Tumour Necrosis Factor Gene Polymorphism and Disease Prevalence

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Abstract

Tumour necrosis factor (TNF), an important proinflammatory cytokine, plays a role in the regulation of cell differentiation, proliferation and death, as well as in inflammation, innate and adaptive immune responses, and also implicated in a wide variety of human diseases. The presence of DNA sequence variations in regulatory region might interfere with transcription of TNF gene, influencing the circulating level of TNF and thus increases the susceptibility to human diseases (infectious, cancer, autoimmune, neurodegenerative and other diseases). In this review, we have comprehensively analysed various published case–control studies of different types of human diseases, in which TNF gene polymorphism played a role, and computationally predicted several single nucleotide polymorphisms (SNPs) lie in transcription factor–binding sites (TFBS) of transcription factors (TFs). It has been observed that TNF enhancer polymorphism is implicated in several diseases, and TNF rs1800629 and rs361525 SNPs are the most important in human disease susceptibility as these might influence the transcription of TNF gene. Thirty-two SNPs lies in TFBS of 20 TFs have been detected in the TNF upstream region. It has been found that TNF enhancer polymorphism influences the serum level of TNF in different human diseases and thus affects the susceptibility to diseases. The presence of DNA sequence variation in TNF gene causes the modification of transcriptional regulation and thus responsible for association of susceptibility/resistance with human diseases.

Introduction

Tumour necrosis factor (TNF) cytokine, produced as the part of host defence against infection. This cytokine is involved in multiple inflammatory and immune responses and plays role in the pathogenesis of many autoimmune and infectious diseases. TNF gene is located on chromosome 6 in the class III region of the major histocompatibility complex (MHC) and is flanked by the lymphotoxin ‘α’ and ‘β’ genes (Fig. 1). A close linkage among HLA class I (HLA-B), class II (HLA-DR) and TNF genes has been reported [1]. TNF gene is tightly regulated at the level of transcription [2, 3].

DNA sequence variation in promoter regions of genes encoding cytokines influences susceptibility to infection and has been associated with a large number of complex human diseases. Reports indicated that polymorphism in the 5’ regulatory region of the gene has been correlated with many infectious and inflammatory diseases [4, 5].

The association of TNF rs1799964 and rs1800630 polymorphisms with advanced-stage endometriosis in the Korean population have been reported. The TNF rs1800750 polymorphism affects the binding of TF OCT-1 and alters the DNA–protein interaction. The in vitro study of TNF promoter polymorphism function was stimulated by several case–control studies of the polymorphism in relation to human disease [6]. The polymorphism rs361525 has been associated with insulin resistance syndrome and obesity [7], and rs1800629 has been associated with various inflammatory and autoimmune diseases [8–13]. Associations between polymorphism (rs1799964, rs1799724, rs1800630) and immune-mediated diseases such as rheumatoid arthritis and Crohn’s disease (CD) have been reported [14, 15]. Limited reports are available showing that variants (rs1800629 and rs361525) are involved in the regulation of cytokine production [16]. The rs1799964 polymorphism has been associated with extra intestinal
manifestations of CD including uveitis, erythema nodosum and large joint arthropathy [17] and Crohn’s disease itself [16]. It is clear that TNF enhancer polymorphism is implicated in several case–control studies. In the present review, the literature regarding the role of TNF-α polymorphism has been studied with respect to different human diseases and different populations. Several single nucleotide polymorphisms (SNPs) in TFBS of different TFs have been predicted computationally. The purpose of this review is to provide an overview of what is currently known about the role of gene level polymorphism of TNF and susceptibility/resistance to human diseases and to highlight directions that are likely to see major advances.

Infectious diseases

Bacterial diseases

Pulmonary tuberculosis. Mycobacterial 

tuberculosis is the leading cause of mortality in India as well as in the world. Approximately one-third of the world’s population is suffering from Mycobacterial diseases [18, 19]. Pulmonary tuberculosis, caused by M. tuberculosis, is a granulomatous disease of the lungs. The host genetic factor plays a significant role in determining susceptibility to developing the active form of the disease [20, 21]. A number of genes have been identified, which are important in tuberculosis [22–24]. Elevated serum tumour necrosis factor-α (sTNF-α) levels have been reported in patients with advanced tuberculosis in comparison with those with mild tuberculosis and healthy controls. Several polymorphisms within the promoter region of TNF-α and the intron 1 of LT-α have been associated with altered circulating levels of TNF-α [25, 26]. Some of these polymorphisms have been determine susceptibility or resistance to tuberculosis in several ethnic groups [27–33]. Sharma et al. [34] carried out a case–control study, including patients with pulmonary tuberculosis and controls in North India. In this study, five promoter SNPs in TNF-α gene and one SNP rs909253 in LTα gene were detected in patients with tuberculosis and controls samples collected from North India (Fig. 2). No significant differences in allele frequencies between the patients with tuberculosis and controls were reported. Serum TNF-α levels showed a significant difference between patients with tuberculosis and controls, and none of the polymorphism affects the serum TNF levels. Ates et al. [27] reported that IL10 and TNF promoter polymorphisms have been associated with altered levels of circulating IL10 and TNF-α. A significant association was reported between TB and rs1800896 G-allele. IL10 GCC and ACC haplotypes distribution showed a significant difference between patients with TB and controls. No statistically significant association was detected between rs1800871, rs1800629, rs1800750, rs361525 polymorphisms, functional TNF-α/IL-10 genotypes and TB.

Leprosy. Leprosy is a mycobacterial disease, caused by Mycobacterium leprae that initially affects the peripheral nervous system and patients displaying contrasting clinical, immunological and pathological manifestations. Many factors and metabolic pathways including TLR/LIR-7, VDR, TNF-α and TGF-β have been reported to play role in disease. Goulart and Goulart [35] reviewed the complex molecular interactions in affected individuals influenced by the pathogenetic background. A significant association between the TNF rs1800629 A-allele and multibacillary leprosy has been reported from India [36]. In Brazil, this allele was associated with resistance against multibacillary leprosy [37, 38]. A significant association of TNF rs1800629 was found in borderline tuberculoid leprosy patients with the magnitude of in vivo delayed type hypersensitivity skin test reactivity to cutaneously injected M. leprae antigens. It has been reported that signalling deficient mutations in

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certain Toll-like receptors (TLR2; act upstream of TNF) can be strongly correlated with lepromatous leprosy. TNF rs1800629 regulatory polymorphism plays an important role in patients with leprosy in a Brazilian population [39], and in patients with leprosy, higher frequency of TNF rs1800629, GG genotype, and a decreased frequency of GA/AA genotypes were reported as compared to the control group. The GG genotype was particularly higher in patients with tuberculoid (TT) and borderline (BB) leprosy. A lower frequency of GCC/GCC haplotype of IL-10 in patients with lepromatous leprosy (LL) than in controls was also reported. TNF-alpha polymorphism rs361525 and rs1800629, and its association with the outcome of different clinical forms of leprosy have been reported by Vanderborght et al. [39]. TNF polymorphism rs361525 and rs1800629 have shown differences in the frequency of the haplotypes among the ethnic groups, but no statistical differences were observed in haplotype frequencies between patients with multibacillary (MB) and paucibacillary (PB). A lower bacteriological index (BI) among the TNF polymorphism rs1800629 carriers was reported, while higher BI in rs361525 carriers [40].

Recurrent acute otitis media. Acute otitis media (AOM) is caused by bacterial infection in children. Genetic variations in immune response genes are reported to influence susceptibility to infectious diseases [41], and increased expression of TNF-α, IL-1β, IL-6 and IL-10 was observed during experimental otitis media in animals. Polymorphism in immune response genes such as IL10, IL6 and IL4 has been associated with altered cytokine expression levels [42]. The polymorphism rs361525 was associated with the otitis-prone condition and lower specific anti-pneumococcal antibody levels after vaccinations in comparison with carriers of the A-allele. TNF polymorphism rs1800630 A-allele was associated with lower specific anti-pneumococcal IgG levels compared with children carrying C/C genotype of rs1800630.

Typhoid fever. Typhoid fever is caused by Salmonella enterica infection with serotype Typhi and 22 million cases of typhoid fever occur worldwide per year, resulting in 200,000 deaths. Indonesian study suggested a protective role of DRB1*12021 for complicated typhoid fever. Keuter et al. [43] found a lower level of TNF-α in the patients with acute phase of typhoid fever than in convalescence. The seven polymorphisms have been found within the genesBAT1 (a member of the DEAD-box protein family encoding an ATP-dependent RNA helicase and a negative regulator of inflammation), LTA and TNF. All three genes, or haplotypes spanning these genes, have been associated with a variety of infectious and inflammatory diseases. Dunstan et al. [44] genotyped eighty SNPs in a region of 150 kb in Vietnamese individuals. Thirty-three SNPs with a minor allele frequency of greater than 4.3% were used to construct haplotypes. Fifteen SNPs which tagged the 42 constructed haplotypes were selected. The haplotype-tagging SNPs (T1–T15) were genotyped, and allelic frequencies of seven SNPs (T1, T2, T3, T5, T6, T7 and T8) have shown a significant difference between typhoid cases and controls. Haplotype-based analysis of the tag SNPs provided positive evidence of association with typhoid. The analysis detected a low-risk cluster of haplotypes that each carries the minor allele of T1 or T7, but not both, and otherwise carries the combination of alleles *12122*1111 at T1–T11. Individuals who carry the typhoid fever–protective haplotype *12122*1111 also produce a relatively low TNF-α response to LPS.

Severe sepsis in trauma patients. In the non-coronary intensive care unit, sepsis is the prevalent cause of death. A restriction fragment length polymorphism (RFLP) present in TNF gene is correlated with increased level of TNF-α in plasma and a high mortality rate in patients with severe sepsis. This non-synonymous polymorphism in the first intron of the TNF-β gene (1064–1069 position) is responsible for an amino acid change at position 26 (asparagine for the TNFB1 sequence and threonine for the TNFB2 sequence) [45]. Previously, the mortality rate in severe sepsis was found to be significantly increased in patients homozygous for the allele TNFB2 of the Nco1 polymorphism compared with heterozygous patients [46]. A statistically highly significant association was obtained between the genotype of the biallelic Nco1 polymorphism of the TNF-β gene and the development of severe sepsis after severe blunt trauma. Subjects homozygous for the allele TNFB2 have a significantly increased risk of the development of severe post-traumatic sepsis.

Sepsis-induced organ failure causes death because of activation of a mediator cascade initiated by microbial components [47]. Mira et al. [48] reported the association of TNF2 (rs1800629 SNP with A-allele) with Septic Shock Susceptibility and Mortality. This polymorphism has been correlated with enhanced spontaneous and stimulated TNF-alpha production both in vitro and in vivo and has been associated with morbidity and mortality of severe forms of cerebral malaria [49], fulminans purpura [9], and mucocutaneous leishmaniasis (MCL) [10]. Variation in TNF2 allele frequencies between the controls and patients with septic shock was reported. The patients with septic shock had significantly greater TNF2 allele frequency in comparison with those who had died.
NcoI polymorphism. NcoI is a restriction enzyme used in the typing of polymorphism. The presence of A-allele eliminates the restriction site for the enzyme NcoI, while G-allele creates restriction site for NcoI restriction enzyme.

Mediterranean spotted fever. Cytokines plays important role in the protective immune response against Rickettsia conorii. A significantly elevated levels of IFN-γ, TNF-α, IL-10 and IL-6 in serum was observed in patients with acute-phase Mediterranean spotted fever (MSF) compared with the levels found during the convalescent phase of the disease or in healthy controls. Forte et al. [50] carried out genotyping of the TNF-alpha (rs1800629), interleukin-10 (rs1800896, rs1800871 and rs1800872) and IFN-gamma (rs2430561) in a group of Sicilian patients affected by MSF. No significant differences in TNF-α polymorphism in Sicilian patients compared to CC genotype.

Viral diseases

Clearance of hepatitis B virus infection. Hepatitis B virus (HBV) infection is a global public health problem, and more than 350 million people are infected with HBV worldwide. Tumour necrosis factor-alpha (TNF-α) plays an important role in host immune response to HBV. Kim et al. [51] carried out a case–control study of hepatitis B-infected patients and controls and genotyped seven TNF-α polymorphism in Korean. The results of the study showed that the presence of the rs1800629 A-allele or the absence of the rs1800630 A-variant was strongly associated with the resolution of HBV infection. The two TNF-α haplotypes were significantly associated with HBV clearance, showing protective antibody production and persistent HBV infection. Thus, those variations that affect the level of gene product might influence the outcome of disease. SNP rs1800629 A is common in Iranian population, but has no association with development of chronic HBV infection [52].

SARS-CoV infection. Severe acute respiratory syndrome (SARS) disease is caused by a novel coronavirus-SARS-CoV.

Host genetic factors may play a role in the occurrence and progress of SARS-Cov infection. Wang et al. [53] conducted case–control study including SARS-infected patients, health care workers and controls. They found no differences in TNF-α genotype distribution at the rs1799964, rs1800630, rs1800629 and rs361525 among the three populations. The CT and CC genotypes of rs1799964 were associated with a risk effect on femoral head necrosis. The rs1800630 AC genotype was another risk effect associated with femoral head necrosis in cured SARS-infected patients compared to CC genotype.

Severe dengue virus infection. Cascade of cytokine produced TNF and LTA in severe dengue virus (DENV) infections. The TNF rs361525 A polymorphism marking the TNF-4, LTA-3 haplotype, was significantly increased in patients with secondary dengue haemorrhagic fever (DHF) compared to those with secondary dengue fever (DF) in Thais [54]. Two extended MHC haplotypes containing TNF-4 and LTA-3, together with HLA-B48, B57 and DBP1*0501, have been reported only in patients with secondary DHF. These observations indicate that polymorphism in functionally distinct MHC-encoded proteins contributes to the risk of developing severe secondary DENV infection.

Guivier et al. [55] found that two SNPs within the TNF-alpha promoter (~302GG/GG and ~296A/A) were associated with higher TNF-α gene expression and were more frequent in non-endemic areas among European populations of bank voles.

Protozoan parasites

Plasmodium falciparum malaria. Malaria is the most common parasitic disease of the tropics caused by the sporozoa of the genus Plasmodium, is endemic in more than 90 countries, and together with HIV and tuberculosis constitutes one of the major causes of death by infectious diseases worldwide. During P. falciparum malarial infection, TNF has been described as both protective and pathogenic, and at low levels, TNF kills the parasite by macrophage activation and subsequent release of cytokines, whereas high TNF level has been associated with severe manifestations like acute respiratory distress and cerebral malaria. It has been reported that SNPs (rs1799964, rs1799724, rs1800750, rs1800629 and rs361525) in the proximal enhancer of the TNF gene have different associations with malaria in different populations [49, 56–58]. Sinha et al. [59], genotyped these SNPs in patients with P. falciparum malarial infection and controls in Indian population. They found association of the rs1799964 and rs1800630 with increased risk of severe malaria. TNF enhancer haplotype CAGCG (rs1799964, rs1800630, rs1799724, rs1800629 and rs361525) correlated with enhanced plasma TNF levels in both patients with falciparum malarial infection and controls and were associated with increased susceptibility to severe malaria. No association between
rs1800629 polymorphism and susceptibility to cerebral malaria among central Sudanese children was reported [60].

**Mucocutaneous leishmaniasis.** Leishmania braziliensis infection is responsible for MCL. It is a severe form of American cutaneous leishmaniasis (ACL). High serum TNF-α levels have been reported in this disease. So TNF regulatory polymorphism may have some putative role in circulating level of TNF-α and thus in disease manifestation. In Venezuelan case–control study, homozygotes for allele 2 of a polymorphism in intron 2 of the TNF-β gene showed a high relative risk of MCL disease, and a significantly higher frequency of allele 2 of rs1800629 polymorphism was predicted in patients with MCL compared with endemic controls. Polymorphism affecting TNF-α production may be associated with susceptibility to the mucocutaneous disease [10].

**Chagas disease.** The parasite Trypanosoma cruzi causes chronic Chagas disease cardiomyopathy (CCC), affecting 18 million individuals in Latin America. One-third of patients with CCC develop heart failure, and their survival is reduced by 50% compared to patients with other cardiomyopathies. Aguiar and Prestes [61] reported the role of TNF polymorphism in this disease. Elevated TNF-α levels in plasma and heart tissues were observed in patients. The TNF-α such as TNFa2, TNFa microsatellite allele 2 and the TNF2 rs1800629, TNF promoter polymorphism allele 2 were genotyped. Patients positive for TNF2 or TNFa2 alleles display a significantly shorter survival time compared with those carrying other alleles. No association of TNF-α polymorphism with Chagas disease in Brazilian patients have been found [62]. The TNFα microsatellite and rs1800629 polymorphism in an association study were detected. The patients with CCC were grouped in three categories according to degree of left ventricular (LV) dysfunction into severe, mild to moderate and absent. No significant differences between either CCC and asymptomatic (ASY) patients or patients with CCC, according to severity of cardiomyopathy with respect to TNFα or rs1800629 TNF promoter polymorphism, were reported.

**Chronic beryllium disease and beryllium sensitization.** Sato et al. [63] detected the role of TNF-α polymorphism in development of chronic beryllium disease (CBD). They genotyped five TNF-α promoter polymorphism in patients with CBD, sensitized subjects and control subjects and measured TNF-α production in beryllium-stimulated and beryllium-unstimulated BAL. A significantly increased TNF-α production was reported in patients with CBD compared with those only sensitized in beryllium-stimulated, but not beryllium-unstimulated, BAL cell. No significant association has been reported between TNF promoter polymorphism or haplotypes and CBD-sensitized patients, and controls. The rs1799724 T allele has been shown to be associated with BAL cell TNF-α production.

**Human African trypanosomiasis and host inflammatory cytokine response profile.** Lean et al. [54] identified two trypanosomiasis with dramatically different disease virulence profiles in Uganda and Malawi. The two disease profiles were associated with markedly different levels of TNF-α and transforming growth factor β (TGF-β) in plasma. In Uganda, but not Malawi, early-stage TNF-α was elevated, while in Malawi, but not Uganda, early-stage TGF-β was elevated. Rapid progression of disease in Uganda is associated with TNF-α-mediated inflammatory pathology.

**Invasive pulmonary aspergillosis.** The role of TNF-α and lymphotoxin-alpha (LT-α) in fungal infection diseases has been reported [64]. The presence of polymorphism in TNF-α and LT-α genes or their receptors might increase the susceptibility of haematologic patients to develop invasive pulmonary aspergillosis (IPA). SNPs in TNF-α, LT-α and tumour necrosis factor receptor 2 (TNFR2) and a variable number of tandem repeats (VNTRs) in TNFR2 were investigated in haematologic patients and controls. Similar genotype and alleles frequencies were detected between patients and controls. TNF-α and LT-α polymorphisms were not associated with the presence of IPA. A strong association of IPA with VNTR in the promoter region of the TNFR2 gene was found.

**Cancer**

**Breast cancer**

Cancer is the major health problem and leading cause of death. Several genetic polymorphisms have been reported to associated with disease. The genetic factors play important role in the epidemiology and pathogenesis of cancer. TNF genetic polymorphism can regulate gene expression and have been associated with inflammatory and malignant conditions. Azmy et al. [65] have been detected the role of TNF-α rs1800630 and rs361525 polymorphisms in breast cancer susceptibility and severity. Breast cancer cases and controls have shown similar allele frequencies for both polymorphisms. No association was found between rs1800629, rs361525 and susceptibility to breast cancer in North European population.
Role of TNF rs361525 in breast cancer risk was investigated by Gaudet et al. [66], in breast cancer cases and controls, in European, from 30 studies in the Breast Cancer Association Consortium. Jung et al. [67] have detected 12 SNPs in 11 apoptosis-related genes in the apoptosis pathway.

Cervical cancer

Human papillomavirus (HPV) 16 infection is an important factor for cervical cancer. Alteration in local levels of TNF in the cervix may affect the immune response of an individual, hence affecting the persistence of HPV. Excess TNF-α can result in harmful inflammatory responses, whereas too little can contribute to persistent infection. TNF-α is one of the primary cytokines released after HPV infection and upregulates the expression of antigen-processing and presentation pathway components for class I HLA. Eleven TNF SNPs were associated with susceptibility to HPV16-associated cervical cancer. A significant difference in genotype distribution of three SNPs between the cases and controls were reported. Haplotype distribution also showed a significant difference between cases and controls. A new association was reported between several TNF-SNPs and susceptibility to cervical cancer [68].

Prostate cancer

The associations between six TNF SNPs (rs1799964, rs1800630, rs1799724, rs1800629, rs361525 and rs1800610) and prostate cancer risk were investigated [69]. No TNF SNP was associated with prostate cancer risk in Prostate, Lung, Colorectal, and Ovarian Cancer Screening Trial (PLCO), while in the Nutrition Cohort, associations were significant for two highly correlated variants (rs1799724, 1800610). In pooled analyses, no single SNP was associated with prostate cancer risk. No differences in haplotype distribution between case/control status in PLCO, but marginal associations in the Nutrition Cohort and the pooled analysis, were reported. The TNF +488A has been reported to be associated with common variable immunodeficiency in addition to prostate cancer. The association between prostate cancer risk and rs1800629 in 296 patients diagnosed with prostate cancer and in 311 healthy controls was studied.

Bladder cancer

Polymorphism at position TNF-859 shows no disease association. TNF regulatory polymorphism may alter the expression and alter the risk of developing bladder cancer and subsequent tumour behaviour. TNF-α polymorphism, TNF +488A and TNF –859T are significantly associated with risk of bladder cancer.

Non-Hodgkin’s lymphoma

Seidemann et al. [70] studied tumour necrosis factor and lymphotoxin-alpha genetic polymorphism and outcome in paediatric patients with non-Hodgkin’s lymphoma (NHL). The study examines the association of TNF-α rs1800629 and LT-α rs909253 polymorphisms with diagnostic NHL. Patients with Burkitt’s lymphoma (BL) and B cell acute lymphoblastic leukaemia patients carrying at least two variant alleles (high-producer haplotypes) had an increased risk of events. TNF-α rs1800629 and LT-α rs909253 polymorphisms were negative prognostic factors in paediatric BL and in B cell acute lymphoblastic leukaemia (B-ALL).

Pancreatic, lung cancer and renal cell carcinoma

A case–control study of pancreatic cancer was conducted in the San Francisco Bay area by Duell et al. [71]. No association between pancreatic cancer risk and TNF rs1800629 polymorphism was reported. Pancreatitis was significantly associated with TNF rs1800629 GA + AA among patients with pancreatic cancer. A significant difference in genotype frequencies of rs1800629 and rs361525 was reported between patients with lung cancer and the healthy controls and also between patients with lung cancers of various stages. The study was carried out by Shih et al. [72], in 202 patients, 205 controls in Taiwan. Individuals with rs1800629 AA/GA genotypes against GG genotype had higher odds ratios (ORs) while individuals with rs361525 AA/GA genotypes against GG genotype had lower ORs for lung cancer. The patients carrying AA or GA genotype at rs1800629, or a GG genotype at rs361525, had a tendency to advanced disease. A significant association between TNF-α rs1800629 and rs361525 polymorphism and the susceptibility to lung cancer was demonstrated. A case–control study of patients with renal cell carcinoma (RCC) and healthy controls was conducted by Basturk et al. [73]. G-allele frequency of rs1800629 was significantly higher in the patients than in controls. These findings suggest that the TNF-α rs1800629 G/G genotype may be considered as potential risk factors for RCC, whereas TNF-α rs1800629 G/A genotypes may play role as a protective factors. Onishi et al. [74], detected the genetic polymorphism of TNF-α (α1, α2) and TNF-β (β1, β2). All patients having TNF-β-1/1 homozygote were alive, and a significantly favourable prognosis in the patients with TNF-β-1/1 homozygote compared with other TNF-β polymorphism was observed. In the Turkish population, rs1800629 polymorphism is associated with an increased risk of hepatocellular carcinoma as this polymorphism plays role in the regulation of expression level. A case–control study was designed by Akkiz et al. [75], and they found
that rs1800629 genotype was significantly associated with the risk of HCC. The presence of the high producer allele rs1800629 A in the TNF-α gene was associated with an increased risk of the development of HCC in Turkish population.

**Acute pancreatitis.** Tumour necrosis factor α (TNFα) plays important roles in the pathogenesis of acute pancreatitis (AP). Ozhan et al. [76] determined two TNF promoter polymorphisms (rs1800629 and rs361525) in patients with AP and healthy controls. The frequencies of these polymorphisms were similar in both patients with mild or severe pancreatitis and in controls.

**Autoimmune diseases**

**Sarcoidosis**

Sarcoidosis is a complex disease with autoimmune basis, a multisystemic granulomatous disorder which occurs in almost all populations. Disease manifestations are localized to lung and skin, but the involvement of other parts such as eyes, lymph nodes, parotid glands, heart, liver and spleen can also occur. Sharma et al. [25] reported for the first time the association of TNF haplotypes and genotypes with sarcoidosis and its prognosis in the Indian population. Five promoter polymorphism in the TNF-α gene and one in LTA gene (rs909253) were genotyped in North Indian patients. They have measured sTNF-α and serum angiotensin-converting enzyme (SACE) levels. Serum TNF-alpha and SACE levels are influenced by rs1800629 and rs361525 polymorphisms. The patients and controls have significant differences in haplotype frequency. The haplotype GTCCGG was identified as the major risk/susceptibility haplotype and was associated with increased SACE levels in the patients.

Cystic fibrosis conductance regulator, tumour necrosis factor, interferon-alpha-10, interferon-alpha-17 and interferon-gamma genotyping as potential risk markers in pulmonary sarcoidosis pathogenesis were detected by Makrythanasis et al. [77], in Greek patients. They have detected a statistically significant increase of CFTR mutation carriers in patients with sarcoidosis than in the control population. A difference was observed within sarcoidosis patients group where patients with CFTR mutations suffered more frequently from dyspnoea than those without.

**Multiple sclerosis**

Tumour necrosis factor (TNF-α), a proinflammatory cytokine, plays an important role in multiple sclerosis (MS) pathogenesis. In Turkish population, Akcali et al. [78] reported that rs1800629 and rs361525 polymorphisms are not associated but rs1799724 allele is associated with the multiple sclerosis disease. A statistically significant increase in rs1799724 CC genotype was found in MS patients than in controls, while rs1799724 CT genotype showed a significant negative correlation with patients with MS. No differences in the distribution of rs1800629 and rs361525 alleles were observed. None of the three polymorphisms (rs1800629, rs361525 and rs1799724) showed relation with disease. Significant difference of rs1799724 CC genotype was identified with the low disease index. Thus, rs1799724 CC genotype may cause susceptibility to MS in the Turkish population. TNF-β and TNF-α gene (rs1800629 and rs361525) polymorphisms and susceptibility to MS were determined in Caucasian patients with MS, and healthy controls from Norway [79]. TNF-β genotypes were significantly associated with MS. TNF-α genotypes were not associated with MS. Huizinga et al. [80], reported TNF-α promoter polymorphism and susceptibility to multiple sclerosis in different groups of patients. TNF-α production in whole blood cultures upon stimulation with LPS was determined in individuals from 61 families. Highest TNF production is characterised in three families, and in contrast, the lowest TNF production is characterised in three families. The difference of highest and lowest TNF production could not be attributed to the promoter polymorphism rs1800629, rs361525 or rs1800750, although rs361525 GA donors produced low TNF upon culture with endotoxin compared with TNF rs361525 GG donors. The frequency of the rs361525 GG genotype was increased in patients with MS in a nursing home compared to patients with MS in an outpatient’s clinic or Dutch controls. TNF-α rs1800629 and rs361525 polymorphisms have no association with MS, but the microsatellite allele a11 is associated with the disease in French patients [81]. In French patients with MS and controls, TNF-α rs1800629 and rs361525 and a microsatellite polymorphisms were investigated. TNF-α rs1800629 and rs361525 polymorphisms have shown no significant differences between patients with MS and controls. Very significant association was found between allele frequency for the a11 allele and MS.

**Rheumatoid arthritis**

Rheumatoid arthritis (RA) is a type of systemic autoimmune disease. Rheumatoid arthritis has both environmental and genetic background, with genetic factors contributing 15–30% of the overall risk. The genetic studies have given different associations in different populations. The TNF +488A have been reported to be associated with rheumatoid arthritis [82], while TNF +489 polymorphism does not contribute to susceptibility to rheumatoid arthritis in Europeans. In Caucasian TNF, rs1800629 polymorphism is not associated with response to TNF-α blockers in patients with rheumatoid arthritis.
and does not serve as a genetic risk factor for RA susceptibility and severity in Americans. TNF2 allele has been associated with susceptibility and severity to a variety of autoimmune and inflammatory diseases, including RA, systemic lupus erythematosus and ankylosing spondylitis. The presence of TNF2 allele increases the production of TNF-alpha and thus increases the host’s resistance to infection. Aguillon et al. [82] suggested that RA is favoured by the presence of the rs1800629 polymorphism and is responsible for increased TNF production. Ten European, three Latin American and one Asian studies were analysed by Lee et al. [83], and no association was found between RA and the TNF-α rs1800629 A-allele in the overall population.

**HLA-B27–positive ankylosing spondylitis**

The association between TNF-α promoter polymorphism and ankylosing spondylitis (AS) susceptibility was reported with inconsistent results. Chung et al. [84] conducted a case–control study including six TNF-alpha promoter polymorphism. They found a significant difference in the allelic and genotypic frequencies at rs1799964, rs1799724 and rs1800750 in patients with HLA-B27 (+) and AS and random controls, but not in patients with AS and HLA-B27 (+) healthy individuals. Haplotype (rs179964 T/rs1799724 C/rs1800630 C/rs1800629 G) increases the risk of susceptibility to AS compared to random controls, whereas haplotype (rs179964 C/rs1799724 A/rs1800630 C/rs1800629 G) have shown to be associated with decreased susceptibility to AS compared to random controls. One Latin American and seven European studies were analysed by Lee and Song [85]. No association between AS and rs1800629 A-allele, AA and AA + AG genotypes were reported.

**Graves’ disease**

In the development of Graves’ disease (GD), a role is played by TNF-α. Gu et al. [86] investigated the association of TNF-α polymorphism rs1800629, rs361525 and rs3093661 with GD in Chinese population. A significant difference in distribution of rs361525 and rs3093661 allelic frequencies between Graves’ disease and control individuals was reported. The G-alleles of rs361525 and rs3093661 SNPs have been associated with higher risk of GD as compared with A-alleles. No significant difference of rs1800629 allelic frequency was observed. The haplotype GGG was associated with an increased risk of GD, whereas the haplotype GAA was protective.

**Type 1 diabetes mellitus**

Type 1 diabetes mellitus (TIDM) is an autoimmune disorder, which involves T cell-mediated destruction of the pancreatic β-cells [87]. Several reports had shown the association of polymorphism with the disease TIDM [87–90]. The proinflammatory cytokines are elevated in patients at the onset of diabetes. A significant increase of rs1800629 G/A and A/A genotypes in North Indian patients with T1DM were reported [91]. Das et al. [92] suggested a significant association of rs1800629 A-allele and G/A genotype with T1DM in North Indians, but no association with rs361525 polymorphism. The same increase in the prevalence of rs1800629 A-allele in patients with diabetes in the Hungarian population was reported [93]. In Chinese and Caucasian populations, rs1800629 variant was not considered as a genetic factor for susceptibility to T1DM and the associations of the TNF gene are because of a Linkage disequilibrium (LD) between TNF and DR3-DQB 1*0201 haplotype reported [94]. Initially, rs1800629 and rs361525 variants show association with T1DM, but after adjusting the data for LD with DRB1-DQB1 and B18-DR3 haplotypes, the association lost its significance [93]. Boraska et al. [95] studied relation of TNF gene promoter polymorphism (rs1800629 and rs361525) with TIDM in a case–control study from South Croatia. Haplotype (rs1800629 A and rs361525 G) was observed more often in patients with TIDM than in controls. SNP rs1800629 was found to be more frequent in patients with TIDM. The author did not find strong evidence of association of TNF promoter polymorphism with TIDM. Independent association of TNF polymorphism with type 1 diabetes susceptibility have been found in Korean [96]. Seven SNPs in the TNF genes (TNFα and TNFβ) were genotyped in a Korean, along with HLA DRB1, DQB1 and MICA (MHC class I chain–related genes). Three SNPs and two common TNF haplotypes showed significant association with the risk of TIDM.

**Type 2 diabetes**

In case of type 2 diabetes, high levels of cytokines have been considered as risk factors. Kubaszek et al. [97] investigated TNF-α and IL-6 polymorphisms and found that TNF-α rs1800629 A-allele was associated with an approximate twofold higher risk of type 2 diabetes compared with the rs1800629 G. The rs1800629 A-allele of TNF-α rs1800629 polymorphism is a predictor for the conversion from IGT (impaired glucose tolerance) to type 2 diabetes.

**Diabetic nephropathy**

In diabetic nephropathy, glucose auto-oxidation and production of free radicals causes protein glycation that increases the concentrations of proinflammatory cytokines. Myeloperoxidase (MPD) is a heme enzyme, participating in microorganism killing by phagocytes. Patients with chronic renal failure results from diabetic
nephropathy show a significant reduction in the intracellular myeloperoxidase level and myeloperoxidase gene promoter polymorphism (-463, G/A) causes a decreased gene expression. In a case–control study, no significant differences in TNF genotype and allele frequencies between the groups and patients with diabetic nephropathy were found. A lower frequency of TNF1/TNF1 genotype has been reported [98]. Significant differences of TNF plasma level in patients with diabetic nephropathy and other renal diseases were reported. A statistically significant difference in MPO genotype frequencies between patients with diabetic nephropathy and patients with other renal diseases was observed. MPO, GG and AA genotypes were significantly more common in patients with diabetic nephropathy. A correlation between the MPO genotype and an earlier onset of the disease was observed while such a relationship was not found for the TNF genotype. It has been found that in patients with diabetic nephropathy, TNF variants were more frequent than in non-diabetic patients with chronic renal failure.

Inflammatory bowel disease

Crohn’s disease is a chronic inflammatory disease of the intestines. This is closely related to another chronic inflammatory condition that involves only the colon called ulcerative colitis. Together, CD and ulcerative colitis are referred to as inflammatory bowel disease (IBD). The chromosomal region, 6p21, IBD3, has been identified as an IBD-susceptibility locus [99–101]. IBD3 region encompasses the tumour necrosis factor 2 (TNF) gene. TNF-alpha is considered as a strong candidate gene for IBD. Levels of TNF are elevated in the serum, mucosa and stool of patients with IBD. TNF production is under a strong genetic influence [102]. The positive association of TNF rs1799724 C with UC was reported in Caucasians and is also supported by a small Japanese case–control study. The same study reported an association of TNF rs1799724 T with Japanese CD, although a significant effect of this allele was not observed in a larger patient cohort [103].

TNF-α and Fc-gamma receptor polymorphism with infliximab in Crohn’s disease

The associations between TNF-alpha and Fc-gamma receptor (Fc-gammaR) polymorphism with infliximab (IFX) treatment for CD are not well known. Patients with CD were given IFX 5 mg/kg intravenously and followed prospectively for 8 weeks, and the Crohn’s disease activity index (CDAI) was measured before and after 8 weeks of treatment [104]. On the basis of predicted CD activity index, patients were grouped as responders or non-responders. The TNF-alpha, Fc-gammaRIIA and Fc-gammaRIIIA genotype distribution was not significantly different between responders and non-responders 8 weeks after treatment. Fc-gammaRIIIB genotype distribution has shown significant differences between responders and non-responders after 8 weeks. Fc-gammaRIIIB polymorphism may be an important factor for clinical response to IFX treatment in CD.

Asthma

Asthma is a complex polygenic disease in which gene–environment interactions have shown to play important role. TNFα gene is one of the important candidate genes involved in pathogenesis of asthma. Several studies have investigated TNFα gene polymorphisms (rs1800629 G designated as TNF1 and rs1800629 A designated as TNF2) and asthma susceptibility in different populations. A positive association between TNF2 and asthma [79, 105–112] have been reported. Some studies have been reported a negative association [113–116], and one study reported a positive association between TNF1 and asthma [117]. Gao et al. [118] included 2409 patients with asthma and 3266 controls, in the study. They found that TNF2 allele confers a significant risk of developing asthma.

Risk of primary biliary cirrhosis

Juran et al. [119] recently reported an association between primary biliary cirrhosis (PBC) and (rs231725) polymorphism of the immunoreceptor gene cytotoxic T-lymphotope antigen 4 (CTLA4). They have detected its interaction with the rs1800629 polymorphism in which TNF2A allele has been shown to increase the TNF production. The genotyping of polymorphism was carried out in patients with PBC and in controls from US and Canada. Allele frequency for TNF2A was elevated in patients with PBC, but only borderline significance was detected. TNF2A carriage was significantly increased in CTLA4 ‘A/A’ PBC patients compared with CTLA4 ‘A/A’ controls; no apparent increase in TNF2A carriage was shown in CTLA4 (A/G or G/G) individuals. TNF2A amplifies the CTLA4 (rs231725, A/A) genotype risk of PBC.

Behcet’s disease

Behcet’s disease (BD) is a chronic multisystem inflammatory disorder, the hallmarks of which are recurrent oral and genital ulceration, skin lesions, and uveitis. It has been reported that rs1799964 polymorphism has been associated with Behcet’s disease [120].

Dermal diseases

Atopy on irritant contact dermatitis in health care workers

Davis et al. [121] studied the effects of TNF-alpha G to A rs1800629 polymorphism on chronically damaged skin
of healthcare workers. They have genotyped TNF-alpha rs1800629 polymorphism and measured the epidermal response. Excess hand erythema decreased with hand hygiene exposure and increased during time off for AA/GA genotypes, but had opposite effects for GG. AA/GA had smaller reductions in dryness with lotion treatment and larger reductions in excess erythema than GG. Repeated exposure to water and sodium laurel sulphate produced higher erythema in normal skin for AA/GA than for GG genotype. The study suggested that the TNF-alpha rs1800629 polymorphism and an atopic history influence the severity of irritation and recovery from exposure.

Risk of psoriasis vulgaris

Several studies have given different association between TNF-α polymorphism and psoriasis risk. The rs1800629 and rs361525 polymorphisms have been reported to influence the transcription of the TNF-α gene and have been implicated in psoriasis risk. Li et al. [122] conducted psoriasis case and control study. The rs361525 GA + AA genotypes had significantly increased risk, compared with the GG genotype, whereas a significantly reduced psoriasis risk was associated with rs1800629 GA + AA genotypes compared with the GG genotype.

Tumour necrosis factor-α antagonists are effective in the treatment for refractory psoriasis. In many diseases such as rheumatoid arthritis, ankylosing spondylitis, and CD, treatment with this therapy results in induction of such as rheumatoid arthritis, ankylosing spondylitis, and many other diseases, including cardiovascular diseases (CVDs). Yilmaz et al. [124] have evaluated the role of cytokine gene polymorphism in carotid intima-media thickness (CIMT) and left ventricular mass index (LVMI) progression in non-diabetic haemodialysis (HD) patients. TNF-α and IL-10 polymorphisms were determined in the study. Risk factors for cardiovascular diseases have no difference between TNF-alpha rs1800629 high-/low-producer genotype groups. CIMT and LVMI progressions were detected at higher levels in patients with high-producer genotypes (AA + AG) than in patients with the low-producer genotype (GG). The rs1800629 polymorphism was strongly associated with C-reactive protein (CRP). Analysis also showed that the combination of high production of TNF-α and low production of IL-10 was associated with higher average IMT, LVMI progression and elevated average CRP levels compared with a combination of low production of TNF-α and high production of IL-10.

Spontaneous deep intracerebral haemorrhage

Association of TNF-α gene with spontaneous deep intracerebral haemorrhage was investigated by Chen et al. [125] in the Taiwan population. Deep parenchymal structure including the basal ganglia, thalamus, brainstem and cerebellum is the most frequently affected site of spontaneous intracerebral haemorrhage (SICH). Rost et al. [126] comprehensively reviewed the candidate genes of SICH reported during 1996–2007. Reported candidate genes that show association with SICH were involved in the pathways of the vessel wall integrity (ACE, APOE, nephrilysin, endoglin, TGFB1), endothelial dysfunction (ACE), inflammation markers (IL-6, TNF) and haemostasis (APOE, CD-14, Factor VII and XIII, VKORC1). Spontaneous deep intracerebral hemorrhage (SDICH) risks were positively associated with TNF (rs1799964 C and rs1800629 A) in men but inversely associated with (rs1800630 A) in females [126]. There were significant interaction effects between gender and SNPs (rs1799964, rs1800630 and rs1800629) on SDICH risks.

TNF-α polymorphism and homocysteine levels in patients with ischaemic strokes

Kim et al. [127] carried out case–control studies including patients with ischaemic stroke, patients with silent
brain infarctions SBIs and controls. Significant differences in the frequency of the TNF-α rs1800629 polymorphism were found between the patients with ischaemic stroke and the control group. The frequency of the TNF-α (rs1800629 GA + AA) genotype was higher in the group having highest homocysteine (tHcy) levels than in the group having lowest tHcy levels. The tHcy levels were significantly and inversely correlated with folate levels in the TNF-α (rs1800629 GG) and TNF-α (rs361525 GG) genotypes in the ischaemic stroke, SBI and control groups. TNF-α rs1800629 polymorphism is responsible for susceptibility to ischaemic stroke and is associated with high tHcy levels in Koreans. No relationship between TNF-α polymorphism and SBI susceptibility was found in this study.

Neurological/behavioural disorders

Alzheimer disease

Alzheimer’s disease (AD) is one of the most common types of chronic neurodegenerative diseases. Vascular dementia, AD and stroke are all associated with inflammation, but they have different initiating factors. Polymorphism in the TNF and apolipo protein E (APOE) was reported to increase AD risk. Laws Simon et al. [128] conducted a case–control study and investigated −850C>T, rs1800629 in TNF and the APOE polymorphism in controls and patients with sporadic AD. The frequency of (−850C/T) genotypes and T allele was significantly different in AD individuals, while the (rs1800629) SNP was not associated with AD. T allele of (−850) polymorphism significantly modified risk associated with possession of the APOE ε4 allele only, and (−850) T allele was found to be associated with lower levels of CSF Aβ42. In a Southern China population, patients with sporadic Alzheimer’s disease (SAD) have a significantly increased frequency of rs1800629 A-allele as compared with controls. The carriers of A-allele have a significantly increased risk of SAD. Level of TNF-α in serum of SAD group was much higher than that in control group, and the elevated serum TNF-alpha level was closely associated with the risk of SAD detected by Yang et al. [129]. Seventeen studies that investigated the association between five TNF-α polymorphism (−850, rs1800629, rs1800630, rs361525 and rs1799964) and AD were retrieved and analysed [130]. The presence of T allele significantly increased the risk of AD associated with carriage of the apolipoprotein E epsilon 4 allele in Caucasian Australians and Northern Europeans. A significant association of (−850) polymorphism with AD risk and non-significant difference in genotype distribution of (rs1800629) polymorphism in AD was found. For the (rs361525 and rs1799964) polymorphism, Di Bona et al. [130] did not find an association with AD. Only four studies investigated rs361525 variant, and the results were not significant. Current findings suggested an association between (−850C>T) polymorphism and the risk of developing AD. No positive associations between TNF-alpha promoter haplotypes and AD disease in Italian population have been reported by Tedde et al. [131].

Event-related potential indices of attention and mental rotation

Tumour necrosis factor plays an important role in glutamatergic neural transmission [132] and serve essential functions in neural plasticity [133] and cognitive processes like learning and memory [134, 135]. SNPs in TNF have profound impact on this disease. A-allele of rs1800629 fastens cognitive processing speed in a visual task, compared with G-allele carriers [136]. Mental rotation describes the cognitive process of imagining an object turning around. Mental rotation is usually examined using objects (e.g. letters) that are rotated by certain degrees clockwise or counter-clockwise from the vertical upright. This angular displacement allows a parametrical modulation of task difficulty and the demand of processing in working memory. Working memory processes are closely interrelated to attentional processes as attention permits information to be further stored and processed in working memory. Attentional processes are reflected by the visual N1 event-related potential (ERP)-component. The visual N1 may reflect effects of attention on sensory processing or an integrated process of perception and attention. The visual N1 is an exogenous potential that is modulated by attentional processes modifying the magnitude of neural responses to incoming information.

Beste et al. [136] examined the association of the TNF-α rs1800629 polymorphism with attention and mental rotation performance in an event-related potential (ERP) study in healthy participants. The results show that carriers of rs1800629 A-allele display elevated attentional processes as compared to the GG genotype group. Carriers of the rs1800629 A allele performed better than the GG genotype group. The finding of enhanced attentional and mental rotation performance in A-allele carriers supports recent findings that the A-allele of this SNP enhances cognitive performance on a general measure of cognitive processing speed.

Labile anger

Interferon-alpha increases the expression of TNF-α. During interferon-alpha therapy in psychiatric symptoms, TNF-α polymorphism played a role in susceptibility to this disorder. Recently role of TNF-α rs1800629 polymorphism in labile anger and depression was investigated by Lotrich et al. [137]. A-allele of rs1800629 was associated with worsened labile anger and fatigue during
treatment but not with major depression incidence or increased Beck Depression Inventory II. Labile anger was not predicted by the serotonin transporter polymorphism. During treatment with an exogenous cytokine, vulnerability to worsening labile anger distinct from major depression is associated with genetic variability in TNF-α.

Neuropathic pain

Tumour necrosis factor-alpha has been reported to play a role in neuropathic pain. Leung and Cahill [138] described the role of TNF-α in neuropathic pain. Neuropathic pain is pathological pain where nociceptive responses persist beyond the resolution of damage to the nerve or its surrounding tissue. Animal models of neuropathic pain based on various types of nerve injuries have persistently implicated a pivotal role for TNF-α at both peripheral and central levels of sensitization.

Increased risk of new haemorrhage

Achrol et al. [139] identified SNPs associated with increased risk of new intracranial haemorrhage (ICH) after brain arteriovenous malformation (BAVM). Achrol et al. [125] investigated four promoter SNPs in interleukin-6 and tumour necrosis factor (rs1800629, rs361525). An association has been found between TNF-α rs361525 polymorphism and increased risk of new ICH after diagnosis. The patients with TNF-α rs361525 AG genotype had increased risk of new ICH. No other SNP was found to be associated with new ICH.

Others

Endometriosis

Genetic factors play role in endometriosis [5, 140]. Elevated levels of TNF-α in peritoneal fluid of women with endometriosis have been reported, suggesting that TNF-α may be involved in the development of endometriosis. Asghar et al. [5] investigated the possible association between endometriosis and the TNF-α gene promoter polymorphism rs1799964, rs1799724, rs1800629, rs361525 and rs1800630 in a Japanese population. No significant differences in frequencies between the crude endometriosis cases and controls were reported for the above-studied polymorphism. Division of endometriosis group in a subgroup of women with stage IV disease only, the frequency of rs1799964 C allele, was significantly lower in this subgroup than controls. Therefore, the study suggested that the TNF-α rs1799964 polymorphism might be associated with susceptibility to endometriosis.

Muscle phenotypes

During ageing, there is 2- to 4-fold increase in plasma levels of inflammatory mediators such as TNF-α, IL-6, interleukin 1 receptor antagonist (IL-1Ra), soluble TNF-α receptor (sTNFR), acute-phase proteins, such as C-reactive protein (CRP), and neutrophils has been reported. This low-grade inflammation may play an important role in age-related diseases such as Alzheimer’s disease, atherosclerosis, type 2 diabetes, osteoporosis, as well as sarcopenia. TNF-α played role in many age-related inflammatory changes, whereas other cytokines like IL-6, IL-1Ra, sTNFR, as well as acute-phase proteins (APPs) like CRP, reflect responses to upregulated local or generalized TNF-α activity [141]. The authors have detected five TNF promoter SNPs, including rs1799964, rs1799724, rs1800629, rs361525 and rs1800630. The rs1799964 and rs1800630, putative high-expression alleles individually or in the haplotype rs1799964 C-rs1800630 A- rs1799724 C- rs1800629 G- rs361525 G, were associated with lower muscle mass in men. Carriers of rs1799964 C, compared with non-carriers, exhibited lower arm muscle mass also tending to be lower. Similarly, rs1800630 A allele carriers (linked with rs1799964), compared with non-carriers, exhibited lower arm muscle mass. Carriers of the haplotype rs179964 C-rs1800630 A- rs1799724 C- rs1800629 G- rs361525 G, compared with non-carriers, exhibited lower arm muscle mass and trunk muscle mass.

Paget’s disease

Interleukin (IL)-6, a cytokine, plays an important role in the differentiation and activation of osteoclasts and might be involved in osteoblast stimulation in Paget’s disease of bone (PDB). Corral-Gudinol et al. [142] investigated the association of IL-6, IL-8 and TNFα (rs1800629 and rs361525) polymorphism in patients with PDB and healthy controls in Spanish population. No significant association between genotype and allele distribution of any of the cytokines polymorphism and PDB was observed. The study concluded that Paget’s disease of bone is not associated with polymorphism in interleukin-6, interleukin-8 and tumour necrosis factor-alpha genes.

Proliferative vitreoretinopathy disease

Genetic factors have role in proliferative vitreoretinopathy (PVR). Eleven SNPs in the lymphotoxin-alpha (LTA), TNF-α, leucocyte-specific transcript 1 (LST1) and the activating natural killer receptor p30 (NCR3) were analysed, and a significant association in the non-synonymous polymorphism rs2229094 (T/C causes cysteine to arginine change in the signal peptide) in the LTA gene has been reported [143]. This marker was also present in all significant
haplotypic associations and was not observed in any non-significant associations. The strong association found in the rs2229094 (T/C) of the LTA gene may indicate an important role of this polymorphism in the development of PVR.

Risk of fracture in older women

Tumour necrosis factor-α is a proinflammatory cytokine that promotes osteoclastic bone resorption. Moffett et al. [64] detected the association between TNF rs1800629 polymorphism and osteoporosis phenotypes in older women. Women with the A/A genotype had greater subperiosteal width and endocortical diameter than those with the G/G genotype, and there was a greater distribution of bone mass away from the neutral axis of the femoral neck in women with the A/A genotype, resulting in greater indices of bone bending strength. TNF rs1800629 polymorphism was not associated with a reduced risk of other fractures. A potential role has been played by TNF-α polymorphism in the aetiology of osteoporosis.

Susceptibility to oral lichen planus

Kimkong et al. [144] investigated the association between oral lichen planus (OLP) susceptibility and clinical type in the Thai population and found a higher proportion of TNF-α, rs1800629 AA genotype (high producer genotype) among patients with PDB when compared to healthy controls. For polymorphism (rs1800630 and rs361525), no significant association with OLP development was reported. Thus, in Thai population, TNF-alpha rs1800629 AA genotype might play a role in the susceptibility and severity of OLP.

Idiopathic recurrent miscarriage

Reports indicated that approximately, 1–3% of healthy women experienced recurrent miscarriage (RM), defined as three or more consecutive pregnancy losses prior to the twentieth week of gestation. Zammiti et al. [145] reported that high expression of tumour necrosis factor (TNF)-α and lymphotoxin-α (LT-α) was associated with pregnancy complications, including idiopathic recurrent

Figure 3. Tumor necrosis factor binding to its receptors eliciting an immune response.
| Polymorphism | Disease associations/no association (infectious) | P-values and OR | Population | Cases/controls (n) | References |
|--------------|-----------------------------------------------|---------------|------------|-------------------|------------|
| rs1799964, rs1800629, rs361525, rs1800630 and rs909253 | No association with tuberculosis | 0.78 | North Indian | 185 patients and 155 controls | Sharma et al. [34] |
| rs1800629 and rs361525 | No association with tuberculosis | ns | Turkey | 128 TB patients and 80 controls | Ates et al. [27] |
| rs1800629 | No statistical differences in haplotypes frequences between patients with MB and PB | ns | Afro-and Euro-Brazilians | 631 patients with leprosy | Vanderborght et al. [39] |
| rs1799964, rs1800629, rs361525 | Higher symptom scores of rAOM | <0.02 | USA | 128 subjects | McCormick et al. [167] |
| rs1800750, rs361525 | Significant differences in allele frequences between controls and patients with typhoid fever | >0.05 | Vietnam | 380 patients and controls | Dunstan et al. [44] |
| rs1800629 | No significant differences in TNF-α genotype frequences. The rs2430561 TT genotype, associated with an increased production of IFN-gamma, and significantly less frequent in patients with MSF than in the control group | ns, for inter group comparison | Iranian | 100 patients 89 healthy controls | Somi et al. [52] |
| rs1800629 G (TNF1), rs1800629 A (TNF2) | TNF2 is not associated with predisposition to CM. The rs1800629 genotypes and alleles did not differ significantly between patients with CM and controls | 0.0009 | Thais | 435 with Dengue infection | Vejbaesya et al. [170] |
| rs1799964, rs1799724, rs1800750, rs1800629 and rs361525 | Significant differences in TNF levels were seen between genotypes of both ~1031 and ~863 in malaria cases and controls | rs1799964 P = 0.0178, rs1800630, P = 0.0481 | Indian | 121 patients and 121 controls | Sinha et al. [59] |
miscarriage (RM). TNF-\(\alpha\) (rs361525, rs1800629) and LT-\(\alpha\) (rs909253) polymorphism were investigated in RM and control women. Higher frequency of rs361525 A, but not the rs1800629 A or the LT-\(\alpha\) rs909253 G, allele was reported in patients. The rs361525 G/G was lower in patients. Association of the rs361525 SNP with idiopathic RM was confirmed by regression analysis. Haplotypes rs1800629 A\(⁄\)rs361525 G\(⁄\)rs909253 G and rs1800629 G\(⁄\)rs361525 A\(⁄\)rs909253 G played a susceptible role in idiopathic RM. Palmirotta et al. [146] reported that TNFA gene promoter polymorphism and susceptibility to recurrent pregnancy loss in Italian women.

Discussion

Tumour necrosis factor a pleiotropic cytokine regulating a broad range of biological activities including inflammation (Fig. 3). The inflammatory effects of TNF are mediated mainly by activation of the transcription factor NF-\(\kappa\)B and mitogen-activated protein kinases (MAPKs). TNF binds to TNFR1 (receptor subunits, p55–p60) and TNFR2 (receptor subunits, p75-p80). TNFR1 is constitutively expressed by most cell types, and TNFR2 expression is induced mainly in immune and endothelial cells. Therefore, any factor that affects the serum level of TNF might be important in outcome of disease. The presence of polymorphism in regulatory region may alter regulation expression level and thus affect disease manifestation.

Regulation of gene expression at the level of transcription is the most important. Several mechanisms play important role in gene regulation. Promoter hypermethylation and the presence of polymorphism in regulatory region are the two most important factors that interfere with the gene regulation, and our hypothesis is that during disease conditions, there is upregulation or downregulation of gene (transcriptional dysregulation). For example, infection of Plasmodium falciparum upregulates the expression of TNF-\(\alpha\). In certain cancers, there is downregulation of genes. Promoter polymorphism lies in transcription factor–binding sites (cis elements) and hypermethylation of CpG (CG dinucleotide) islands can contribute to dysregulation. CpG islands are present in the promoters regions of almost half of mammalian genes [147], leading to gene silencing. DNA methylation could repress transcription by either or both of the following two mechanisms: (1) the methyl group may disrupt the binding sites of the transcription factors (TFs) and result in the failure of transcription [148–150]. (2) Methylated cytosines may attract methyl-CpG-binding domain proteins (MBD) which would bring repressors to silence the gene [151]. Methylation in CpG Island interferes with gene regulation and thus responsible for outcome of disease. There is an inverse correlation of DNA methylation
with gene expression. High levels of DNA methylation in CpG-rich promoters are strongly associated with downregulation of gene expression [151]. The several reports observed a strong anticorrelation of DNA methylation with CpG density and GC content, indicating that DNA methylation declines the more the sequence resembles a CpG island [45, 151, 152]. Methylation levels of 1320 CpG sites in regulatory regions of 416 genes in cells from acute lymphoblastic leukaemia (ALL) children were studied by Milani et al. [153].

A large number of diseases have been reported, in which promoter polymorphism affects the susceptibility to diseases (Tables 1–3). DNA sequence variations (polymorphism) that affect the transcription of genes play important roles in the pathogenesis of many complex diseases [154]. While most discovered genetic defects create missense or non-sense substitutions in protein-coding sequences, there remain disease-associated genes for which there is no difference in protein-coding information between individuals of different phenotypes. Examples of the latter include CD36 type II deficiency [155], phenotypic variances in plasma dopamine-beta-hydroxylase [156] and serum angiotensin I converting enzyme levels [157]. The gene for TNF is polymorphic. Several TNF promoter SNPs have been reported to be associated with disease in humans. DNA sequence variations modifying transcriptional regulation of gene [154] play important role in many complex diseases. The first 200 bp of the promoter are highly conserved across a range of species, with the murine, bovine and porcine promoters showing approximately 80% homology with the human promoter; while further upstream, there is far less conservation between species. It has been reported that TNF rs1800630 polymorphism was associated with reduced level of serum TNF-α, because this polymorphism is strongly influence the binding of nuclear proteins [158].

In gene expression, the multiple TFs first assemble at the promoter site and the recruit RNA polymerase. These TFs bind to their cognate binding sites in the promoter region. The presence of polymorphism in regulatory region affects the interaction of TFs with transcription factor–binding site (TFBS), influencing the expression of gene and thus susceptibility/resistance to disease. We have also predicted several SNPs in the promoter of

### Table 2: Details of polymorphism, P-values and OR, cancer risk association, population type and case studies with references.

| Polymorphism | Disease associations/No associations (cancer risk) | P-values and OR | Population | Cases and controls (n) | References |
|--------------|--------------------------------------------------|----------------|------------|------------------------|------------|
| rs1800630, rs361525 | No association with breast cancer | OR = 0.82 (0.64–1.04), P = 0.637 | North European | 709 patients and 498 controls | Azmy et al. [65] |
| rs361525 | Evidence against an overall association between invasive breast cancer risk and TNF rs361525 | OR 1.00 (0.95–1.06) | Europe, USA and Australia | 30,000 cases and 30,000 controls | Gaudet et al. [66] |
| rs1800629 | Genotype distribution of 3 SNPs differed significantly between case subjects and 1 or both of the control groups | <0.05 | New Mexico | 541 cases and 241 controls | Deshpande et al. [68] |
| rs199724, rs1800610 | Significant association with prostate cancer risk | | | | |
| rs1800629 | No overall association between pancreatic cancer risk and TNF-α (rs1800629 G/A) | | | | |
| rs1800630, rs361525 | rs1800630, rs361525 were significantly different between the patients with lung cancer and controls | 0.0001 | Taiwan | Patients 202, controls 205 | Shih et al. [72] |
| rs1800629 | rs1800629 G-allele significantly higher in RCC patients compared to controls | Significant | Turkey | 29 patients, 50 controls | Bağtırık et al. [73] |
| rs1800629 | rs1800629 was significantly associated with the risk of HCC | <0.001, odds ratio [OR] = 4.75 | Turkish | 110 patients and 110 control | Akkiz et al. [75] |
| rs1800629, rs361525 | Both SNPs of TNF-α are not genetic risk factor for AP susceptibility | OR = 1.63; 1.13–4.01 for rs1800629 and OR = 0.86; 0.75–1.77 for rs361525 | Turkey | 105 patients with acute pancreatitis (AP) and 92 controls | Ozhan et al. [76] |
| rs1800629, rs1143627, rs1800896, rs361525 | rs1800629 confer a higher risk of HCC. Other polymorphisms were not related to risk of HCC in this study | OR = 1.74 | 20 studies | 2763 patients with HCC and 4152 controls | Yang et al. [166] |

RCC, renal cell carcinoma; HCC, hepatic cell carcinoma; AP, acute pancreatitis; TNF-α, tumor necrosis factor-alpha.
| Polymorphism          | Disease associations/No associations (others)                                                                                                                                                                                                 | P-values     | Population | Cases and controls (n) | References            |
|----------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|--------------|------------|------------------------|-----------------------|
| TNF, IFNA10, IFNA17, IFNG genes | A statistically significant increase of CFTR mutation carriers in the population of patients with sarcoidosis versus the control population was found                                                                                           | 6.1 x 10^-8 | Greek      | 89 Greek patients with sarcoidosis and 212 control           | Makrythanasis et al. [77] |
| rs1799724, rs1799724, rs1800750 | Significant increase in rs1799724 CC genotype in ms. The rs1800629 and rs361525 Not associated with MS                                                                                                                                   | <0.001       | Turkish    | 86 patients and 150 controls                                     | Akcali et al. [78]       |
| rs1800629, rs361525, rs3093661 | Haplotype rs1799964 T/rs1800630 C/rs1799724 C/rs1800629 G increased the risk of susceptibility to AS compared to random controls G-alleles of rs361525 and rs3093661 SNPs have been associated with higher risk of Graves’ disease as compared with A-alleles. No significant difference of rs1800629 allele frequency | P (corr) < 0.001, OR = 2.756 | Chinese    | 119 patients, 95 healthy controls, and 135 random healthy controls | Chung et al. [84]       |
| rs1800629, rs361525 | rs1800629 A-allele is associated with overall susceptibility to asthma OR = 1.37, P = 0.04                                                                                                                                            | OR = 1.37, P = 0.04 | China       | 2409 patients and 3266 controls                                | Gao et al. [118]         |
| rs1800629 | A-allele increases the production of TNF-a. TNF2A allele frequency was elevated in patients with PBC                                                                                                                                  | OR = 1.21, P = 0.042 | US and Canada | 866 patients with PBC and 761 controls                        | Juran et al. [119]       |
| rs1799964 | rs1799964 associated with bowel disease 0.00004 in UK white Caucasian USA 68 healthcare workers with irritant hand dermatitis                                                                 | OR = 2.60 | Various ethnicities | 997 cases, 943 control for rs361525 and 1156 cases, 1083 control for rs1800629 | Davis et al. [121]       |
| rs1800629, rs361525 | Significantly increased risk was associated with the variant GA + AA genotypes of rs361525, compared with the GG genotype and significantly reduced psoriasis risk was associated with the variant GA + AA genotypes of the rs1800629, compared with the GG genotype OR = 2.60 | Significant Swedish | 1209 subjects with MI | Mellick [62] |
| rs1800630, rs1800629 | Association between TNF haplotype and plasma levels of plasminogen activator factor inhibitor 1 (PAI-1)                                                                                                                                  | Significant | Swedish     | 1209 subjects with MI | Mellick [62] |
| rs1800629 | Cardiovascular risk factors did not differ between TNF-alpha rs1800629 high-/low-producer genotype groups                                                                                                                                 | P = 0.03 and P = 0.005, respectively | Turkish    | 102 non-diabetic patients                                      | Yilmaz et al. [124]     |
| rs1800630, rs1800629, rs1799964 | SDICH risks were positively associated with the minor alleles rs1799964 C and – rs1800629 A in men but inversely associated with rs1800630 A in females (P = 0.03)                                                              | OR = 2.635, P < 0.01 | Taiwan      | 260 SDICH patients and 368 controls                          | Chen et al. [125]       |
| rs1800629, rs361525 | Association with homocysteine levels in patients with ischemic strokes and silent brain infarctions <0.05                                                                                                                               | OR = 2.635, P < 0.01 | Koreans     | 257 patients with SBIs, and 216 control                     | Kim et al. [127]         |
| rs1800629 | A significantly increased risk of SAD was observed in the carriers of A-allele                                                                                                                                                    | OR = 2.635, P < 0.01 | Southern China | 112 patients and 121 controls                                | Yang et al. [129]        |
| rs1800630, rs1800629, rs1799964 | No significant difference in genotype distribution of rs1800629 in AD was found and no association for rs1800630 and rs1799964. A significant association between ~850 polymorphism and AD risk TT vs. TC + CC: pooled odds ratio [OR], 1.61; 1.08-2.29; P = 0.02 | Significant | Caucasian     | Meta analysis    | Di Bona et al. [130] |

Table 3 Details of polymorphism, other disease association, P-values and OR, population type and case studies with references.
| Polymorphism            | Disease associations/No associations (others)                                                                 | P-values and OR | Population    | Cases and controls (n) | References |
|------------------------|---------------------------------------------------------------------------------------------------------------|-----------------|---------------|------------------------|------------|
| Promoter SNPs          | No positive associations were found with AD                                                                 |                 |               |                        |            |
| rs1800629              | Carriers of the A-allele demonstrated better attentional processes as compared to G-allele carriers. The distribution of rs1800629 genotypes did not significantly differ from the Hardy–Weinberg equilibrium | 0.202           | Caucasian     | 67 genetically unrelated healthy participants |            |
| rs1800629              | rs1800629 A-allele was associated with worsened labile anger                                                  | <0.05           | USA           | 105 patients with hepatitis C |            |
| rs361525               | Associated with increased risk of new ICH                                                                    | 0.003           | Northern California | 280 patients |            |
| rs179964               | Variability of the rs179964 polymorphism may be associated with susceptibility to endometriosis. The frequency of the rs179964 C allele was significantly lower in stage IV endometriosis cases than controls | P = 0.04; OR = 1.75, 1.019–3.01 | Japanese     | 185 female neonates born at the Hayashi Clinic in Kobe, Japan |            |
| rs179964, rs180063,    | Alleles −1031 and −863, individually or in combination in the haplotype −1031C −863A −857C −308G 238G, were associated with lower muscle mass in men. Specifically, carriers of −1031C, compared with non-carriers, exhibited lower arm muscle mass | 0.01            | European descent | 1050. Most participants are of European descent | Liu et al. [141] |
| rs1799724, rs1800629,  | rs361525                                                      |                 |               |                        |            |
| rs361525               | No association with Paget’s disease of bone                                                                  | rs              | Spanish patients | 172 patients and 150 healthy controls |            |
| rs2229094 of LTA       | Strong association between rs2229094 and development of Proliferative Vitreoretinopathy                     | 0.0283          | Spain         | 450 patients and 312 controls | Rojas et al. [143] |
| rs1800629, rs361525,   | The rs361525 and rs909253 but not rs1800629, polymorphic variants are associated with early RM. Haplotype rs1800629 G/rs361525 A/rs909253 G show association with RM | Confirmed by regression analysis | Tunisia       | 372 RM women and 274 age-matched parous control women | Zammiti et al. [145] |

SAD, sporadic Alzheimer’s disease; ICH, intracranial haemorrhage; MS, multiple sclerosis; PBC, primary biliary cirrhosis; AD, Alzheimer’s disease; SDICH, spontaneous deep intracerebral hemorrhage; RM, recurrent miscarriage; CFTR, cystic fibrosis conductance regulator; ns, non-significant; AS, ankylosing spondylitis; IBD, inflammatory bowel disease; MI, myocardial infarction; TNF, tumor necrosis factor.
TNF-alpha, computationally, which lies in TFBS of several TFs in upstream region of TNF-alpha (Table 4). Therefore, we hypothesized that predicted SNPs interfere with gene regulation and will increase the susceptibility to disease.

Tumour necrosis factor promoter polymorphism and susceptibility to falciparum malarial infection and pulmonary tuberculosis have been carried out in Indian population. In malaria, TNF-α rs1799964 C and rs1800630 A-alleles as well as homozygotes for the TNF enhancer haplotype CACGG correlated with enhanced plasma TNF levels in both patients and controls. Significantly, higher TNF levels were observed in patients with severe malaria. In tuberculosis, no significant differences of the allele frequencies between the patients with tuberculosis and controls have been reported but a significant difference in the serum TNF-α level in the patients and the controls has been found. Two TNF polymorphisms rs1800629 and rs361525 show association in most of the diseases (if any association found). Probably, these polymorphisms affect the transcription of gene.

Polymorphisms of TNF are likely to contribute to disease, the complex pattern of associations that has been revealed could also be attributable to LD with another susceptibility locus in the vicinity of the gene. By examining LD patterns, we determined that the effect of TNF is independent of the known HLA–A and HLA–DRB1 associations (Fig. 4). The chromosomal region surrounding TNF, however, is abundant in genes of immunologic relevance. To identify true susceptibility genes, the genetic variation of the region must be studied, and extended haplotypes must be constructed and analysed.

Linkage disequilibrium patterns were estimated to determine whether the observed TNF SNP effects are attributable to the known HLA associations with disease or are independent contributors to disease. All of the associated TNF SNPs were tested against all HLA–A and HLA–DRB1 alleles. The results showed that none of the polymorphic positions in TNF are in LD with any of the associated HLA–A or HLA–DRB1 alleles (HLA–A*02, HLA–DRB1*08 or HLA–DRB1*1).

Acute anterior uveitis case–control study was carried out by Kuo et al. [159] in UK population. The association of the SNPs of TNF-α, LT-α, TNF-R1 and TNF-R2 genes in patients with idiopathic acute anterior uveitis (IAU) was investigated in this study. In addition, there

Table 4 Detected single nucleotide polymorphisms lies in transcription factor–binding site for transcription factors in tumor necrosis factor-alpha upstream region.

| SNP ID   | Position | TF     | Allele 1 | Allele 2 | Conservation |
|----------|----------|--------|----------|----------|--------------|
| rs5875327| 31648480 | NRF-2  | aGagaagaag| aGagaagaag| Yes          |
| rs4647194| 31648484 | NRF-2  | aGagaagaag| aGagaagaag| Yes          |
| rs2857713| 31648535 | Thng1-E47 | TTTGAGGTGT | TTTGAGGTGT | No           |
| rs1041981| 31648763 | AML-1  | CCTGCCACCCC | CCTGCCACCCC | No           |
| rs4647195| 31648936 | Chop-EBP | AGAGGAAGAGC | AGAGGAAGAGC | No           |
| rs3093544| 31649758 | Spz1   | AAGGAAAA | AAGGAAAA | Yes          |
| rs4645834| 31649818 | Broad-complex_4 | AGAGGAAGGGC | AGAGGAAGGGC | No           |
| rs3093547| 31649827 | Hunchback | GAaaaaaa | GAaaaaaa | Yes          |
| rs4248157| 31650122 | NF-xB  | GAGAACCTC | GAGAACTCTC | No           |
| rs9282875| 31650245 | Thng1-E47 | AGTCCTGGCAG | AGTCCTGGCAG | Yes          |
| rs6471398| 31650314 | COUP-TF | TGACCCCCCCTCCCTG | TGACCCCCCCTCCCTG | No           |
| rs2507961| 31650413 | TBP    | CTTGGGGGTCC | CTTGGGGGTCC | No           |
| rs1800630| 31650455 | NF-xB  | GGGACCCCCCA | GGGACCCCCCA | Yes          |
| rs4645836| 31650496 | p50    | GGGGGGGTCC | GGGGGGGTCC | No           |
| rs4248158| 31650512 | E74A   | CTGGAAGG | CTGGAAGG | No           |
| rs9870866| 31650536 | bZIP910 | GGGACAT | GGGACAT | No           |
| rs4248159| 31650539 | SOX17  | CAAATGGGG | CAAATGGGG | No           |
| rs2857712| 31650632 | HHF-2  | TAATGCTGGTGT | TAATGCTGGTGT | Yes          |
| rs2736196| 31650634 | HHF-2  | TAATGCTGGTGT | TAATGCTGGTGT | Yes          |
| rs2736195| 31650670 | NF-xB  | GGCCCTGCACC | GGCCCTGCACC | Yes          |
| rs4248160| 31650672 | Snail  | CAGGCT | CAGGCT | Yes          |
| rs4248162| 31650748 | Spz1   | GGGGATGAGG | GGGGATGAGG | No           |
| rs2857711| 31650749 | Spz1   | GGGGATGAGG | GGGGATGAGG | No           |
| rs1800195| 31650745 | Spz1   | GGGGATGAGG | GGGGATGAGG | No           |
| rs3093548| 31650799 | Myf    | CCCCAGCTCCTC | AAGGAGCCTGGG | No           |
| rs4248163| 31650807 | E74A   | CCCAGC | CCCAGC | No           |
| rs4248164| 31651327 | Her-1  | CAGCAGAACGTC | CAGCAGAACGTC | Yes          |
| rs4645838| 31651391 | E74A   | CTGAAAA | CTGAAAA | No           |
| rs2515924| 31651470 | CREB   | GAGGGTGACGGG | GAGGGTGACGGG | No           |
| rs2228088| 31651584 | Myf    | AAGCACCAGCCTG | AAGCACCAGCCTG | Yes          |
| rs4645839| 31651804 | ARNT   | ATctgt | ATctgt | No           |
| rs1800610| 31651806 | ARNT   | ATctgt | ATctgt | No           |
was very little linkage disequilibrium between TNF-α -857 and the other TNF SNPs, suggesting that the effect is largely attributable to TNFα-857. Results suggest that the uncommon TNF-α -857T allele is a susceptibility marker for IAU (Fig. 5).

We have checked the conservation pattern in the promoter region and found that most of the region in the promoter is conserved (Fig. 6). It is also suggestive of the fact that if a polymorphism or any variation in the DNA sequence occurs in the conserved region, then it affects the interaction of TF with TF binding largely. Understanding the conservation and change of regulatory sequences is critical to our knowledge of the unity as well as diversity of animal development and phenotypes. It can be hypothesized from these data that as the number of organisms increases, the per cent conservation decreases although certain position in the sequence remains constant throughout. These conserved sequences are thought to be the essential sites that are controlling the regulatory activity for the normal expression of the gene.

**Contribution of promoter polymorphism in expression divergence, fitness and evolution**

Wittkopp [160] reported that natural selection has played some role in expression divergence, but the relative frequency of adaptive and neutral changes remains unclear. Bradley et al. [161] observed differences in TFBS between species that were similar in regions of the genome. DNA sequence variation in TFBS affects gene expression, gene expression to phenotypic variation and phenotypic variation to fitness in the wild [161]. The variations in the DNA region alter the interaction between TF and TFBS, thereby modulating the host–parasite interaction. The genome tries to selects those variations, which provide

![Figure 4. The Linkage disequilibrium (LD) between the four SNPs mentioned in the review (rs1799964 (−1031); rs1800630 (−863); rs1800629 (−308); rs361525 (−238)). This LD plot is taken from falciparum malaria case control study in Indian population (Sinha et al. [59]).](image1)

![Figure 5. Linkage disequilibrium (LD) between TNF-1031, −863, −857, −307, and −237. Linkage disequilibrium was weak between TNF-857 and the other SNPs in the gene. The strongest evidence for LD was between TNF-857 and TNF-307 (r = 0.19). From acute anterior uveitis case control study in UK population (Kuo et al. [159]).](image2)
resistance against the disease. Malaria is a classic example of evolutionary selection, in which sickle cell anemia is selected against the pressure of malaria in endemic region. There is evidence of positive selection in early HIV-1 infection, which appears to be driven in many cases by escape from early cytotoxic T-lymphocyte (CTL) responses via mutations in the APOBEC sequence, suggesting a role for APOBEC in determining the pathway of immune escape [162].

The recruitment of different combinations of TFs to different genes allows expression of each gene to be regulated independently. Those changes that alter the activity or availability of TFs and the cis-regulatory sequences, to which they bind, can change gene expression. Both types of changes result in evolution. The studies suggested that those changes that affect the cis-regulatory activity are the predominant source of expression divergence between species [160, 163–165]. Bradley et al. [161] detected binding of the same TFs to regions of DNA in D. melanogaster and D. yakuba that have a common evolutionary origin; however, the relative affinity of these binding sites often differed between species. This suggests that evolutionary changes in the DNA sequence of cis-regulatory regions have occurred that alter the strength of the interaction between TFs and their binding sites without eliminating binding.

In the light of these facts, we have hypothesized that TNF enhancer polymorphism plays important role in susceptibility/resistance to diseases and those polymorphism that lie in transcription factor–binding sites might play role in expression divergence, fitness and evolution.

Conclusion

As the TNF gene is tightly regulated at the level of transcription. The presence of polymorphism in the 5’ regulatory region might affect transcription of TNF gene. We concluded that low-level TNF provides host defence, whereas high TNF level has been associated with severe manifestations. Alterations in the circulating levels of TNF might be a reason for differential association with diseases in different populations and also affect the expression divergence, fitness and evolution.

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