words < 1% "The interleukin-10 gene promoter polymorphism –1087AG does not correlate with clinical outcome in non-Hodgkin's lymphoma", Genes & Immunity, 2007
- Eliana Peresi, Larissa Ragozo Cardoso Oliveira,
 Weber Laurentino da Silva, Érika Alessandra
 Pellison Nunes da Costa et al. "Cytokine Polymorphisms, Their
 Influence and Levels in Brazilian Patients with Pulmonary
 Tuberculosis during Antituberculosis Treatment", 'Hindawi
 Limited', 2013
 Internet
- Enam Reyaz, Niti Puri, Angamuthu Selvapandiyan. $_{6 \text{ words}} < 1\%$ " Global Remodeling of Host Proteome in Response to Infection ", ACS Infectious Diseases, 2023
- Marcos V. da Silva, Vladimir J. Massaro Junior,
 Juliana R. Machado, Djalma A. A. Silva et al. "

 Expression Pattern of Transcription Factors and Intracellular
 Cytokines Reveals That Clinically Cured Tuberculosis Is
 Accompanied by an Increase in -Specific Th1, Th2, and Th17
 Cells ", BioMed Research International, 2015
 Crossref
- Merja Helminen. "IL-10 gene polymorphism at $_{-1082}$ A/G is associated with severe rhinovirus bronchiolitis in infants", Pediatric Pulmonology, 04/2008

- Petr Nemec. "Association of the -1082 G/A promoter polymorphism of interleukin-10 gene with the autoantibodies production in patients with rheumatoid arthritis", Clinical Rheumatology, 03/24/2009
- Ying-Ju Lin, Yu-Ching Lan, Lei Wan, Ting-Hsu Lin, Da-Yuan Chen, Chang-Hai Tsai, Chiu-Shong Liu, Kai-Chung Hsueh, Fuu-Jen Tsai. "Serological surveillance and IL-10 genetic variants on anti-HBs titers: Hepatitis B vaccination 20years after neonatal immunization in Taiwan", Clinica Chimica Acta, 2011 Crossref

EXCLUDE QUOTES ON

EXCLUDE BIBLIOGRAPHY ON

EXCLUDE SOURCES

OFF

EXCLUDE MATCHES

OFF