

Polymorphism of genes associated with infectious lung diseases in Northern Asian populations and in patients with community-acquired pneumonia

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Abstract: The innate immune system is the first to respond to invading pathogens. It is responsible for invader recognition, immune-cell recruitment, adaptive-immunity activation, and regulation of inflammation intensity. Previously, two single-nucleotide polymorphisms of innate-immunity genes – rs5743708 (Arg753Gln) of the *TLR2* gene and rs8177374 (Ser180Leu) of the *TIRAP* gene – have been shown to be associated with both pneumonia and tuberculosis in humans, but the data are contradictory among different ethnic groups. It has also been reported that rs10902158 at the *PKP3-SIGGIR-TMEM16J* genetic locus belongs to a haplotype race-specifically associated with tuberculosis. Meanwhile, a gradient of its frequency is observed in Asia. The aim of this work was to assess the effect of selection for the genotypes of the above-mentioned SNPs on the gene pools of populations living in harsh climatic conditions that contribute to the development of infectious lung diseases. We estimated the prevalence of these variants in white and Asian (Chukchis and Yakuts) population samples from Northern Asia and among patients with community-acquired pneumonia (CAP). Carriage of the rs5743708 A allele was found to predispose to severe CAP (odds ratio 2.77, $p = 0.021$), whereas the GG/CT genotype of rs5743708/rs8177374 proved to be protective against it (odds ratio 0.478, $p = 0.022$) in white patients. No association of rs10902158 with CAP (total or severe) was found among whites. Stratification of CAP by causative pathogen may help eliminate the current discrepancies between different studies. No significant difference in rs5743708 or rs8177374 was found between adolescent and long-lived white samples. Carriage of the alleles studied is probably not associated with predisposition to longevity among whites in Siberia. Both white and Asian populations studied were different from Western European and East Asian populations in the variants' prevalence. The frequency of the rs8177374T (Ser180Leu) variant was significantly higher in the Chukchi sample ($p = 0$, $\chi^2 = 63.22$) relative to the East Asian populations. This result may confirm the hypothesis about the selection of this allele in the course of human migration into areas with unfavorable climatic conditions. Key words: community-acquired pneumonia; pulmonary tuberculosis; genetic predisposition; genetic polymorphism; *TLR2*; *TIRAP*; *PKP3-SIGGIR-TMEM16J*; long-lived people.

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Полиморфизм генов, ассоциированных с инфекционными заболеваниями легких, в популяциях Северной Азии и среди пациентов с внебольничными пневмониями

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Аннотация: Врожденный иммунитет первым отвечает на инфекцию. Он участвует в распознавании патогена, привлечении к месту заражения иммунных клеток, а также активирует адаптивный иммунитет и регулирует интенсивность воспалительного ответа. Для двух однонуклеотидных полиморфизмов (ОНП) генов врожденного иммунитета – rs5743708 (Arg753Gln) гена *TLR2* и rs8177374 (Ser180Leu) гена *TIRAP* – была показана ассоциация одновременно с пневмонией и туберкулезом, однако полученные данные различаются для разных этнических групп. Для rs10902158, расположенного в генетическом локусе *PKP3-SIGGIR-TMEM16J*, ранее было показано, что он входит в гаплотип, распецифически ассоциированный с туберкулезом. При этом на тер-

ритории Азии наблюдается градиент его частоты. Целью нашей работы была оценка влияния отбора по генотипам названных ОНП на генофонды популяций, живущих в климатических условиях, неблагоприятных по инфекционным заболеваниям легких. Мы оценили распространение этих ОНП в выборках европеоидного и монголоидного (чукчи и якуты) населения Северной Азии и среди пациентов с внебольничной пневмонией. Носительство аллеля А rs5743708 гена *TLR2* предрасполагало к развитию тяжелой внебольничной пневмонии ($OR = 2.77, p = 0.021$), а генотип GG/CT по rs5743708/rs8177374 оказался протективным против нее ($OR = 0.478, p = 0.022$) у европеоидов. Ассоциации rs10902158 с внебольничной пневмонией (как в целом, так и ее тяжелой формы) в европеоидной выборке обнаружено не было. Дифференцировка внебольничных пневмоний по их этиологии, возможно, позволит устранить наблюдаемые противоречия в данных разных исследователей об ассоциации исследованных ОНП с заболеванием. Не выявлено достоверных различий по частоте rs5743708 гена *TLR2* и rs8177374 гена *TIRAP* между европеоидными выборками подростков и долгожителей. Вероятно, эти ОНП не влияют на предрасположенность к долгожительству в европеоидных популяциях, проживающих на Севере Евразии. Исследованные нами европеоидные и монголоидные популяционные выборки отличались по частотам вышеперечисленных вариантов от западноевропейских и восточноазиатских популяций. Частота варианта rs8177374 T (Ser180Leu) гена *TIRAP* в выборке чукчей была достоверно выше ($p = 0, \chi^2 = 63.22$), чем в популяциях Восточной Азии, что может служить подтверждением нашего предположения об отборе этого варианта в ходе миграции человека в районы с неблагоприятными климатическими условиями. Ключевые слова: внебольничная пневмония; туберкулез легких; генетическая предрасположенность; генетический полиморфизм; ген *TLR2*; ген *TIRAP*; генетический район PKP3-SIGGIR-TMEM16J; долгожители.

Introduction

Innate immunity constitutes the first barrier against microorganisms and viruses by destroying infected cells and activating adaptive immunity. Nonetheless, an excessive nonspecific immune reaction (inflammation) may be life threatening because it can completely disrupt the functioning of vital organs. Community-acquired pneumonia (CAP) and pulmonary tuberculosis (PTB) are infectious diseases characterized by high mortality, and according to WHO, are ranked consistently among the top 10 leading causes of death in the world (<https://www.who.int/en/news-room/fact-sheets/detail/the-top-10-causes-of-death>).

Pneumonia is an inflammatory lower-respiratory-tract disease caused by viruses, bacteria, fungi, and parasites. In addition, it may be due to noninfectious processes or have a combined cause. For a long time, *Streptococcus pneumoniae* infection has been considered the main cause of CAP; however, it was shown recently that CAP develops mainly as a result of viral infections (influenza A and B viruses, parainfluenza viruses, adenovirus, respiratory syncytial virus, or coronaviruses) (Choi et al., 2012; Hong et al., 2014; Self et al., 2017). *Streptococcus pneumoniae*, *Haemophilus influenzae*, *Staphylococcus aureus*, *Mycoplasma pneumoniae*, *Chlamydia pneumoniae*, *Legionella pneumophila* and other microbes may be causative agents of bacterial pneumonia (Choi et al., 2012; Hong et al., 2014; Self et al., 2017). After invasion of the airway epithelium by pathogens, these cells start to produce reactive oxygen species, cytokines, and other mediators to recruit immune cells. Being most abundant in lungs, alveolar macrophages ingest bacteria and apoptotic cells and can present antigens on MHC II to other immune cells. Proinflammatory M1 macrophages produce cytokines TNF α , IL-6, IL-1 β , IL-12, and IL-23 to enhance inflammation for elimination of the invaders. Anti-inflammatory M2 macrophages produce cytokines IL-4, IL-13, and TGF- β to induce completion of the inflammatory reaction and remodeling of damaged tissue (Moldoveanu et al., 2009; Arango Duque, Descoteaux, 2014; Kumar, 2019).

Depending on the set of present chemokines and cytokines, different cells responsible for humoral and cellular immunity are attracted to the site of infection (Kumar, 2019). Severe

pneumonias are more likely to develop in coinfections; it has been demonstrated that a viral infection (in particular influenza) facilitates the development of pneumococcal infection by damaging the epithelium and reducing the amount of a surfactant (McCullers, 2014; Aguilera, Lenz, 2020). Human respiratory syncytial virus, metapneumovirus, adenovirus, and influenza viruses A and B prefer a cold season (Price et al., 2019). The seasonal increase in the incidence of pneumococcal pneumonia coincides with seasonal outbreaks of influenza; *S. pneumoniae*, *H. influenzae*, and *S. aureus* infections have been reported to be associated with significant influenza pandemics (McCullers, 2014; Bystritskaya, Bilichenko, 2017; Morris et al., 2017).

PTB is a pulmonary infectious disease caused mainly by *Mycobacterium tuberculosis* (*Mtb*). According to the WHO, approximately one-quarter of the world's population is estimated to be infected by *Mtb*, and 5–15 % of these people will fall ill with active tuberculosis. In Russia, most of these patients (95 %) have PTB (<https://minzdrav.gov.ru/ministry/61/22/stranitsa-979/statisticheskie-i-informatsionnye-materialy/statisticheskiy-sbornik-2018-god>). The pathogenesis of pulmonary tuberculosis is based on *Mtb* survival after phagocytosis by alveolar macrophages. These bacteria can modulate a host immune response to protect the infected cells, change their metabolism, induce IL-10, suppress IL-12 and TNF α synthesis, and to inhibit MHC II expression and antigen presentation. *Mtb* makes macrophages unresponsive to interferon (IFN) γ and inhibits autophagy. It allows the mycobacteria to establish a persistent or latent infection in macrophages. Mycobacteria are believed to use the general mechanism of negative feedback regulation that restricts excessive inflammation (Harding, Boom, 2010; Richardson et al., 2015; Gopalakrishnan, Salgame, 2016). With the loss of immunity-driven control over mycobacterial reproduction, foamy macrophages accumulate in granulomas, and lung tissue necrosis begins (Liu C.H. et al., 2017). Vitamin D deficiency is known to negatively affect the effectiveness of the immune response in tuberculosis (Wilkinson et al., 2000; Aibana et al., 2019).

These data suggest that in Northern Asia, a region with low temperature and reduced insolation during most of the year,

signs of purifying selection for genes associated with lung infections may be noticeable. For many genes of innate immunity, an association with viral, bacterial, and autoimmune diseases has been proven. Despite differences in the pathogenesis between CAP and PTB, it has been shown that minor alleles of rs5743708 (the *TLR2* gene) and of rs8177374 (the *TIRAP* gene) can have a pathogenic and protective effect, respectively, in both of these lung diseases in humans. Nevertheless, data obtained by different research groups are contradictory.

Toll-like receptors (TLRs) play a pivotal role in host defense. Being membrane-anchored (TLRs 1, 2, 4–6, and 10) or endosomal (TLRs 3 and 7–9) in human immune cells (e.g., macrophages, monocytes, dendritic cells, and some leukocytes), they are involved in the recognition of structurally conserved surface molecules of microorganisms and viruses as well as viral nucleic acids (Barbalat et al., 2009; Kawai, Akira, 2010; Kumar, 2019). TLR2 participates in the recognition of a large number of diverse lipoproteins and peptidoglycans of gram-positive and gram-negative bacteria, fungi, and virus-infected cells. After ligand binding to the receptor, TIR (Toll-interleukin 1 receptor) domains of TLR2 and TLR1 or TLR2 and TLR6 dimerize via the formation of an extensive hydrogen-bonding network and hydrophobic interactions (Jin et al., 2007; Takeda, Akira, 2015). Homodimerization of the cytoplasmic domains of TLR2 does not induce TNF α production *in vitro* in murine macrophages, and the formation of the TLR2–TLR2 dimer is not detectable even in the presence of an agonist (Ozinsky et al., 2000; Shukla et al., 2018). Therefore, the existence of TLR2–TLR2 homodimers *in vivo* is being questioned. After ligand binding, reorientation of the TIR domains and triggering of a cascade of intracellular reactions lead to the activation of proinflammatory NF- κ B and MAPK pathways, synthesis and a release of proinflammatory cytokines (IL-1 β , IL-12, TNF- α , and IL-6) and various chemokines into extracellular space, and the development of an inflammatory response at the pathogen entry site (Liu C.H. et al., 2017; Tapader et al., 2018). In inflammatory monocytes, TLR2 induces type I IFN production in response to a viral ligand (Barbalat et al., 2009). It is reported that prolonged stimulation of TLR2 (more than 24 h) causes PI3K/Akt pathway activation in alveolar macrophages. It limits the production of NF- κ B, TNF- α , and IL-12 and activates the synthesis of anti-inflammatory IL-10. This mechanism is assumed to prevent excessive inflammation (Richardson et al., 2015; Liu Y. et al., 2016).

The *TLR2* gene is located in 4q31.3, has five exons, and expresses few splicing isoforms, but all of the coding sequences are contained within exon 3. The protein consists of 784 amino acid residues (aa) and includes extracellular leucine-rich repeat domains, which are primarily responsible for ligand recognition (aa 54–524), followed by the leucine-rich repeat C-terminal domain (aa 525–579) and intracellular TIR domain (aa 639–782), which mediates downstream signaling (<https://www.uniprot.org/uniprot/O60603>). It is expressed constitutively on macrophages and dendritic cells and can be induced in epithelial cells or B-cells. Its overexpression in patients with pneumococcal disease had been documented (Siebert et al., 2018).

Single-nucleotide polymorphism (SNP) rs5743708 (of the *TLR2* gene) causing the Arg753Gln substitution is located

in the TIR domain of the protein. This SNP is associated simultaneously with resistance to Lyme disease (Schröder et al., 2005) and with predisposition to tuberculosis (Guo, Xia, 2015; Patarčić et al., 2015), whereas the association with predisposition to PTB is race-specific (Caws et al., 2008; Guo, Xia, 2015; Hu et al., 2019). It is believed that TLR2 signaling may be nonessential to control acute tuberculosis but important during chronic tuberculosis (Gopalakrishnan, Salgame, 2016). A meta-analysis has shown the *TLR2* rs5743708 minor allele to be associated with CAP, Legionnaires' disease, and pneumococcal disease; however, the data obtained in different studies are contradictory (Moens et al., 2007; Patarčić et al., 2015; Smelaya et al., 2016).

The *TIRAP* (TIR domain-containing adaptor protein) gene also known as *Mal* (MyD88 adapter-like) encodes one of the five adapter proteins that are involved in signal transduction from activated TLRs to protein kinases at the plasma membrane (Bonham et al., 2014). It is located in 11q24.2, consists of six exons, and encodes a protein of 221 aa. The TIRAP protein includes an N-terminal PEST domain (aa 15–35) responsible for binding to special sites in the plasma membrane, followed by an AB-loop mediating MyD88 and TLR4 binding. A binding site for TRAF6 (TNF receptor-associated factor 6) is located within the region aa 188–193 (Bernard, O'Neill, 2013). TIRAP is expressed in many cell types (Narayanan, Park, 2015), and its isoforms resulting from alternative splicing have unknown functions. There are different opinions about whether TIRAP forms a complex with the TIR domain of TLR6 for signal transmission; however, it has been proven that TIRAP mediates TLR2 and TLR4 signaling by facilitating the recruitment of the MyD88 adaptor protein to the TLRs (Nagpal et al., 2009; Bernard, O'Neill, 2013).

Activation of NF- κ B, MAPK1, MAPK3, and JNK results in cytokine secretion and an inflammatory response. SNP rs8177374 (the *TIRAP* gene) is located in exon 5 and represents the Ser180Leu substitution in the encoded protein. It is located close to the TLR-binding site of TIRAP. In carriers of this substitution, modulation of TLR1, TLR2, TLR4, and TLR6 but not TLR9 signaling has been shown (Khor et al., 2007; Ferwerda et al., 2009; Siebert et al., 2018). Ser180Leu in a heterozygous state has a protective effect against PTB and invasive pneumococcal disease in white and African samples and against malaria in African and Asian samples (Khor et al., 2007; Panda et al., 2016). Carriage of heterozygous Ser180Leu protects children from pneumococcal lower-respiratory-tract infections, whereas carriers of the homozygous 180Leu polymorphism alone or in combination with some TLR1 and TLR6 polymorphisms may be susceptible to recurrent pneumococcal infections (Siebert et al., 2018). Simultaneous carriage of the TIRAP 180Leu variant and some SNPs in the *TLR4* gene as well as 180Leu homozygosity increases susceptibility to severe hospital-acquired infections (Kumpf et al., 2010).

The opposite results have been obtained as well. The rs8177374 T allele (180Leu) increases the risk of PTB in a sample of Iranian population (Naderi et al., 2014). A meta-analysis of nine published case-control studies did not reveal a significant association of 180L with tuberculosis risk (Miao et al., 2011). There are controversial opinions about the mechanism behind the observed protective effect of Ser180Leu heterozygosity. They are based on differences in observed

effects of the SNP at the level of proinflammatory cytokines. Depending on the model used, some research groups showed an increased level (Ferwerda et al., 2009; Panda et al., 2016) and others a decreased level (Khor et al., 2007; Kumpf et al., 2010; Siebert et al., 2018) of cytokines after their induction in 180 Leu/Leu carriers. Accordingly, homozygosity of the minor variant of rs8177374 is thought to cause either an excessive inflammatory reaction or the absence of an adequate immune response. It is supposed that selection pressure on the *TIRAP* gene provides a balance between protection against excessive inflammation and effective defense during infectious diseases (Khor et al., 2007; Ferwerda et al., 2009).

Besides the polymorphisms in genes *TLR2* and *TIRAP*, in this paper, we focused on the *PKP3-SIGGIR-TMEM16J* gene region. An association of its haplotypes with different types of tuberculosis has been shown among children in Vietnam and South Africa (Horne et al., 2012; Gupta et al., 2016). It is believed that the impact of the haplotypes on immunity is determined by SIGIRR (single immunoglobulin interleukin 1 receptor; synonym: IL-1R8), which is a negative regulator of TLR signaling (Molgora et al., 2016). Carriage of rs10902158 GG and rs7111432 AA in introns of *PKP3* and *TMEM16J*, respectively, acts additively with a vitamin D deficiency and “pathogenic” genotypes of rs5743708 (*TLR2*) and rs8177374 (*TIRAP*) on tuberculosis predisposition (Horne et al., 2012; Gupta et al., 2016). rs10902158 located in intron 2 of the *PKP3* gene has been analyzed. Encoded desmosomal plaque protein plakophilin 3 is involved in intracellular adhesion (Gurjar et al., 2018). Of note, rs10902158 has a frequency gradient in Asia; according to the Genome Aggregation Database (GnomAD) (<https://gnomad.broadinstitute.org/>), it is absent in South Asia and is found with a frequency of ~50 % in Southeast Asia. Nonetheless, functional significance of genetic variants in noncoding parts of the *PKP3-SIGGIR-TMEM16J* gene region, including rs10902158, is not clear.

In this work, we analyzed the frequencies of rs5743708, rs8177374, and rs10902158 (for which conflicting data on the association with respiratory infections have been reported previously) in white and Asian samples from Northern Asia and among CAP patients. According to the statistics of the Ministry of Health of the Russian Federation, Novosibirsk Oblast and Yakutia are characterized by an increased incidence of PTB, whereas Chukotka Autonomous Okrug is the leader in both pneumonia and PTB morbidity in Russia (<https://minzdrav.gov.ru/ministry/61/22/stranitsa-979/statisticheskii-i-informatsionnye-materialy/statisticheskii-sbornik-2017-god>; <https://minzdrav.gov.ru/ministry/61/22/stranitsa-979/statisticheskii-i-informatsionnye-materialy/statisticheskii-sbornik-2018-god>). According to the WHO, the highest death rate from pneumonia is observed before the age of 5 and after 75–80 years (https://www.who.int/medicines/areas/priority_medicines/Ch6_22pneumo.pdf). Therefore, we assumed that long-lived people of the Siberian Federal District may differ from adolescents in the frequency of rs5743708 and rs8177374, and we assessed the prevalence of the pathogenic variants in the sample of long-lived people.

Materials and methods

The study protocol was approved by the local Ethics Committee of the Institute of Internal and Preventive Medicine (branch

of the Institute of Cytology and Genetics of the Siberian Branch of the Russian Academy of Sciences, Novosibirsk, Russia; approval No. 22.06.2008). Written informed consent to be examined and to participate in the study was obtained from each patient. For individuals younger than 18 years, the informed consent was signed by a parent or legal guardian.

The white sample consisted of 451 adolescents (95 % Russians, 197 males, 253 females, aged 14–17) from Oktiabr'skii district of Novosibirsk (55°01' N 82°55' E) and 289 Russian settlers (120 males, 169 females, aged 45–64, mean age 52.5) in towns Tommot (58°58'00" N 126°16'0" E), Neryungri (56°39'30" N 124°43'30" E), Ust-Nera (64°34'05" N 143°14'10" E), and Yakutsk (62°01'38" N 129°43'55" E), who lived in Yakutia for more than 16 years or were born there. The adolescent sample was described previously (Zavyalova et al., 2011). Asian samples (220 individuals) consisted of 130 Chukchis (66 males, 64 females, aged 18–73, mean age 40) from the Kanchalan village (65°10'41" N 176°44'52" E) of Chukotka Autonomous Okrug and 132 Yakuts (53 males, 79 females, aged 44–64, mean age 49) from the Kylay village (63°13'34" N 132°08'06" E) and towns Tommot, Neryungri, and Ust-Nera of Yakutia. The sample of long-lived people was collected in cities Novosibirsk, Tomsk, and Tumen and consisted of 188 individuals (180 females, 8 males) aged 90–105, mean age 92.

Ethnicity of individuals was identified using questionnaires and additional cross-examination with elucidation of the nationality of ancestors (at least in three generations). Persons of mixed origin were excluded from the analysis. The CAP patient sample (406 whites) was collected in offices of pulmonary hospitals of Novosibirsk and Yakutsk in 2003–2005 before the COVID-19 outbreak. The sample consists of 120 patients with severe CAP (aged 18–80, mean age 52) and 286 patients with nonsevere (mild to moderate severity) CAP (aged 16–92, mean age 39). The diagnosis of pneumonia was made on the basis of radiologically confirmed “fresh” lung tissue infiltration and clinical data (fever, cough, sputum production, chest pain, and shortness of breath) in the absence of an obvious diagnostic alternative. CAP of various etiologies was regarded as severe if the CURB65 rating scale index was 4–5 points (Lim et al., 2003).

Genomic DNA was extracted from peripheral blood leucocytes by the standard phenol–chloroform method (Sambrook, Russell, 2006). Genotyping was performed using polymerase chain reaction (PCR) with an analysis of restriction fragment length polymorphism by electrophoresis in a 5 % polyacrylamide gel after visualization with an ethidium bromide solution.

The rs5743708 SNP (*TLR2*) was identified by amplification of DNA with primers 5'-GCCATTCTCATTCTTCTGG*AGC-3' and 5'-GGGAACCTAGGACTTTATCGCA-3' (* denotes a nucleotide changed for restriction site creation). The 168-bp PCR product was digested with the Pst I restriction endonuclease (SibEnzyme, Novosibirsk) for 2 h at 37 °C. The rs5743708 A (753Q) allele was revealed by the presence of fragments of 20, 45, and 103 bp, whereas the rs5743708 G allele by fragments of 20 and 148 bp.

The detection of rs8177374 (*TIRAP*) was performed by amplification of genomic DNA with primers 5'-GGCTGC ACCATCCCCCA*GC-3' and 5'-CCGTTCCCCTTCTCCCT

CCTGTAG-3' (* denotes a nucleotide changed for restriction site creation). The 162-bp PCR product was digested with the AccB7 I restriction endonuclease (SibEnzyme, Novosibirsk) for 2 h at 37 °C. The rs8177374 T (180L) allele was identified by the presence of fragments of 21 and 141 bp, whereas in case of rs8177374 C, the PCR product was not cut.

Primers 5'-TGGCAAGGATTGGAGAAGCTC*C*TGTC-3' and 5'-CAGGGCCAGTGCCTCCCC-3' (* denotes nucleotides changed for restriction site creation) were used for the amplification of the *PKP3* intron 2 sequence containing rs10902158. The resulting 192-bp amplicon was digested with the BstEN I restriction endonuclease (Sibenzyme, Novosibirsk) for 2 h at 65 °C. In the presence of the rs10902158 A allele, the PCR product was not cut, whereas in the case of the rs10902158 G allele, fragments of 24 and 168 bp were observed.

Statistical analysis was performed in the SPSS 16.0 software.

Results

Genotype distributions were consistent with the Hardy–Weinberg equilibrium among all the population samples (data not shown). Minor allele frequencies for rs5743708, rs8177374, and rs10902158 are represented in Table 1.

In the sample of Novosibirsk adolescents, allele frequencies of rs5743708, rs8177374, and rs10902158 differed from those of the non-Finnish European sample in GnomAD ($\chi^2 = 6.621$, $p = 0.013$; $\chi^2 = 19.541$, $p = 0$; $\chi^2 = 54.554$, $p = 0$, respectively).

For the frequency of the rs5743708 A (Arg753Gln) allele, there was a tendency for a decrease in Russian settlers in Yakutia and among long-lived people compared with the Novosibirsk sample.

The rs8177374 T (Ser180Leu) frequency did not differ among the studied white samples.

In the sample of Russian settlers of Yakutia, males and females differed in the frequency of rs10902158 ($p = 0.037$, $\chi^2 = 4.86$). Moreover, the frequency of this SNP among males was closer to that observed in non-Finnish Europeans according to GnomAD data, and the frequency among females was closer to that observed in the Novosibirsk sample. Perhaps there were sex differences during recent migration to Yakutia from different regions of Russia. Genetic analysis of a larger sample and estimation of this SNP's frequency in western regions of Russia are required for explaining the observed differences.

The two analyzed Asian samples differed from each other and from GnomAD East Asian cohorts. Among Yakuts, the rs5743708 A (Arg753Gln) variant, which is very rare in other Asian populations, was found at a frequency of 0.015 ± 0.007 (mean \pm SD). Chukchis differed significantly from GnomAD East Asians in rs10902158 allele frequency ($p = 0$, $\chi^2 = 63.22$) (see Table 1).

Frequencies of polymorphisms rs5743708, rs8177374, and rs10902158 were not different among adolescents and total white CAP patient samples. By contrast, after the sample was divided into patients with severe and nonsevere CAP, differences were found for rs5743708 ($p = 0.021$, $\chi^2 = 6.24$). Next, genotype frequencies were estimated for rs5743708, rs8177374, and rs10902158 in the examined samples (except for long-lived people regarding rs10902158). For the latter SNP (in the *PKP3* gene), no difference in frequency was detectable within any group (data not shown). The observed distribution of rs5743708 and rs8177374 genotypes among the studied samples is presented in Table 2.

Table 1. Minor allele frequencies for rs5743708, rs8177374, and rs10902158 in the studied samples and in the Genome Aggregation Database (GnomAD)

Group, sample size	rs5743708 A (<i>TLR2</i>)		rs8177374 T (<i>TIRAP</i>)		rs10902158 A (<i>PKP3</i>)	
	Allele number	Allele frequency	Allele number	Allele frequency	Allele number	Allele frequency
Asians						
Chukchis, 130	0	0	17	0.065	116	0.446
Yakuts, 132	4	0.015	6	0.023	100	0.379
East Asia (GnomAD)		0.0002		0.012		0.562
Whites						
Adolescents (Novosibirsk), 451	39	0.043	93	0.103	198	0.220
Long-lived people, 188	11	0.029	29	0.077	–	–
Russians (Yakutia) total sample, 289	13	0.022	50	0.087	119	0.206
Russians (Yakutia), females, 169	7	0.020	29	0.086	80	0.237
Russians (Yakutia), males, 120	6	0.025	21	0.088	39	0.163
Total CAP, 406	25	0.031	80	0.100	200	0.246
Severe CAP, 120	13	0.054	17	0.071	62	0.258
Nonsevere CAP, 286	12	0.021	63	0.109	138	0.241
Non-Finnish European (GnomAD)		0.029		0.157		0.153

Note: CAP, community-acquired pneumonia.

Table 2. Combined genotype frequencies of rs5743708 and rs8177374 in the studied populations and patient groups

Samples/genotypes	Sample size	GG/CC		GG/CT		GG/TT		AG/CC		AG/CT		AA/CC		AA/CT	
		<i>n</i>	<i>q</i>												
Adolescents, Novosibirsk	451	335	0.742	75	0.166	4	0.009	27	0.060	8	0.018	1	0.002	1	0.002
Russians, Yakutia	289	230	0.796	44	0.152	2	0.007	10	0.035	3	0.010	0	0	0	0
Long-lived people	188	150	0.798	25	0.133	2	0.011	11	0.058	0	0	0	0	0	0
Total CAP	406	308	0.759	71	0.175	2	0.005	22	0.054	3	0.007	0	0	0	0
Severe CAP	120	93	0.775	13	0.108	1	0.008	11	0.092	2	0.017	0	0	0	0
Nonsevere CAP	286	215	0.752	58	0.203	1	0.003	11	0.038	1	0.003	0	0	0	0
Chukchis	130	113	0.869	17	0.131	0	0	0	0	0	0	0	0	0	0
Yakuts	132	122	0.924	6	0.045	0	0	4	0.030	0	0	0	0	0	0

Note: *n*, number of genotype carriers; *q*, genotype frequency.

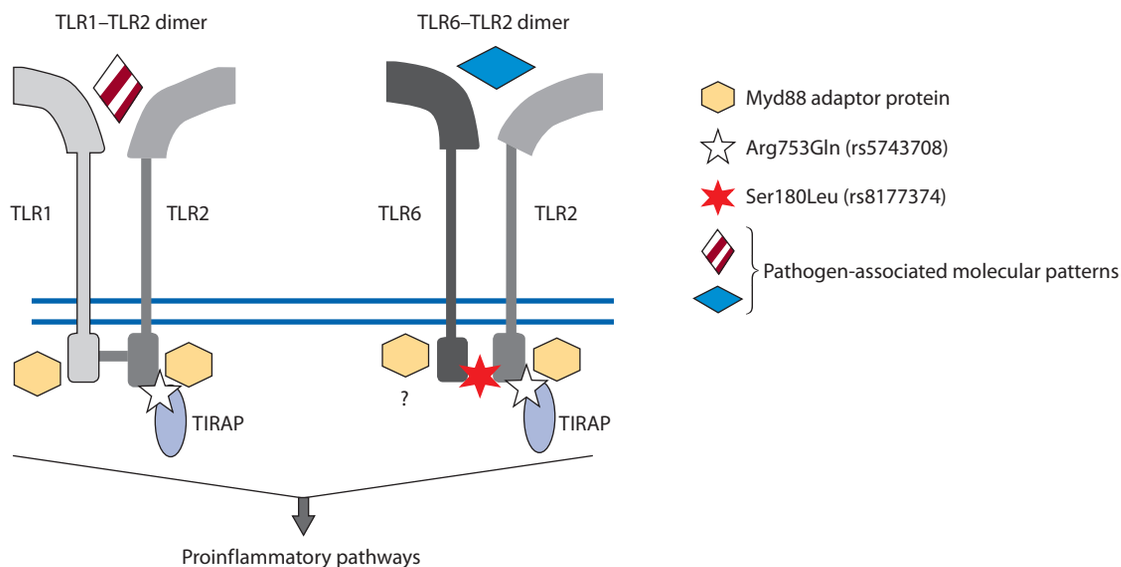
The frequencies of genotypes of rs5743708 and rs8177374 did not differ among the following samples: adolescents of Novosibirsk, long-lived people of Siberia, all patients with CAP, and all Russians in Yakutia. Possibly, the carriage of the studied alleles is not associated with predisposition to longevity in Siberia and does not significantly affect the probability of resettling of migrants in more unfavorable climatic conditions at present. This notion is consistent with WHO findings that in Eastern Europe, in contrast to Western and Central Europe, the pneumonia mortality rate does not increase significantly after age 80 (https://www.who.int/medicines/areas/priority_medicines/Ch6_22Pneumo.pdf). As for pneumonia, carriage of the rs5743708 A allele predisposed to severe CAP (AG/CC+AG/CT vs all: odds ratio 2.77, 95 % confidence interval 1.227–6.272, $p = 0.021$). The heterozygous genotype of rs8177374 in combination with the GG genotype of rs5743708 had a protective effect against severe CAP (GG/CT vs all: odds ratio 0.478, 95 % confidence interval 0.251–0.909, $p = 0.022$).

Discussion

It was shown here that carriage of none of the three studied SNPs, rs5743708, rs8177374, and rs10902158, is associated with the predisposition to CAP in total. By contrast, we found that the Arg753Gln variant of *TLR2* predisposes to severe CAP, and the heterozygous Ser180Leu variant of *TIRAP* in combination with the 753 Arg/Arg variant of *TLR2* has a protective effect against it in the white population. These data partially explain the contradictions in the data from different researchers. Most likely, the contribution of the alleles of genes *TLR2* and *TIRAP* to CAP predisposition is determined by pneumonia etiology. A substantial proportion of severe pneumonia cases are known to be caused by combined viral and bacterial infections (McCullers, 2014; Morris et al., 2017; Aguilera, Lenz, 2020). *TLR2* is responsible mainly for the recognition of bacteria-associated molecular patterns; Ser180Leu of the *TIRAP* gene modulates signal transduction only from *TLR2* and *TLR4* recognizing molecular patterns of bacteria as well (Nagpal et al., 2009). Most likely, combined and bacterial but not viral pneumonias are associated with *TLR2* and *TIRAP* gene variants.

The studied Asian ethno-geographical groups showed an increased frequency of the protective rs8177374 T (Ser180Leu) variant of *TIRAP* relative to neighboring East Asian populations. In the Chukchi sample, the difference was significant ($p = 0$, $\chi^2 = 63.22$). It may be a consequence of the natural selection that has promoted protection from excessive inflammation during pulmonary diseases. The hypothesis about the selection of the Ser180Leu variant along with the out-of-Africa migration to a harsh environment has been advanced earlier (Khor et al., 2007; Ferwerda et al., 2009). An increased frequency of the Arg753Gln variant of *TLR2* and a decreased frequency of Ser180Leu of *TIRAP* as compared to non-Finnish Europeans may indicate higher genetic predisposition of the Siberian white population to PTB and severe CAP. Nevertheless, there are a lot of genes associated with CAP and PTB independently. Apparently, during the settlement of peoples in Northern Eurasia, the formation of gene pools had been determined by the selection that facilitated adaptation to specific infections (Lime disease among them), parasites, and the climate. It would be interesting to determine why two mutations changing the same *TLR2* signaling have opposite effects on the predisposition to severe CAP. One possible explanation is the difference in the structure and functions of heterodimers *TLR2*–*TLR1* and *TLR2*–*TLR6* (see the Figure).

Existing data on the roles of *TLRs* 1, 2, and 6 in the activation of proinflammatory and anti-inflammatory signaling are conflicting. Overexpression of *TLR2* carrying the Arg753Gln variant has been demonstrated to cause a significantly stronger impairment of cytokine induction by *TLR2*/*TLR1* ligands as compared with *TLR2*/*TLR6* ligands in the HEK293 cell line (Schroder et al., 2005). In later papers, it has been shown that the Arg-to-Gln substitution at position 753 of *TLR2* changes the size, charge, and hydrophobic properties of this site and reduces the ability of *TLR2* to form a heterodimer with *TLR6* (Basith et al., 2011; Xiong et al., 2012). This SNP significantly alters agonist-inducible association of *TLR2* with adaptor proteins *TIRAP* and *MyD88* and impairs NF- κ B signaling and IL-8 mRNA expression in the HEK293 cell line (Xiong et al., 2012). Genes *TLR1* and *TLR2* have different expression activators (Lancioni et al., 2011). In the *TLR2*–*TLR1* dimer, *TLR1* and *TLR2* are responsible for NF- κ B and MAPK pathways



Locations of the Arg753Gln substitution in TLR2 and Ser180Leu in TIRAP on the protein complexes.

and for PI3K pathway activation, respectively; therefore, upregulation of proinflammatory cytokines is TLR1-dependent, whereas upregulation of type I IFN is TLR2-dependent (Raieli et al., 2019). TLR2–TLR6 binding disruption possibly causes an increase in the number of TLR2–TLR1 dimers in which TLR1 drives the activation of the NF- κ B inflammatory signaling cascade. On the contrary, the Ser180Leu variant of TIRAP weakens proinflammatory signal transmission. Besides, TIRAP acts as an adaptor protein for TLR4 homodimers. This receptor is primarily responsible for the recognition of lipopolysaccharides of gram-negative bacteria and fungi (Takeda, Akira, 2015). It is believed that TLR4 takes part not only in MyD88-dependent proinflammatory signaling but also in MyD88-independent anti-inflammatory signaling. Weakening of TLR4 signaling through TIRAP probably enhances the signaling through adapter proteins TRIF and TRAM, causing the secretion of anti-inflammatory cytokines (Li et al., 2013).

It remains unclear why the SNPs in *TLR2* and *TIRAP* have similar effects on the predisposition to or protection against acute (CAP) and chronic (PTB) lung infections. Perhaps this phenomenon is due to an impact on inflammation in both diseases. The severity of CAP is determined by life-threatening acute inflammation; in PTB, the development of chronic inflammation masks the infection from the host immune system (Liu C.H. et al., 2017).

Conclusions

In Northern Asian populations, the observed difference in rs8177374 frequency may reflect consequences of natural selection during the settlement of peoples on territories with unfavorable climatic conditions. As for pneumonia, carriage of the rs5743708 A allele (the *TLR2* gene) predisposes to severe CAP; the heterozygous genotype of rs8177374 (the *TIRAP* gene) in combination with the GG genotype of rs5743708 (the *TLR2* gene) has a protective effect against it. Stratification of CAP by causative pathogen may help to eliminate the current discrepancies among research groups. Regional differences in a set of pathogens along with the genetic characteristics of

ethno-geographical groups can determine the associations of various genetic variants of innate immunity with the prevalence and severity of pneumonia.

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