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MOLECULAR BIOLOGY APPLICATIONS OF THE RED KING CRAB DUPLEX-SPECIFIC NUCLEASE

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Duplex-specific nuclease (DSN) from hepatopancreas of the craboid Paralithodes camtschaticus (red king crab) has a unique combination of properties. Along with thermal stability and a high optimal temperature of catalysis, this enzyme exhibits high substrate selectivity, cleaving only DNA in duplexes (DNA-DNA or DNA-RNA). Accordingly, it digests neither single strands (nor single-stranded regions) of DNA, nor RNA strands with any secondary structure. Such properties make it possible to create unique protocols based on DSN, which is also an important object of fundamental research in the field of nuclease evolution. The review considers diverse applications of the red king crab DSN in modern methods of molecular biology.

Keywords: duplex-specific nuclease, DSN, crab hepatopancreas, Paralithodes camtschaticus, red king crab

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Author contribution: DA Shagin — preparation of the manuscript; DV Rebrikov — editing of the manuscript.

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This review considers diverse applications of DSN in modern molecular biology.

Single nucleotide polymorphisms (SNP) genotyping

SNP genotyping is used in the diagnosis of genetic predispositions, pharmacogenetics, forensic science, molecular genealogy, population genetics, and other research areas [5–9]. An SNP genotyping protocol known as duplex-specific nuclease preference (DSNP) approach is based on the unique property of DSN to cleave perfect (i.e. fully matched) short double-stranded DNA substrates with much higher efficiency than their imperfect analogs [10].

SNP genotyping by DSNP-analysis requires two specific 10-mer oligonucleotides with a fluorophore at the 5'-end and a quencher at the 3'-end — the so-called FRET-labeled probes (FRET, fluorescence resonance energy transfer). One of the probes corresponds to wild-type allele, the other one corresponds to variant allele. In the case of perfect duplex formed

Crab duplex-specific nuclease (DSN) from hepatopancreas of the red king crab was firstly characterized in 2002 [1]. The enzyme with a molecular weight of 41.5 kDa consists of 407 amino acid residues (Genbank AAN86143) and exerts a unique set of functional properties [2, 3]:

- DSN exhibits maximum activity at pH 6.6 at 60–65 °C; DSN remains active after heating at 90°C or incubation at pH within the range of 4–12;
- DSN is Mn²⁺, Co²⁺ and Mg²⁺ dependent;
- DSN is resistant to proteases (including proteinase K and papain);
- DSN cleaves only double-stranded DNA, leaving single strands intact;
- DSN shows negligible activity towards RNA of any secondary structure, while effectively cleaving the DNA chain in DNA-RNA hybrids.

The unique properties of DSN, which is also an important object of fundamental research in the field of nuclease evolution [4], inspires creation of molecular protocols on their basis.
between the probe and the target DNA, the probe is hydrolyzed by DSN, which results in the fluorophore-quencher uncoupling and emission of fluorescence at a specific wavelength. In the absence of probe hydrolysis, no fluorescence is emitted.

Before DSNP analysis, the studied polymorphic region must be amplified to a high concentration by polymerase chain reaction (PCR) with specific primers. The crude (unpurified) PCR product is mixed with the probes and incubated with DSN at 60°C for 5–10 min. During the incubation, the DNA substrate becomes amplified due to combined activity of DSN and thermostable DNA polymerase; the latter is introduced into the reaction mixture as a component of the crude PCR product used as a template. DSN cleaves double-stranded DNA producing fragments that can serve as primers for DNA polymerase. At the same time, due to the hydrolysis of amplicons, short DNA fragments are formed that are capable of efficient hybridization with the signaling probes. At the final step of the analysis, the reaction mixture is incubated at 30–35°C, which ensures hybridization of the probes with the target DNA and the emission of fluorescence due to DSN activity.

The optimal length of PCR products for DSNP-analysis was tested empirically: fragments of various lengths containing C- or T-variants of the human mitochondrial COX1 C7028T were hybridized with a T-specific probe. Clear and unambiguous results were obtained for all tested products, proving the possibility of using DSNP analysis for a wide range of amplicon lengths.

Diverse approaches for SNP genotyping have been proposed, based on the difference in physicochemical properties of the variants [11, 12]. With regard to other published protocols, DSNP has several advantages, starting from its overall convenience (the use of crude PCR product, no cleanups/centrifugations, 5 min hands-on and results within 1 hour). Secondly, the protocol allows analysis of both alleles simultaneously in one tube. Thirdly, the specific fluorescence can be recorded using standard laboratory equipment. Finally, the protocol is applicable for virtually any length of the PCR product harboring the polymorphic position.

The study of multiple allelic variants in one tube implies the use of probes with fluorophores that emit at different wavelengths. Effects of different fluorophores on the efficiency of hydrolysis were negligible, unless the mismatches were positioned at the termini. The efficiency of hydrolysis for imperfect duplexes containing an unpaired nucleotide in the midportion did not depend on the type of fluorophores. Importantly, when using probes for different alleles labeled with identical fluorophores, the analysis must be carried out in separate tubes.

DSNP was successfully applied for genotyping of variants involved in a number of diseases or predispositions, including TP53 C309T; F2 G20210A, MTHFR 677T; KRAS G34A, G35T, G35A, and G38A; NRAS G34A, G35C, and G35A; HRAS G35T; APOE C388T; F5 G1698A; and BRCA1 S382insC. Allelic status of the studied samples was confirmed by Sanger sequencing.

The resulting model experiments showed that DSNP allows reliable differentiation between mutant and wild-type alleles in both homozgyous and heterozygous samples. In addition, the example of BRCA1 S382insC demonstrates that, apart from point substitutions, the method is also applicable to single-nucleotide indels. The suitability of the same standard reaction conditions for different genomic positions indicates the universality of the approach.

Clustered occurrence of point mutations in certain genomic regions is well described [13]. The majority of available PCR systems cannot afford accurate determination of such closely located mutations by routine genotyping. On an example of KRAS, with a mutagenesis hotspot at positions 34 and 35 (G34A, G35A, and G35T [14]), it has been demonstrated that DSNP analysis is suitable for genotyping of closely-spaced point mutations even in multiplex. Simultaneous use of up to four FRET-probes, inclusive, produced specific signal only when the probe was fully complementary to the target.

Occasionally, it may be important not only to detect a variant, but also to measure the allelic ratio for a sample. Such tasks may be relevant for tumor tissue samples or pooled genomic DNA samples from multiple donors [12, 15]. Experiments with KRAS G35A as a model proved the possibility of using DSNP for semi-quantitative determination of mutant alleles in complex samples.

Thus, the natural properties of DSN have qualified this enzyme as a basis for a genotyping protocol termed DSNP, fairly simple and automatable. Most prominent advantages of this protocol include:

1) the use of crude PCR products with arbitrary fragment lengths;
2) no purification/separation required;
3) rapidity (1 hour, starting from PCR products);
4) the allelic ratio assessment option;
5) universal applicability: the method is suitable for determination of single-nucleotide substitutions and indels, at clustered positions or not, regardless of the context.

The protocol requires no special equipment apart from that ubiquitously found in the labs. Even fluorescence signals can be assessed with the use of ordinary instruments: for instance, with fluorescein, the signal can be recorded with a conventional UV lamp used in gel-doc systems.

DSNP disadvantages compared with real-time PCR include:
1) preparative PCR amplification of the target fragment;
2) endpoint detection;
3) risk of contamination associated with the need to open tubes with amplicons.

**cDNA normalization**

Heterogeneous gene expression levels in a cell complicate the full-scale analysis of transcriptomes and gene hunting. Normalization of cDNA libraries prior to analysis allows to increase the sensitivity towards rare transcripts.

The classical principle of cDNA normalization involves hybridization kinetics. As hybridization rate is proportional to the squared concentration of molecules in a sample, high-copy fragments renature faster than low-copy fragments [16]. Separation of reassociated double-stranded fragments after denaturation of a complex cDNA sample affords a library with equalized concentrations of abundant and rare transcripts [17–19].

The existing normalization protocols differ by the means of separation of the normalized single-stranded (ss) and double-stranded (ds) fractions. The possibilities include physical separation of fractions using hydroxyapatite chromatography [17, 20] or paramagnetic beads [19, 21], dsDNA digestion with restriction endonucleases [18], and selective amplification of ssDNA using the PCR suppression effect [22]. Unfortunately, these possibilities are hardly adaptable for normalization of cDNA samples enriched with full-length sequences.

The unique properties of DS enabled a highly efficient and easy-to-perform method, known as DSN-normalization and now a routine at many laboratories in Russia and across the world, universally applicable for normalization of both fragmented and full-length cDNA. Like most its predecessors and counterparts, this method is based on the kinetics of cDNA reassociation, but differs in the way the normalized ssDNA fraction is separated [23].
Table. Representation of repetitive elements in human genomic DNA before and after DSN normalization

<table>
<thead>
<tr>
<th>Families of repetitive sequences</th>
<th>Control DNA sample (no normalization)</th>
<th>Normalized DNA sample</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>b.p.</td>
<td>%</td>
</tr>
<tr>
<td>Total length</td>
<td>6 643 277</td>
<td>100</td>
</tr>
<tr>
<td>LINE L1P</td>
<td>371 253</td>
<td>5.6</td>
</tr>
<tr>
<td>ERV-K</td>
<td>20 597</td>
<td>0.3</td>
</tr>
<tr>
<td>SSR</td>
<td>68 367</td>
<td>1</td>
</tr>
<tr>
<td>Alu</td>
<td>631 112</td>
<td>9.5</td>
</tr>
<tr>
<td>Satellite sequences</td>
<td>293 827</td>
<td>4.4</td>
</tr>
<tr>
<td>ERV1</td>
<td>165 461</td>
<td>2.5</td>
</tr>
<tr>
<td>ERV-L</td>
<td>323 437</td>
<td>4.9</td>
</tr>
<tr>
<td>LINE L1M/HAL</td>
<td>496 398</td>
<td>7.5</td>
</tr>
<tr>
<td>DNA transposons</td>
<td>161 304</td>
<td>2.4</td>
</tr>
<tr>
<td>MIR</td>
<td>118 927</td>
<td>1.8</td>
</tr>
<tr>
<td>LINE L2</td>
<td>104 535</td>
<td>1.6</td>
</tr>
<tr>
<td>LINE CR1/L3</td>
<td>10 226</td>
<td>0.1</td>
</tr>
<tr>
<td>LINE L4-L5</td>
<td>3 508</td>
<td>0.05</td>
</tr>
</tbody>
</table>

After hybridization, the reaction mixture is treated with DSN to remove the non-target fraction of dsDNA. Since DSN is a thermostable enzyme active at 70 °C, the hydrolysis occurs at the same temperature as hybridization. The high temperature affords minimization of non-specific binding and thereby prevents the loss of transcripts prone to formation of secondary structures. The normalized ss cDNA fraction is amplified by PCR.

The method is also applicable to non-amplified first strand cDNA. Abundant transcripts (‘majors’) are sponged through renaturation of the first strand cDNA with the poly(A)+ RNA that has served as a template for its synthesis. This protocol is applicable with large amounts of biomaterial available so that it is possible to isolate the poly(A)+ RNA fraction from total RNA. It should be noted that in the normalized libraries, the content of clones corresponding to certain highly represented transcripts is sometimes lower than the number of clones corresponding to rare transcripts. Such “supernormalization” can be explained by the continued dominance of major RNA species, which serve as guides for elimination of complementary DNA molecules. Upon the release after DSN-mediated hydrolysis of the complementary DNA strand, major RNAs can form new hybrids with single-stranded DNA molecules, thus promoting their hydrolysis, and so on.

For the subsequent use in a variety of applications, the normalized first strand cDNA must be amplified. For this reason, preparation of first strand cDNA for DSN normalization necessarily involves ligation of adapter sequences, which will provide annealing sites for oligonucleotide PCR primers. As is well-known, in PCR, short fragments are amplified more efficiently than longer fragments. Accordingly, amplification of complex normalized cDNA is fraught with the loss of long transcripts and decreased average length of the library. To preserve the fraction of long cDNA molecules during library preparation, short inverted repeats are included in the design of the adapters. PCR amplification with a primer matching the inverted repeat (albeit shorter) favors amplification of long cDNA molecules against the background of suppressed amplification of short molecules [24]. According to experimental data, normalization of amplified cDNA, although generally less efficient than the first strand cDNA normalization, also provides significant leveling of over-represented transcripts to enable the search for rare mRNA species under circumstances when only total RNA is available.

To date, DSN normalization provides both the simplest and most effective means for cDNA normalization. By contrast with many other protocols, it involves no physical separation of DNA fractions. Furthermore, it can be used to normalize both amplified cDNA and the non-amplified first strand of cDNA enriched with full-length molecules. In addition, DSN-normalization preserves the average lengths as well as the length distributions of cloned cDNA libraries.

Specific enrichment and normalization of genomic DNA libraries

Song et al. (2016) developed a protocol for enrichment with minor alleles (including those with mutations of clinical or biological significance) through selective elimination of wild-type alleles in mixed (pooled) clinical samples, termed Nuclease-Assisted Minor-Allele enrichment using Overlapping Probes, NaME-PrO [25]. The simultaneous removal of the excess of wild-type DNA for a virtually unlimited number of target genomic sequences is performed before amplification. The unique properties of DNS ensure priority cleavage of wild-type DNA regardless of genomic context.

For each target sequence, a pair of oligonucleotide probes is designed to bind the target region on opposite strands, with a 10–15 bp overlap between them. The probes are added in excess to the fragmented genomic DNA denatured at 98 °C. When the temperature is lowered to 67 °C, DNA remains single-stranded due to its low concentration and slow reassociation kinetics. The probes anneal to their target sites in DNA, whereby they create pinpoint mismatches on complementary strands upon their contact with mutated DNA, resulting in imperfect duplexes. Upon exposure to DSN, which preferentially cleaves perfect duplexes, the wild-type DNA is cleaved whereas mutant DNA remains substantially intact. Because the two probes match the target sequence on opposite strands, both strands of wild-type DNA undergo preferential cleavage within the region covered by the probes. Thus, if at least one of the DNA strands containing the mutation is preserved after DSN cleavage, then subsequent DNA amplification will lead to exponential amplification providing the multiplex enrichment for all mutated sites simultaneously. This approach abolishes the need for deep sequencing to detect rare mutations.
As demonstrated by the authors, NaME-PRO affords 50 to 200-fold enrichment for a variety of target mutations found in clinical samples (exemplified by KRAS mutations).

In connection with the rapid technological progress, next generation sequencing (NGS), in particular whole-genome sequencing, is becoming an increasingly common approach in basic science and clinical laboratory diagnostics. In eukaryotes, a significant proportion of genomic DNA consists of highly homologous repetitive elements. Their presence not only increases the cost of genome sequencing, but also makes the bioinformatics processing and interpretation of the data extremely difficult.

The problem can be solved by several approaches. In particular, selective repetitive elements can be suppressed through the employment of the pronounced tendency of repetitive sequences to hypermethylation. Yuan et al. (2002) used selective cleavage of hypermethylated regions with restriction endonucleases sensitive to cytosine 5’-methylated [26, 27]. Similarly, Palmer et al. (2003) used the methylation-dependent endonuclease McrBC from the E. coli K-12 strain in the construction of maize genomic libraries, thereby limiting the cloning of highly methylated DNA [28]. Such technical solutions, however, are not applicable to organisms with other methylation patterns.

An alternative solution to the problem, the so-called Cot filtration, is based on the kinetics of DNA renaturation. Genomic DNA is fragmented, heat denatured and cooled. Since low-copy DNA fragments rehybridize slower than repetitive elements, over a certain time the single-stranded fraction becomes enriched with low-copy sequences [29, 30]. Next, the double-stranded fraction containing repetitive elements is separated from the single-stranded fraction (in the classic version, by hydroxyapatite chromatography). Although Cot filtration may work with any complex mixture of heterogeneously represented repetitive elements, it is possible to employ the pronounced tendency of repetitive sequences to hypermethylation.

Shagina et al. (2010) investigated the possibility of using DSN normalization to eliminate the highly homologous repetitive elements from genomic sequencing libraries. DNA is subjected to fragmentation, supplemented with adapter sequences through ligation, and denatured by heating. During the renaturation process, the sample is treated with DSN. The preserved single-stranded fraction of genomic DNA enriched in low-copy sequences is amplified with primers corresponding to the adapter sequences [31].

The method was tested in a model experiment on normalization of human genomic DNA before sequencing in a 454 GS FLX system (Roche). To enhance the sponging of non-target sequences, hybridization was carried out in an excess of the Cot-1 fraction of human genomic DNA. For the normalized and control samples, 29,240 and 31,789 reads were obtained with a total coverage of 6,269,460 and 6,643,277 nucleotides, respectively. Representation of diverse repetitive elements in the sequencing data was determined using the RepeatMasker software available at repeatmasker.org/cgi-bin/WEBRepeatMasker. According to the results, normalization reduced the content of repetitive elements from 40% to 25%.

The Table shows representation of different families of repetitive elements in non-normalized (control) and normalized samples of human genomic DNA. Significant effects can be observed for Alu, LINE L1P, ERV-K, and ERV1 repeats, as well as satellite sequences. At the same time, certain families of repetitive elements show resistance to normalization by this method.

Reciprocally, control samples (no normalization) contained about 10% of sequences sharing 100–91% similarity and about 20% of sequences sharing 90–71% similarity, whereas the remaining 70% of sequences had identical nucleotides in less than 71% positions. DSN normalization reproducibly reduced the content of low-divergent repetitive elements (100–91% identical) 15-fold and medium-divergent repetitive elements (81–90% identical) 2-fold. Concentrations of other sequences in the samples did not decrease with DSN-normalization. Preservation of single-copy genomic sequences during DSN normalization was demonstrated by real-time PCR assay on a panel of 11 unique genes.

These findings indicate that DSN normalization can effectively reduce the content of the evolutionary young low-divergent repetitive sequences in genomic DNA samples. The cut-off threshold can be lowered by using milder reassociation conditions (e.g., by lowering the temperature and/or increasing the cation concentration), albeit with the risk of partial loss of unique sequences due to increased non-specific interactions.

**MicroRNA studies**

MicroRNA molecules are increasingly considered as promising biomarkers for diagnosis and monitoring of various pathologies, including cancers and autoimmune disorders. They are found in the blood plasma both in a freely circulating form and as part of the exosomal fraction. MicroRNAs are easy to isolate, resistant to degradation, and show reproducible and characteristic expression patterns.

The unique properties of DSN, particularly its indifference to RNA substrates, can be used to create specific chemiluminescent and fluorescent sensors for miRNA [32, 33]. A system developed by Shen et al. (2015) contains biotinylated DNA molecules (probes) labeled with fluorescent and immobilized on magnetic beads. The microRNA and the beads, the medium contains DSN, which recognizes and cleaves the duplexes formed upon binding of microRNA with the probe. After the cleavage, the labeled outer fragment of the probe drifts into the medium, while its microRNA partner finds and binds the next immobilized DNA molecule, promotes its cleavage, and so on. The fluorescently labeled cleavage products eventually accumulate in the medium. In the end, the beads with immobilized unreacted probes are separated from the reaction medium with a magnet. The labeled cleavage products remain in the medium for the endpoint detection of fluorescence. The system is sensitive enough to detect femtomolar microRNA concentrations. Noteworthy, in contrast to protocols that use quantitative PCR, amplification of the signal is carried out isothermally at 40 °C, which makes the proposed method even more attractive [33].

**CONCLUSION**

Duplex-specific nuclease from hepatopancreas of the king crab exerts a unique combination of properties including the exquisite substrate specificity (selectively digests double-stranded DNA without affecting single-stranded DNA or RNA), the high optimal temperature of catalysis (60–65 °C), and the thermal stability (retains activity at 90 °C). Since its original characterization in 2002, the crab nuclease has been featured in diverse molecular protocols, which still evolve and are continually updated. The enzyme has been successfully used in a wide range of applications, including genotyping of single nucleotide polymorphisms (in both experimental and clinical samples), normalization of cDNA and genomic DNA libraries, selective elimination of non-target sequences, and miRNA studies.
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DYNAMIC CHANGES IN THE CONCENTRATION OF ANTI-SARS-COV-2 ANTIBODIES WITHIN 12 MONTHS AFTER RECOVERY FROM COVID-19

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Generation and maintenance of immunity to SARS-CoV-2 is essential for overcoming the pandemic of the novel coronavirus infection COVID-19. The study was aimed to assess the dynamic changes in the levels of IgG antibodies against the SARS-CoV-2 receptor-binding domain (RBD) with the use of the enzyme-linked immunosorbent assay (ELISA) kits, calibrated using the International Standard for anti-SARS-CoV-2 immunoglobulin (IS-SARS-CoV-2). The concentrations of anti-RBD-IgG were measured in the cohort of individuals, who had recovered from COVID-19, with an interval of a month for 6 months, and at a time point of 12 months, using the ELISA kits, calibrated with the use of IS-SARS-CoV-2; the results were expressed in binding antibody units (BAU) per 1 mL. A total of 97 blood serum samples, obtained from 20 individuals with SARS-CoV-2 infection, confirmed by PCR, were collected. The geometric mean titer (GMT) of anti-RBD-IgG was 433 BAU/mL (range 36-25,900 BAU/mL) within a month after the infection. The concentration of anti-RBD-IgG gradually decreased with time and reached the GMT value of 68 BAU/mL by the 12th month; anti-RBD-IgG persisted in 13 individuals (93%) out of 14, examined 12 months after the infection. The standardized quantitative serological data play a vital part in monitoring the immune response and make it easier to compare the studies, providing the basis for seeking the common serological correlate of the protective immunity to SARS-CoV-2.

KEYWORDS: SARS-CoV-2, anti-RBD IgG, dynamic changes, concentration, BAU/mL

Author contribution: Mayansky NA — concept, data processing, manuscript writing; Brzhozovskaya EA — sample collection, ELISA, data processing, making illustrations; Stoyanova SS — sample collection, data processing, making illustrations; Frolov AV — data processing, manuscript preparation; Lebedin YuS — concept, ELISA, manuscript editing.

Compliance with ethical standards: the study was approved by the Ethics Committee of the Pirogov Russian National Research Medical University (protocol № 197 dated May 21, 2020).

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The natural infection and vaccination against COVID-19 result in the production of antibodies against viral antigens, playing a vital part in the immune response monitoring [1]. Although it is expected that the naturally acquired immunity against SARS-CoV-2 would last long [2-3], serological equivalent of antiviral host defense has not yet been discovered. The lack of a standardized approach to laboratory testing is one of the obstacles to defining such correlates, which probably explains the conflicting literature data on the serological assessment of SARS-CoV-2 infection. The WHO have recently introduced the International Standard for anti-SARS-CoV-2 immunoglobulin (IS-SARS-CoV-2), which makes it possible to unify the results of measuring the levels of anti-SARS-CoV-2 antibodies using the IS-SARS-CoV-2 units, namely the binding antibody units (BAU) [1]. The study was aimed to measure the concentration of IgG against the SARS-CoV-2 receptor-binding domain.
(RBD) at different times over the 12-month period in a cohort of healthcare professionals, who have recovered from the SARS-CoV-2 infection, using the enzyme-linked immunosorbent assay (ELISA) kits, calibrated using the IS-SARS-CoV-2.

METHODS

The study, carried out from May 2020 to June 2021, involved the staff members of the Russian Children’s Clinical Hospital, Pirogov Russian National Research Medical University. Inclusion criteria: positive PCR test result for COVID-19. There were no exclusion criteria.

In April and May 2020, after returning to work, individuals with positive PCR test results for COVID-19 gave the blood serum samples with an interval of a month for the anti-RBD-IgG measurement. The samples were collected on a monthly basis for 6 months, and the last sample was obtained 12 months after the positive PCR test result; a total of 4–7 samples was obtained from each subject. The samples collected were stored at a temperature of –80 °C.

In July 2021, all the samples were assayed in one batch using the ELISA kit for the anti-RBD-IgG quantification (XEMA; Russia) [4], calibrated using the IS-SARS-CoV-2. The analytical measuring range was 15–240 BAU/mL; samples with the anti-RBD-IgG concentration exceeding 240 BAU/mL were further diluted 10–100 times and measured repeatedly in a separate batch. The samples were considered positive when the level of anti-RBD-IgG was 15 BAU/mL.

Statistical processing was performed using the IBM SPSS Statistics 27.0 software package (IBM Corp.; USA).

RESULTS

A total of 97 serum samples were obtained from 20 individuals during the study, including 14 women (70%) with the SARS-CoV-2 infection, confirmed by PCR. The median age was 50 years (Q1–Q3, 40–57 years). All the subjects had mild to moderate COVID-19, there were no severe cases of the disease.

In three individuals, blood serum samples were obtained a month before the SARS-CoV-2 infection; no anti-RBD-IgG were found in these samples. A month after the positive PCR test result for SARS-CoV-2, anti-RBD-IgG were found in all subjects in a concentration exceeding the threshold value of 15 BAU/mL, with the geometric mean titer (GMT) of 433 BAU/mL (95% CI 123–1,527 BAU/mL; range 36–25,900 BAU/mL) (see Figure). At the time point 2 of the month the median GMT was similar, 456 BAU/mL (95% CI 154–1,353 BAU/mL). Later the concentration of anti-RBD-IgG gradually decreased and reached the median GMT value of 68 BAU/mL (95% CI 35–131 BAU/mL) by the 12th month. All samples were anti-RBD-IgG-positive during the period between the first and the sixth months. Among 14 individuals assessed 12 months after the SARS-CoV-2 infection, 13 individuals (93%) were still seropositive for anti-RBD-IgG with the GMT values exceeding 15 BAU/mL. The median concentration of anti-RBD-IgG was 6.7 times lower (95% CI 4.4–10.3 times) after 12 months compared with the highest median GMT value, defined during the second month after the SARS-CoV-2 infection.

DISCUSSION

To date, the data of only a few studies, involving the standardized values of the amount of antibodies against SARS-CoV-2 after the natural infection and/or vaccination, have been reported [5–7]. Thus, the live viral neutralization assay showed that anti-RBD-IgG in a concentration of ≥100 BAU/mL ensured the complete neutralization of three SARS-CoV-2 variants of concern a year after infection, which reduced the risk of reinfection with these virus strains [5]. Another report mentioned that on day 14 after vaccination, the anti-RBD-IgG GMT in individuals, vaccinated with mRNA vaccine, was 7,756 BAU/mL [6]. Thus, quantitative results provide the basis for seeking the common serological correlate of the protective immunity to SARS-CoV-2. Moreover, these data are important for monitoring the natural immunity and facilitate the comparison of immune responses to various vaccines [1]. Significant differences and systematic error in the numerical results, expressed in BAU/mL, associated with the use of different test systems [7], encourage further efforts to unify serological tests for detection of antibodies against SARS-CoV-2.

CONCLUSIONS

The study analyzes dynamic changes in the antibody response to SARS-CoV-2 infection during the natural immunity production. The use of IS-SARS-CoV-2 for calibration of the ELISA test system made in possible to demonstrate a change in the concentration of anti-RBD IgG over a wide range of 36–25,900 BAU/mL with the widest disparity within two months after the infection. The levels of anti-RBD-IgG gradually decreased with time, however, positive values persisted throughout 12 months of follow-up in the majority of subjects.
References


Литература


PROPERTIES OF RBD SPECIFIC IGG FROM COVID-19 PATIENTS AND SPUTNIK V VACCINATED INDIVIDUALS

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SARS-CoV-2 specific antibody response is a generally accepted measure of postinfection and vaccination-induced immunity assessment. The dynamics of avidity maturation and neutralizing activity of virus-specific immunoglobulins during the SARS-CoV-2–associated coronavirus infection was studied in cohorts of vaccinated volunteers and COVID-19 patients. 4–6 months after vaccination, neutralization activity was low compared to hospitalized patients (medians 57.4% vs 86.4%). On the opposite, the avidity indices in vaccinated volunteers were significantly higher (median 76.7%) than among hospitalized patients (median 61.4%).

During the acute phase of the disease (14–16 days PI), post-vaccination patients have also higher avidity indices than primary patients (medians 43.5% vs 20.4%). Our results suggest that in long-term perspective antibody affinity maturation rate is higher after vaccination than after a natural infection. We demonstrated that Sputnik V vaccination leads to formation of high-avidity IgG, which persists for at least 6 months of observation. These results also indicate the presence of protective efficacy markers for at least 4–6 months after the vaccination or a previous illness and gives grounds for the half-year time period chosen for booster immunization with Sputnik V in Russia.

Keywords: antibody avidity, virus neutralization, SARS-CoV-2, immune memory, vaccination, Sputnik-V

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Author contribution: LV Generalova, IV Grigoriev — research planning, experiments preparation and execution, data interpretation and paper draft preparation; IS Kruzhkova, LV Kolobukhina — data interpretation and paper draft preparation; DV Vasina, AP Tkachuk, OA Burgasova, VA Gushchin — research planning, data interpretation and paper draft preparation.

Compliance with ethical standards: the study was approved by the ethics committee of the First Moscow Infectious Diseases Hospital (protocol № 11/A dated November 16, 2020); informed consent was obtained from all study participants.

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SARS-CoV-2 specific antibody response is a generally accepted measure of postinfection and vaccination-induced immunity assessment. However, the protective efficacy of virus specific antibodies and their ability to withstand the individual’s reinfection may be influenced not only by antibody quantity but also their quality including neutralizing activity, binding affinity, isotypes spectrum etc., which are not characterized well enough. In general, SARS-CoV-2 antibody avidity (% of antibodies with high affinity) correlated with duration of infection and higher neutralizing titers [1]. Indeed, high avidity antibodies but not the level of spike-binding antibodies has been previously associated with positive clinical outcomes [2]. At the moment there is no published information about the antibody affinity maturation after the Sputnik V vaccination and in recovered patients, as well as antibodies functional transformations during the prolonged period of observation. This study aims to provide the longitudinal assessment of antibody responses dynamics in patients recovered from COVID-19 and Sputnik V vaccinated individuals with or without further infection and to characterize the patterns of sustaining long-term immunity.

METHODS

Study participants

During the study we enrolled 41 participants divided into 3 groups (Table 1): 23 patients, hospitalized in Moscow with different disease severity, were sampled upon the admission to the hospital (acute phase) and 4–6 months after the hospital discharge; 9 vaccinated patients sampled during their hospitalization and 9 healthy vaccinated volunteers at different times from the complete vaccination. Median symptom durations in hospitalized patients before the first sampling were 14 days among unvaccinated individuals and 10 days among vaccinated individuals. For vaccinated volunteers, the absence of COVID-19 was confirmed by the lack of anti-Nc IgG seroconversion.

We characterized antibodies from the three groups of participants. The group of inpatient volunteers was recruited in November through December 2020 at the First Moscow Infectious Diseases Hospital (Moscow, Russia); vaccinated patients were hospitalized and enrolled in March through April 2021 in the same source. Healthy volunteers vaccinated with Sputnik V vaccine were recruited in September through December 2020. Eligible volunteers were adults aged 18–80 years.

All patients were either diagnosed with SARS-CoV-2 infection by RT-PCR of nasopharyngeal swabs or confirmed by IgG seroconversion. For vaccinated volunteers, the periods of sustaining long-term immunity were 14 days among unvaccinated individuals and 10 days among vaccinated individuals. For vaccinated volunteers, the absence of COVID-19 was confirmed by the lack of anti-Nc IgG seroconversion.

Table 1. Study Cohorts. Values are reported as the medians with the range in parentheses

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Patients (n = 23)</th>
<th>Vaccinated volunteers (n = 9)</th>
<th>Volunteers (n = 9)</th>
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<tbody>
<tr>
<td>Age median years, IQR</td>
<td>59 (54–65)</td>
<td>72 (69–79)</td>
<td>34 (33–39)</td>
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<td>Sex, %</td>
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<td>52% male, 48% female</td>
<td>33% male, 67% female</td>
<td>56% male, 44% female</td>
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<tr>
<td>Time from symptom onset to Visit 1 (Me days min-max)</td>
<td>14 (9–17)</td>
<td>10 (5–19)</td>
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<tr>
<td>Time from symptom onset to Visit 2 (Me days min-max)</td>
<td>135 (116–159)</td>
<td>16 (11–24)</td>
<td>NA</td>
</tr>
<tr>
<td>Time from initial vaccination to Visit 1 (Me days min-max)</td>
<td>NA</td>
<td>62 (30–88)</td>
<td>147 (100–263)</td>
</tr>
<tr>
<td>Time from initial vaccination to Visit 2 (Me days min-max)</td>
<td>NA</td>
<td>68 (35–95)</td>
<td>261 (161–349)</td>
</tr>
</tbody>
</table>

Blood sample processing and storage

Blood samples were collected by venipuncture to vacutainers with clot activator and shipped to the laboratory at +4 °C. Centrifugation at 3000 rpm for 10 minutes was applied to obtain serum, which was further aliquoted and stored at −30 °C.

Anti-nucleocapsid and anti-RBD IgG antibody detection

The anti-nucleocapsid (Nc) and anti-RBD IgG antibodies were measured using in house ELISA test-systems and expressed in positivity index (p.i., S/CO). Briefly, for antibody detection we used recombinant receptor-binding domain fragment of S1 SARS-CoV-2 Spike protein (RBD No. 8COV1; HyTest, Russia), expressed in eukaryotic cells and recombinant SARS-CoV-2 Nucleocapsid (Nc) protein, expressed in Escherichia coli and purified in our laboratory.

To perform anti-RBD and anti-Nc ELISAs, 96 well high binding plates (Costar 2592; Corning, USA) were coated overnight with 100 μl of 1 μg/mL recombinant protein solution in PBS. Next day the plates were blocked for 2 hours at room temperature with a blocking buffer, containing 0,5% casein. Serum samples were diluted 1 : 100 with universal ELISA buffer S011 (XEMA; Russia), expressed in eukaryotic cells and recombinant SARS-CoV-2 Nucleocapsid (Nc) protein, expressed in Escherichia coli and purified in our laboratory.

To perform anti-RBD and anti-Nc ELISAs, 96 well high binding plates (Costar 2592; Corning, USA) were coated overnight with 100 μl of 1 μg/mL recombinant protein solution in PBS. Next day the plates were blocked for 2 hours at room temperature with a blocking buffer, containing 0,5% casein. Serum samples were diluted 1 : 100 with universal ELISA buffer S011 (XEMA; Russia), expressed in eukaryotic cells and recombinant SARS-CoV-2 Nucleocapsid (Nc) protein, expressed in Escherichia coli and purified in our laboratory.

Blood samples were collected by venipuncture to vacutainers with clot activator and shipped to the laboratory at +4 °C. Centrifugation at 3000 rpm for 10 minutes was applied to obtain serum, which was further aliquoted and stored at −30 °C.

Neutralization assay (Inhibiting assay)

Detection of IgG neutralizing antibodies was performed with commercial “SARS-CoV-2-anti-RBD-ELISA” kit (MT-
И-С1-04.192; MedipalTech, Russia) in accordance with manufacturer's instructions. Briefly, serum samples were mixed with biotinylated recombinant human ACE-2 receptor and added to the ELISA plate, precoated with recombinant RBD. After incubation and wash, streptavidin-conjugated HRP was added to the wells. Reaction was visualized by adding an HRP-substrate solution. Optical density in wells was inversely proportional to the concentration of antibodies, able to block ACE-2 binding to RBD. Inhibition coefficient (IC) was calculated as ratio of sample OD to negative control OD, subtracted from 1. IC = (1 — ODsample/ONneg) — 100%.

The antibody neutralization criteria were assessed as high neutralizing if serum sample inhibited binding of soluble RBD to ACE-2 on plate by > 30%; intermediate one, if the inhibition rate was between 20 and 30%; and a low level for samples with inhibition < 20%.

**Avidity assay**

The IgG avidity assay targeted the spike protein RBD of SARS-CoV-2 was performed with “SARS-CoV-2-ELISA-IgG plus” kit (No MT-И-С1-03.192; MedipalTech, Russia) in accordance with manufacturer's instructions. Briefly, duplicate anti-RBD IgG positive serum samples were incubated in wells of ELISA plates, precoated with recombinant RBD. Then either PBS (non-denaturing conditions) or urea solution (denaturing agent, DA) was added to the wells. After washing, HRP-conjugated anti-human IgG was added and, finally, reaction was visualized by

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**Fig. 1.** Immune responses to SARS-CoV-2 in the studied groups. (A) Cohort of hospitalized patients (B) Cohort of vaccinated patients (C) Cohort of healthy vaccinated volunteers. Significant differences are shown as asterisks (Wilcoxon signed-rank test). Me — median time of sampling in groups from the symptoms onset or the initial vaccination as indicated in the axis.
adding HRP-substrate solution. Avidity index (a.i.), proportional to antibody denaturation resistance, was calculated as ratio of OD-450 in DA and PBS wells. The antibody avidity criteria were as follows: avidity index (a.i.) > 50% — high avidity; between 40 and 50% — intermediate avidity; < 40% — low avidity.

Data analysis
The results are reported as medians with min-max range and the parameters were compared between groups by the Mann–Whitney nonparametric t test and at different time points using the paired Wilcoxon rank test. P value < 0.05 was considered statistically significant.

RESULTS
All antibody assays were provided for 100% of samples. Samples of 23 hospitalized patients with confirmed COVID-19 demonstrated a common Ab dynamics pattern. Two weeks after the symptoms onset all samples became IgG positive for both RBD (Me S/CO 10.5, min-max, 6.8–9.0) and Nc (Me, 6.8, min-max 5.6–8.9) (Fig. 1, Table 2). Twenty two out of 23 samples demonstrated high neutralization activity in convalescent patients, blood samples were collected 4–6 months later. All samples remained positive for anti-RBD and anti-Nc IgG (Fig. 2, Table 2). Neutralization activity remained at high level without significant difference between first and second time point were obtained after 10–12 days after the symptoms onset, thus, they had not enough time to reach peak of serum antibody concentrations due to heterogeneity of SARS-CoV-2 incubation period and variability in the individual dynamics of Ab production. However, four months later all patients with one exception had similar levels of anti-RBD IgG (Me S/CO 6.8, IQR 6.3–8.9).

Although we detected a 1.5-fold decrease of anti-RBD IgG levels of this cohort, the neutralization activity remained at high level without significant difference between first and second time points which indicates that the quantity of antibodies is not the only determinant of the neutralizing activity. Preservation of high neutralization effects can be due to the compensation of decreased antibody concentration by improved quality (specificity and affinity) during this period. On the other hand, the observed effect may be due to the peculiarities of the test system used in the study. Since the measured values are near the limit of quantification of the system it does not allow to reliably assess the changes in neutralizing activity. Additional experiments are required to access neutralization dynamics more precisely. During the same observation period anti-RBD IgG avidity increased significantly, reaching a high avidity level (>50%) in 78% of patients. Median avidity at the second time point was 61.4% (min-max 18.2–86.9%). Remaining
6 patients (22%) had intermediate (40–50%, “maturation zone”) or low (<40%) anti-RBD IgG avidity. Importantly, five of low- and intermediate-avidity patients had moderate or severe disease (Fig. 3). One patient (Fig 1A, orange circles) had low levels of anti-RBD IgG and neutralization activity at both time points, but his anti-RBD IgG avidity increased from 20.4 to 42.7%. To the end of the hospitalization period most patients possessed strong IgG response to SARS-CoV-2 antigens and their antibodies were effective at preventing RBD binding to human ACE-2. During the next months affinity maturation of anti-RBD IgG response occurred and overall concentration of anti-RBD and anti-Nc IgG decreased, neutralization activity remained at high level.

In a group of healthy Sputnik V vaccinated volunteers (Fig. 1B) 8 out of 9 samples were positive for anti-RBD IgG at the first time point (median of 147 days after the initial vaccination) with 4.9 S/CO (min-max 1–7.9). Four months later (261 days post initial vaccination), the median anti-RBD IgG p.i. dropped significantly to a median level 1.7 S/CO although 7 out of 9 volunteers still remained positive for anti-RBD IgG. None of the volunteers had detectable IgG to Nc, indicating there were no cases of infection with SARS-CoV-2 in this group. The neutralization correlated the IgG dynamics and decreased significantly in all samples from median 57.4% in the first time point to median 30.6% in the second time point. On the opposite, the avidity remained at the same high level indicating that at the first time point (147 days post vaccination), antibody maturation occurred in all vaccinated volunteers. However, for 3 samples (33%) at the second time point it was impossible to estimate the avidity as the level of anti-RBD IgG was too low (Fig. 1B, highlighted in blue).

In a group of Sputnik V vaccinated patients with confirmed COVID-19 (Fig 1C, Table 1) the median time from receiving a first vaccine dose was 62 days (ranging from 30 to 88 days) and the median time of sampling from symptoms onset was 10 days (5 to 19 days). At that time 6 out of 9 samples were positive for anti-RBD IgG with median 4.3 S/CO and only two were positive for anti-Nc IgG. All anti-RBD positive samples and one negative sample were positive in the neutralization ELISA (median 71.6%, min-max 7.4%–98.6%). Most of the samples (66.7%) demonstrated low avidity antibodies with median a.i. 27.6% (min-max 0%–71.1%). The second sampling time point in this group was around the 16th day after the symptoms onset. One week after the first sampling, the median S/CO of anti-RBD IgG in general did not change significantly (median 7 S/CO, min-max 0.8–11.2), however, 7 out of nine patients were seropositive after the infection and/or vaccination. The neutralization did not significantly change compared to the first time point (median 92.9%, min-max 23.2%–99.2%), the same was observed for avidity. Neutralization activity in two samples with low levels of anti-RBD IgG increased significantly to 85 and 89%, probably due to anti-RBD IgM production. (Fig. 1, yellow dots)

DISCUSSION

In COVID-19 patients, neutralizing antibody titers correlate with the severity of the infection [5, 6] and can be achieved even at low somatic hypermutation [7, 8]. The RBD region is found to be immunodominant and it is the target of approximately 90% of the neutralizing antibodies presented in the sera of SARS-CoV-2-infected people. Furthermore, it has been determined that
anti-RBD IgG titers decrease with time post symptom onset, presenting a half-life of approximately 49 days. Importantly, antibody avidity increases over time due to increased maturation (somatic hypermutation, followed by selection in germinal centers). In the serum of hospitalized COVID-19 patients, there is a higher number of IgG against S protein and RBD, compared with that in non-serious and asymptomatic patients [9]. It was previously shown that antibody avidity increased over duration of infection and remained elevated [1]. In convalescent donors, plasma higher neutralizing titer had a stronger positive correlation with anti-spike IgG avidity than with anti-nucleocapsid IgG avidity proposing the anti-RBD IgG to be the main source of neutralizing activity.

Comparing samples of hospitalized patients and vaccinated volunteers 4–6 months after the infection or the vaccination, vaccinated volunteers had significantly lower levels of anti-RBD IgG and neutralization activity, but significantly higher anti-RBD IgG avidity (Fig. 4A, C). All hospitalized patients remained positive for anti-RBD IgG, while in one of the vaccinated volunteers antibody levels dropped below baseline (Fig. 4C).

When comparing the groups of Sputnik V fully vaccinated patients and unvaccinated patients samples, collected during the acute phase of infection (2–3 weeks after symptoms onset), it was shown that, in vaccinated patients, antibody levels to both RBD and Nc antigens were significantly lower, than in unvaccinated patients (Fig. 4B). Anti-RBD IgG avidity was significantly higher in vaccinated patients indicating that despite low antibody levels, vaccination did induce primary immune response and formation of memory B-cells and their antibody production during the infection demonstrates signs of secondary immune response. In the long-term perspective antibody affinity maturation rate is higher after vaccination than after natural infection. Although it is proposed that antibody maturation increases their neutralization potency [10], here we observe no significant correlation between these parameters.

Possibly this may indicate that estimation of RBD-specific antibodies avidity can serve not only as a prognostic marker of protection before the 6-month period. We propose that low anti-RBD IgG avidity two months after vaccination could be one of the potential markers for preliminary revaccination. Our study is under certain significant limitations. First of all, it is based on a small number of samples. Patients in this study had a limited spectrum of COVID-19 clinical manifestations. Particularly, our research does not include asymptomatic and outpatient individuals with a mild course of disease, which represent the most of COVID-19 cases in comparison with other disease manifestations. Second, we could not estimate the peaks of antibody production and maturation as only two timepoints were analyzed in each group.

CONCLUSIONS

We observed the anticipated dynamics of Ab levels in convalescent patients with the peak at acute phase followed by gradual decrease in the subsequent months. But despite the loss of anti-RBD antibodies concentrations, serum neutralization activity remained at high/sufficient level, probably due to improved Ab specificity and increased avidity. Here, we demonstrated the formation and persistence of high avidity IgG for at least 6 months after Sputnik V immunization. It indicates that protectivity markers remain at sufficient levels for at least 4–6 months after the infection or the vaccination. These results give rounds for the half-year period chosen for booster immunization with Sputnik V in Russia. Based on our data one can’t estimate the proportion and characteristics of patients requiring earlier revaccination due to drop of vaccine-induced protection before the 6 months period. We propose that low anti-RBD IgG avidity two months after vaccination could be one of the potential markers for preliminary revaccination. Possibly this may indicate that estimation of RBD-specific antibodies avidity can serve not only as a prognostic marker of
Comparison of immune responses in different groups. (A) Comparison of unvaccinated convalescent patients and healthy vaccinated volunteers 4–6 months after the infection/vaccination. (B) Comparison of vaccinated and unvaccinated patients in the acute phase of infection (2–3 weeks from symptoms onset). (C) Comparison of anti-RBD IgG conversion in the studied groups. (D) Comparison of sera avidity dynamics in the studied groups. Significant differences are shown as asterisks (Mann–Whitney U test).

disease severity, but also to determine individuals, who require revaccination earlier than 6 months after receiving the initial dose. Further study on larger samples is needed to verify this hypothesis.

References


Literatura


GUT MICROBIOTA ALTERATIONS AND THEIR RELATIONSHIP TO THE DISEASE SEVERITY AND SOME CYTOKINE PROFILE INDICATORS IN PATIENTS WITH COVID-19

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Gut microbiota is an essential element of maintaining the immune homeostasis, including in individuals with COVID-19. The study was aimed to assess taxonomic changes in the gut microbiota and their relationship with the disease severity and the levels of IL6, IL10, IL17, and TNFα in patients with COVID-19. A total of 110 patients with COVID-19 (index group) and 98 individuals with no COVID-19 (control group) were enrolled to the comparative cross-sectional study. The gut microbiota composition was determined by shotgun sequencing. Blood serum levels of IL6, IL10, IL17, and TNFα were assessed by enzyme-linked immunosorbent assay. The following significant changes in the gut microbiota composition were observed in patients with COVID-19 in contrast to controls: decreased abundance of B. adolescentis (p = 0.048), E. rectale (p = 0.036), F. prausnitzii (p = 0.0002), B. dorei (p < 0.001), and increased abundance of R. gnavus (p = 0.012), C. hathewayi (p = 0.003), E. faecium (p = 0.0003). Correlations were established between the abundance of B. dorei and the IL6 levels (r = 0.49; p = 0.034), the abundance of F. prausnitzii and the levels of IL10, IL17 (r = 0.44; p = 0.001 and r = 0.52; p < 0.001, respectively). The abundance of R. gnavus correlated with the TNFα levels, and the abundance of E. faecium was related to the levels of IL6 (r = 0.47; p = 0.002) and TNFα (r = 0.56; p = 0.001). The relationship between the abundance of B. dorei, F. prausnitzii, E. faecium and the higher SHOKS-COVID clinical assessment scale scores was also revealed (r = –0.54; p = 0.001, r = –0.60; p < 0.001 and r = 0.067; p = 0.005, respectively). Targeted correction of gut microbiota may improve the COVID-19 treatment efficacy.

Keywords: COVID-19, SARS-CoV-2, gut microbiota, cytokine status

Author contribution: Gumenyuk LN, Sorokina LE — significant contribution to the study concept and design; Golod MV, Silaeva NV, Androschyuk NA — data analysis and interpretation; Ilyasov SS — statistical data processing; Krivoshapko OR, Velilyaev AM, Asanova LN — manuscript writing.

Compliance with ethical standards: the study was approved by the Ethics Committee of the S.I. Georgievsky Medical Academy, VI. Vernadsky Crimean Federal University (protocol № 11 dated November 23, 2021), planned and conducted in accordance with the Declaration of Helsinki. The informed consent was obtained from all study participants.

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World Health Organization declared the pandemic in response to the novel coronavirus infection, caused by the SARS-CoV-2 virus, which was named COVID-19 [1]. Severe forms of this disorder are associated with progressive viral pneumonia and acute respiratory distress syndrome (ARDS). The increasing systemic inflammation plays a vital part in the COVID-19 pathophysiology. Excessive cytokine production, which is caused by SARS-CoV-2 and known as cytokine storm, is closely related to the disease severity [2]. High levels of interleukins IL1α, IL1β, IL6, IL8, IL9, IL10, IL17, C-reactive protein (CRP), tumor necrosis factor (TNFα), are found in patients with COVID-19 [3–5]. Accumulated data of the in vitro and in vivo studies show that gastrointestinal tract (GIT) is also susceptible to COVID-19. For
example, studying the in vitro model, simulating the cellular and spatial intestinal structure, showed the SARS-CoV-2 capability of infecting enterocytes [6]. The frequency of gastrointestinal symptoms in patients with COVID-19 reaches 20% [7]. This is caused by the SARS-CoV-2 ability to enter cells by binding to the angiotensin-converting enzyme 2 (ACE2) receptor, which is intensively expressed on the enterocyte surface in the ileum and colon. The latter results in gastrointestinal symptoms due to the virus-induced immune-mediated damage [8]. Moreover, viral RNA of SARS-CoV-2 can be detected in fecal samples more than 30 days after the disease onset [9]. In this regard the role played by gut microbiota is actively discussed in literature. As is known, gut microbiota is an essential element of maintaining the immune homeostasis. For its part, gut microbiota dysbiosis is directly related to numerous inflammatory disorders [10]. Strong evidence of quantitative changes in the gut microbiota of patients with COVID-19 is provided [11]. In general, a downward trend in the bacterial species diversity is observed in patients with COVID-19, together with the depletion of beneficial commensals and enrichment in pathobionts [12]. However, information about the species composition of gut microbiota is fragmented and conflicting. Taking into account the potential of interactions, realized through the gut–lung axis, to increase the host susceptibility to viral infections and reduce the functional activity of immune cells, thus contributing to systemic hyperinflammation and cytokine storm syndrome, further investigation of the gut microbiota alterations and their relationship with the cytokine status indicators in patients with COVID-19 is relevant.

The study was aimed to assess taxonomic changes in the gut microbiota and their relationship with the disease severity and the levels of IL6, IL10, IL17, and TNFα in patients with COVID-19.

METHODS

A total of 110 patients with COVID-19 (66 females (60.0%), 44 males (40.0%); the average age was 28.6 ± 8.4 years), who had been admitted to hospitals, working in the Obligatory Medical Insurance system in Simferopol (index group, IG), and 98 healthy volunteers with no COVID-19 (61 females (62.2%), 37 males (37.8%); the average age was 29.2 ± 7.6 years), who had their annual health examination at the Gemokod medical center, Simferopol (control group, CG), were enrolled to the comparative cross-sectional study by continuous sampling.

Inclusion criteria for the IG: age 18–45 years; COVID-19, confirmed by PCR test for the SARS-CoV-2 RNA and/or the typical multislice computed tomography (CT) findings of viral pneumonia; mild, moderate or severe COVID-19.

Exclusion criteria for the IG: extremely severe COVID-19; type 1 or 2 diabetes mellitus; obesity; myocardial infarction, severe cardiac arrhythmias, heart failure; history of hypertensive disease, stroke, transient ischemic attack; acute cerebrovascular disease (within six months before the beginning of the study); severe or decompensated concomitant somatic diseases, which could make it more difficult for the patient to participate in the study and affect the study results; irritable bowel syndrome; chronic gastrointestinal and liver diseases; hematological and oncological diseases; history of mental disorders, alcoholism or drug addiction; taking antibiotics, probiotics, prebiotics, antiviral drugs, sibiotics or acid–suppression medications within three months before the beginning of the study; taking medications, affecting the passage of stool, within a month before the beginning of the study; refusal to participate in research.

Inclusion criteria for healthy volunteers: age 18–45 years; no COVID-19, confirmed by PCR test for the SARS-CoV-2 RNA; no chronic disorders or allergic reactions; no infectious or acute disorders within two months before the study; no history of mental disorders, alcoholism or drug addiction; no abnormal passage of stool (constipation/diarrhea); taking no antibiotics, probiotics, prebiotics, antiviral drugs or sibiotics within three months before the beginning of the study; taking no medications, affecting the passage of stool, within a month before the beginning of the study.

Exclusion criteria for healthy volunteers: body temperature above 36.9 °C; refusal to participate in research.

Characteristics of patients with COVID-19 and healthy volunteers are presented in Table 1. The groups were comparable in gender ($\chi^2 = 0.96$; $\chi^2 = 0.92$), age ($\rho = 0.92$; $\rho = 0.96$), and body mass index ($\rho = 0.054$; $\rho = 0.054$).

Clinical characteristics of patients with COVID-19 are provided in Table 2. Individuals with moderate COVID-19 (80 patients (72.7 ± 0.68%)) predominated among patients. Fever (103 patients (94.2%)) and cough (96 patients (86.9 ± 0.61%)) were the most common symptoms of the disease. According to CT images, viral pneumonia was diagnosed in 95 patients (67.1 ± 0.38%).

In all patients, the diagnosis of COVID-19 was confirmed by PCR test for SARS-CoV-2 (specimens were obtained from nasopharyngeal and oropharyngeal swabs) and/or typical multislice CT findings of viral pneumonia. The diagnosis and severity of COVID-19, as well as the extent of pneumonia on CT images, were assessed in accordance with the Temporary Guidelines on Prevention, Diagnosis and Treatment of Novel Coronavirus Infection (COVID-19) of the Ministry of Health of the Russian Federation, versions 6–9.

The original Clinical Assessment Scale for patients with coronavirus infection (SHOKS-COVID) was used to objectify the severity of clinical manifestations [13].

In order to analyze the taxonomic composition of the patients’ gut microbiota, fecal samples were collected on the first day of hospital stay (in the morning, 8 a.m. to 10 a.m.). The samples were frozen and stored in disposable plastic containers at a temperature of ~80 °C before the metagenomic analysis. Total DNA was isolated by the phenol-based extraction. Nucleotide sequence of the DNA isolated was determined by shotgun sequencing using the SOLID 5500 Wildfire system for high-throughput sequencing (AppliedBiosystems; USA) [14]. QiIME software version 1.9.1 was used to filter the reads based on their quality and perform taxonomic classification [15]. The approach, involving the use of two taxonomic databases, was applied to perform the taxonomic assignment of the reads. At the first stage, the reference set of the bacterial

<table>
<thead>
<tr>
<th>Indicators</th>
<th>Index group ($n = 110$)</th>
<th>Control group ($n = 98$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Females / males ($n$, %)</td>
<td>66 (60.0%) / 44 (40.0%)</td>
<td>61 (62.2%) / 37 (37.8%)</td>
</tr>
<tr>
<td>Average age years (M ± CD)</td>
<td>28.6 ± 8.4</td>
<td>29.2 ± 7.6</td>
</tr>
<tr>
<td>Body mass index, kg/m² (M ± CD)</td>
<td>23.4 ± 4.2</td>
<td>23.7 ± 3.6</td>
</tr>
</tbody>
</table>
operational taxonomic units (OTUs) was selected based on the comparison of the 16S rRNA gene reads obtained with the GreenGenes database, version 13.5 [16]. At the second stage, taxonomic assignment of these OTUs was performed using the RDP algorithm based on the custom human intestinal database, HiTdb [17].

Qualitative and quantitative assessment of the gut microbiota composition involved identification of species, genera, and phyla of microorganisms. Assessment of α-diversity by calculating the Chao 1 index, number of the taxa observed (Sobs), and the indicator of species richness (ACE), was performed with the Mothur v.1.22.0 software (http://www.mothur.org).

IL6, IL10, IL17 and TNFα were selected as the cytokine profile markers, which was due to their key role in the pathogenesis of gut microbiota and ARDS [3–5].

Blood samples were collected from peripheral vein during the first day of hospital stay. Blood was taken in the morning (between 7 and 9 a.m.) in a fasting state at rest (for at least 15 min). Test tubes containing blood serum were frozen and stored at a temperature of −20 °C. Serum levels of IL6, IL10, IL17, and TNFα were assessed using the enzyme-linked immunosorbent assay (ELISA) test system (Vector-Best; Vector-Best; Novosibirsk, Russia) with the Elisys Quattro automated ELISA analyzer (Human GmbH; Germany).

Statistical data processing was performed using the STATISTICA 8.0 software package (StatSoft. Inc.; USA). In case of normal distribution, the mean and the standard deviation were defined, and in case of non-normal distribution, the median, 25th and 75th percentiles were calculated. Distributions were tested for normality using the Gaussian distribution. Percentage and absolute values were defined for qualitative traits. Comparative analysis for the normal distribution of quantitative traits was performed using the parametric Student’s t-test. Comparative analysis of non-normal distribution it was performed using the Mann–Whitney U test, and comparative analysis of quantitative traits was carried out using the chi-squared test ($\chi^2$). Spearman’s correlation coefficient was used to assess the relationship between the traits. The differences were considered significant when $p < 0.05$. Correlation analysis and multiple rank correlation were also applied, and the correlations’ significance was tested with the contingency tables.

RESULTS

Assessing the taxonomic composition of gut microbiota revealed a significant decrease in the α-diversity of the bacterial community (Chao1 index; $p = 0.018$) in patients with COVID-19 compared to controls. The ACE and Sobs indices were also slightly decreased in the group of patients with COVID-19 compared to the CG, however, no significant differences were observed ($p = 0.054$; $p = 0.052$, respectively) (Fig. 1).

Comparison of the species composition of gut microbiota revealed a significant decrease in the abundance of Blifdobacterium adolescentis SPM1005-A ($p = 0.048$), Eubacterium rectale ATCC 33656 ($p = 0.036$), Faecalibacterium prausnitzii A2-165 ($p = 0.0002$), Bacteroides dorei DSM 17855 ($p = 0.001$) in patients with COVID-19 compared to controls, together with the increased abundance of Ruminococcus gnavus ATCC 29149 ($p = 0.012$), Clostridium hathewayi DSM-13479 ($p = 0.003$), and Enterococcus faecium W54 ($p = 0.0003$) (Fig. 2).

Serum levels of IL6, IL10, IL17, and TNFα were significantly higher in patients with COVID-19 compared to the CG (Table 3).

Clarification of the relationship between the gut microbiota alterations and some cytokine profile indicators in patients with COVID-19 revealed significant correlations between the abundance of B. dorei and the IL6 levels ($r = −0.49; p = 0.034$), abundance of F. prausnitzii and the IL10, IL17 levels ($r = −0.44; p = 0.001$ and $r = −0.52; p = 0.001$, respectively). The abundance of R. gnavus correlated with the levels of TNFα ($r = 0.51; p = 0.036$), and the abundance of E. faecium was related to the levels of IL6 ($r = 0.47; p = 0.002$) and TNFα ($r = 0.56; p = 0.001$). Correlation analysis also revealed the relationship between the abundance of B. dorei, F. prausnitzii, E. faecium, and the higher SHOKS-COVID scores ($r = −0.54$ and $p = 0.001$; $r = −0.60$ and $p < 0.001$; $r = 0.67$ and $p = 0.005$, respectively).

Table 2. Clinical characteristics of patients with COVID-19

<table>
<thead>
<tr>
<th>Indicator</th>
<th>Index group ($n = 110$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mild course ($n, %$)</td>
<td>7 (6.3)</td>
</tr>
<tr>
<td>Moderate course ($n, %$)</td>
<td>80 (72.7)</td>
</tr>
<tr>
<td>Severe course ($n, %$)</td>
<td>23 (21.0)</td>
</tr>
<tr>
<td>Fever</td>
<td>103 (94.2)</td>
</tr>
<tr>
<td>Sore throat</td>
<td>68 (61.8)</td>
</tr>
<tr>
<td>Cough</td>
<td>96 (86.9)</td>
</tr>
<tr>
<td>Shortness of breath</td>
<td>63 (56.9)</td>
</tr>
<tr>
<td>Nausea</td>
<td>37 (33.6)</td>
</tr>
<tr>
<td>Diarrhea</td>
<td>24 (22.1)</td>
</tr>
<tr>
<td>Temperature, median [25%; 75%]</td>
<td>37.8 [36.7; 37.9]</td>
</tr>
<tr>
<td>RR per minute, median [25%; 75%]</td>
<td>20.0 [17.0; 22.0]</td>
</tr>
<tr>
<td>HR, beats/min (m ± CD)</td>
<td>89.9 ± 16.1</td>
</tr>
<tr>
<td>SBP, mm Hg, median [25%; 75%]</td>
<td>120 [120; 132]</td>
</tr>
<tr>
<td>SO2, %, median [25%; 75%]</td>
<td>96.0 [94.0; 98.0]</td>
</tr>
<tr>
<td>Severity of the condition, SHOKS-COVID score, points, Me [25%; 75%]</td>
<td>3.4 [1.6; 6.1]</td>
</tr>
<tr>
<td>CT stage — 0 ($n, %$)</td>
<td>25 (22.6)</td>
</tr>
<tr>
<td>CT stage CT–1 ($n, %$)</td>
<td>41 (37.4)</td>
</tr>
<tr>
<td>CT stage CT–2 ($n, %$)</td>
<td>32 (29.1)</td>
</tr>
<tr>
<td>CT stage CT–3 ($n, %$)</td>
<td>12 (10.9)</td>
</tr>
</tbody>
</table>

Note: RR — respiratory rate, HR — heart rate, SBP — systolic blood pressure, SO2 — oxygen saturation, CT — computed tomography.
DISCUSSION

In a number of previous studies, alterations in the composition of gut microbiota in patients with COVID-19 have been reported [7, 11, 12]. Our findings have also shown significant differences in the composition of gut microbiota between patients with COVID-19 and individuals with no COVID-19. According to our data, patients with COVID-19 have a lower bacterial diversity compared to individuals with no COVID-19, which is confirmed by the significantly lower Chao1 index and is consistent with the results of previous studies [11]. In patients with COVID-19, gut dysbiosis is characterized by the decreased abundance of bacteria having the immunomodulatory potential, the type representatives B. adolescentis, E. rectale, F. prausnitzii, B. dorei, that are known to be the main butyrate-producing bacteria (butyrate is a powerful anti-inflammatory metabolite). At the same time, we have revealed the increased abundance of potential pathobionts: the bacteria R. gnavus, Cl. hathewayi, and E. faecium. Among them, E. faecium is noteworthy, the presence of which in fecal samples of patients with COVID-19 has been previously reported by Italian researchers [18]. High levels of E. faecium in the gut microbiota of critically ill patients could be of some clinical significance due to the E. faecium pathogenic potential, resistance to common antimicrobial drugs, and the ability to rapidly acquire genetic material or alter gene expression, allowing the bacteria to acquire the
resistance determinants to almost all antibacterial agents [19, 20]. Independently from the strain of the genus Enterococcus, gut microbiota of patients with COVID-19 can therefore function as a reservoir of the potentially antibiotic-resistant opportunistic pathogens, able to migrate through the damaged epithelial barrier into the systemic circulation, as has already been shown in the context of other disorders [20]. Our findings are to some extent consistent with the previously reported research data. For example, in one of the studies, patients with COVID-19 were characterized by the decreased abundance of B. dorei, and the increased abundance of E. faecium [18].

In another study, patients with COVID-19 were characterized by the decreased abundance of B. adolescentis, E. rectale, F. prausnitzii, and the increased abundance of bacteria R. gnavus [21]. The contrasting data patterns could be due to the fact, that the studies were carried out in different geographic regions with the use of different inclusion criteria. The groups varied considerably in age: 28.6 ± 8.4 years in our study vs. 73.0 [59.0; 85.0] in the previously reported study [18], and 36.4 ± 18.7 years in the study [21]. Moreover, in contrast to the listed above researchers, we did not study patients with somatic comorbidities and patients, taking antibacterial and/or antiviral medications, in order to mitigate their influence on the study results.

Despite the fact that some bacteria identified could be common to a number of other disorders, both gastrointestinal and systemic, the determined relationship between the decreased levels of B. dorei, F. prausnitzii, increased levels of E. faecium and the higher SHOKS-COV scores suggests that changes in the abundance of these bacteria may be typical for this cohort of patients with COVID-19. We have compared the associations determined with the results of earlier studies. Thus, depletion of members of the genus Bacteroides was observed in patients, admitted to the intensive care unit [18]. It is interesting that in another study [7], the abundance of B. dorei negatively correlated with the SARS-CoV-2 load in the faeces of patients with COVID-19. The researchers noted that, taking into account the association of this bacterium with the decreased ACE2 receptor expression in the murine colon, B. dorei could potentially provide protection against the SARS-CoV-2 virus. Data are presented on the negative correlation between the bacterium F. prausnitzii, known for its anti-inflammatory effect, and the COVID-19 severity [7]. At the same time, higher abundance of strains of the genus Enterococcus \( p = 0.0001 \) was observed in patients with COVID-19, admitted to the intensive care units, compared to patients, admitted to the general medicine units [18]. It can be assumed that therapeutic upscaling of B. dorei, F. prausnitzii together with the E. faecium downscaling is effective for mitigation of the disease severity. However, further research with appropriate design is required to confirm this hypothesis. Some authors point out that probiotics and/or prebiotics may be used as a potential (additional) vector of the COVID-19 therapy to reduce the disease severity and minimize the risk of secondary bacterial infections [22, 23].

A number of studies have proven the impact of gut microbiota on the susceptibility to infectious and noncommunicable diseases [24]. It has been specified that the intestinal flora immunomodulatory effect is realized through activation of genes, encoding a number of cytokines in the immune and epithelial cells, which determines heterogeneity of their immunomodulatory properties [25]. It has been reported that heterogeneity of the immunoregulatory effects is typical for both dominant and associative microsymbionts [26]. Thus, cumulative effects of the normal bacterial flora representatives on the secretion of cytokines by human immune cells in the conditions of eubiosis ensure the cytokine balance, characterized by moderate levels of pro-inflammatory cytokines (IL6, TNFα) and regulated by the suppressing effects of antiinphlogogenic cytokines. Thus, in the context of increasing antigen load in dysbiosis through activation of toll-like receptors, the increased production of a whole range of pro-inflammatory cytokines is observed. These cytokines promote both local inflammation and the effector immune response in the gut-associated lymphoid tissue, protecting the body from pathogens [27].

As previously reported, COVID-19 infection is associated with the increased levels of cytokines IL6, IL10, IL17, and TNFα [4]. In our study, significant differences in the levels of IL6, IL10, IL17, and TNFα between patients with COVID-19 and healthy volunteers were also observed.

It is assumed that the COVID-19 severity results from the cytokine storm [4]. It is noteworthy that some of the listed above cytokines correlate with the gut microbiota pattern, the specific profile of which is capable of inducing the cytokine storm. Our study revealed the correlation between the abundance of bacteria B. dorei and the levels of IL6 in patients with COVID-19.

We have revealed the relationship between the abundance of bacteria F. prausnitzii and the levels of IL10, which could be mediated by the decreased secretion of metabolites, blocking the NF-kB transcription factor activation and the IL8 production. For example, the in vivo study showed that activation of the peripheral blood mononuclear cells by F. prausnitzii resulted in the significantly reduced secretion of IL12 and TNFα, together with the increased secretion of IL10 [28]. Furthermore, the abundance of bacteria F. prausnitzii negatively correlated with the IL17 levels. When performing literature analysis, we have found no publications on studying the relationship between gut microbiota and IL17 in patients with COVID-19. At the same time, it has been shown that F. prausnitzii inhibits IL17 production in rats [29]. Studying the experimental colitis model has demonstrated that bacteria F. prausnitzii exert their anti-inflammatory effects due to production of butyrate, which maintains the balance between the pro-inflammatory T helper 17 cells (Th17) and immunoregulatory T cells (Treg) via inhibition of histone deacetylase 1. In turn, imbalance between Th17 and Treg promotes autoimmune inflammation [30].

We have also revealed the relationship between the abundance of the bacteria Ruminococcus gnavus and the TNFα levels. These data are consistent with the previously reported research results [31], showing that higher levels of R. gnavus in patients with COVID-19 correlate with the surge of pro-inflammatory cytokines IFNγ and TNFα, resulting in the cytokine storm and activation of the type 1 helper T cells.

We have discovered that the abundance of bacteria E. faecium correlates with the levels of IL6 and TNFα. We
have found the reports of similar correlations in patients with ulcerative colitis [32] and AIDS/HIV [33].

Thus, in the context of the COVID-19-associated gut dysbiosis, orientation of immunoregulatory effects towards proand anti-inflammatory cytokines is limited in the normal flora representatives, which contributes to the impaired homeostasis and the development of inflammatory and autoimmune responses.

CONCLUSIONS

The relationship between the decreased levels of bacteria B. dorei, F. prausnitzii together with the increased levels of bacteria E. faecium and the higher SHOKS-COVID scores has been defined in patients with COVID-19, which is indicative of the pathognomonic nature of these taxonomic alterations of intestinal microflora in individuals, infected with SARS-CoV-2. Significant correlations between the abundance of gut bacteria B. dorei, F. prausnitzii, R. gravis, E. faecium and the levels of IL6, IL10, IL17, TNFα have been found in the SARS-CoV-2-infected patients, which confirms the gut microbiota capability of being involved in systemic inflammation and maintaining the immune tolerance in COVID-19. Our findings in combination with the data reported in literature demonstrate the key role in the COVID-19 pathogenesis played by microbiota. The more systematic and detailed research in this area would enable the development of new approaches, as well as selective probiotics for microbiota correction in patients with COVID-19 and treatment of post-COVID effects.

References


izy182.
Pseudomonas aeruginosa is a significant opportunistic pathogen and a serious burden to public health and economy [1]. Especially dangerous are carbapenem-resistant strains of P. aeruginosa regarded by WHO as critical priority pathogens [2]. This breeds the need for understanding mechanisms underlying bacterial resistance to carbapenems. Research into the molecular genetic underpinnings of carbapenem resistance focuses mostly on β-lactamase-associated mechanisms that are determined by plasmid genes and therefore can be acquired through horizontal gene transfer. However, there is another contributor whose role should not be overlooked: induced mutations in the core genome of P. aeruginosa resulting in high-level carbapenem resistance [3]. There are two approaches to the study of mutations conferring resistance to carbapenems. The first involves the analysis of drug-resistant isolates obtained from clinical, agricultural or environmental sources. In the second approach, the evolution of antibiotic resistance is modeled in vitro. For that, bacteria are grown in antibiotic concentration gradients. A smart method for studying mutational resistance has been proposed in [4]. Its authors...
created a spatiotemporal model that enabled migration of *Escherichia coli* in trimethoprim and ciprofloxacin gradients and generated a variety of mutants for further analysis. Interestingly, some of the *E. coli* clones carried mutations that were not linked to trimethoprim or ciprofloxacin resistance [4]. So, we became curious to explore the direction and implications of such mutations. Specifically, we were interested in the clinically significant phenomenon of cross-resistance, in which a mutation induced by exposure to an antibiotic could confer resistance to other antibiotics [5, 6]. The aim of this study was to test the hypothesis that *P. aeruginosa* can develop cross-resistance to other antibiotics while adapting to meropenem.

**METHODS**

**Bacteriological study**

In our experiment, we used the spatiotemporal model of antibiotic resistance in motile bacteria [4]. The reference ATCC 27853 strain of *P. aeruginosa* was precultured in semi-solid LB agar (0.28% agarose) in a Petri dish at 37 °C for 24 h. After 24 h, the cells were harvested from the propagating colony front and seeded onto another Petri dish with semi-solid LB agar. The procedure was repeated 3 times. Then, 10 µl of the grown bacterial culture was picked up with an inoculation loop and introduced into the top layer (semi-solid agar) of the culture medium contained in a device shown in the Figure. The medium had a sandwich composition. The bottom layer was LB Miller broth (Becton Dickinson; USA) supplemented with 1.6% agarose, 30 µg/ml kanamycin sulfate, 100 µg/ml cycloheximide, and meropenem taken at one of the concentrations shown in the Figure. The optimum thickness of the bottom layer equaled three-fifths of the total medium thickness (~2.0 cm). The bottom layer was distributed into 5 isolated compartments of the dish containing different concentrations of meropenem. The middle layer (one-fifth of the total medium thickness) was LB Miller broth supplemented with 2.0% agarose, 30 µg/ml kanamycin sulfate, 100 µg/ml cycloheximide, and ink (4.0 ml per 1L culture medium) added as a contrasting background for photography purposes. The middle layer spread over the bottom layer was solid. The top layer (one-fifth of the total medium thickness) was semi-solid agar (Miller LB broth) with 0.3% agarose, 30 µg/ml kanamycin sulfate and 100 µg/ml cycloheximide.

The cells were incubated in air at 37 °C for 216 h. Every 12 h, *P. aeruginosa* samples were collected from the propagating colony front and reseeded on Mueller–Hinton agar (Becton Dickinson; USA) to obtain a sufficient amount of bacteria for the subsequent analysis of their phenotypic traits (antibiotic resistance profiles) and genomic changes.

Resistance to meropenem and imipenem was tested using the agar dilution method described in [7]. Resistance to colistin was assessed using the broth microdilution method following the guidelines of the Clinical and Laboratory Standards Institute (CLSI) [8].

Bacterial DNA was isolated from the 24-h culture of *P. aeruginosa* grown on Mueller–Hinton agar (Becton Dickinson; USA) using a QiAamp DNA Mini Kit (Qiagen; Germany) according to the manufacturer’s protocol. The obtained DNA samples were stored at −20 °C.

To prepare genomic DNA libraries, 400 g of the isolated bacterial DNA was sheared in an ultrasonicator (Covaris; USA). The fragments were then end-repaired and ligated to MGI adapters (MGI; China). The libraries were purified on Agencourt AMPure XP beads (Beckman; USA). Concentrations of the bacterial DNA and DNA libraries were measured using a Qubit 4 fluorometer (Thermo Fisher Scientific; USA).

Whole-genome sequencing was performed using the MGISEQ-2000 platform (MGI; China). Read length was 250 bp.

The quality of the raw sequence data was tested in FASTQC; the reads were trimmed in Trimmomatic v.0.38. Bacterial genomes were assembled de novo using SPAdes 3.14 [9]. The assembled sequences were tested for contamination using Contest16S. The obtained genome assemblies were evaluated in QUAST 5.0 [10]. Genetic similarity between the assembled genomes was assessed in MUMmer [11]. The genomes were annotated using RAST [12] and Prokka software [13]. To detect the presence of single nucleotide polymorphisms (SNPs), the short reads were mapped to the reference genome in Snippy [14]. ATCC 27853 was used as a reference genome. The detected variants were annotated and their influence on the genes was predicted in SnpEff [15]. The search for antibiotic resistance genes in the genomes assembled de novo, their analysis and validation of the detected SNPs were all carried out using the program PRINSEQ [16].

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**Table. Characteristics of carbapenem-resistant strains of *P. aeruginosa* with reduced susceptibility to colistin**

<table>
<thead>
<tr>
<th>Methodology</th>
<th>Characteristic</th>
<th><em>P. aeruginosa</em> ATCC 27853, reference</th>
<th><em>P. aeruginosa</em>, isolate E82</th>
<th><em>P. aeruginosa</em>, isolate E74</th>
</tr>
</thead>
<tbody>
<tr>
<td>Phenotypic antibiotic susceptibility assessment</td>
<td>meropenem</td>
<td>0.25 µg/ml, S</td>
<td>16 µg/ml, R</td>
<td>16 µg/ml, R</td>
</tr>
<tr>
<td></td>
<td>imipenem</td>
<td>0.001 µg/ml, S</td>
<td>128 µg/ml, R</td>
<td>256 µg/ml, R</td>
</tr>
<tr>
<td></td>
<td>colistin</td>
<td>0.5 µg/ml, I</td>
<td>2 µg/ml, I</td>
<td>4 µg/ml, R</td>
</tr>
<tr>
<td>Changes in genome</td>
<td>oprD</td>
<td>wt</td>
<td>Mutation resulting in G307D</td>
<td>Mutation resulting in G307D</td>
</tr>
<tr>
<td></td>
<td>mexD</td>
<td>wt</td>
<td>Mutation resulting in E89K</td>
<td>Mutation resulting in E89K</td>
</tr>
<tr>
<td></td>
<td>phoQ</td>
<td>wt</td>
<td>Nonsense mutation resulting in Y290stop</td>
<td>Nonsense mutation resulting in Y290stop</td>
</tr>
<tr>
<td>Polymorphism</td>
<td>SNP</td>
<td>—</td>
<td>134</td>
<td>177</td>
</tr>
</tbody>
</table>

**Note:** S — susceptible; I — intermediate; R — resistant; wt — wild type, matches the reference genome; SNP — single nucleotide polymorphism; " — according to the CLSI criteria, the term “susceptible” cannot be applied to describe the susceptibility of *P. aeruginosa* to colistin; all *P. aeruginosa* strains for which colistin MIC ≤ 2 µg/ml are classified as susceptible at increased exposure.
The P. aeruginosa phoQ strains with simultaneous resistance to oprD32 study proves that carbapenems and colistin. The phenomenon observed in our through consecutive or simultaneous therapeutic exposure to literature. It is possible that they acquire their resistance profiles [18]. The evolution of such isolates is rarely described in the meropenem-resistant isolates were unsusceptible to colistin and increased resistance to meropenem and imipenem. Phenotypic and genotypic characteristics of these 2 strains are shown in the Table.

For Е62, meropenem and imipenem MICs were 16 μg/ml and 128 μg/ml, respectively; for Е74, they were 16 μg/ml and 256 μg/ml, respectively, which satisfied the CLSI criteria for antibiotic resistance. According to CLSI criteria, the Е62 isolate was characterized as susceptible to colistin at increased exposure; for this strain, colistin MIC was 4 times higher than for the baseline strain. According to the CLSI criteria, the Е74 strain was characterized as resistant to colistin (MIC: 4 μg/ml).

Both strains carried a mutation in the porin gene (oprD) resulting in the substitution of glycine with aspartic acid at position 307 of the protein. Besides, both isolates had a missense mutation in the mexD gene (this gene encodes the subunit of the MexCD-OprJ efflux pump). Also, both Е62 and Е74 had a nonsense mutation in the phoQ gene resulting in the premature termination of protein synthesis (289 out of 448 amino acids).

**DISCUSSION**

P. aeruginosa strains with simultaneous resistance to carbapenems and polymyxins are not rare. For example, among multidrug resistant P. aeruginosa representatives, 22.2% of meropenem-resistant isolates were unsusceptible to colistin [18]. The evolution of such isolates is rarely described in the literature. It is possible that they acquire their resistance profiles through consecutive or simultaneous therapeutic exposure to carbapenems and colistin. The phenomenon observed in our study proves that P. aeruginosa can reduce their susceptibility to colistin following exposure to meropenem. The hypothetical mechanisms underlying induction of cross-resistance to colistin by meropenem fall into the “all roads lead to resistance” concept, meaning that in P. aeruginosa any stressor causes hypermutability and leads to the emergence of multiple clones with novel properties [3]. Such a mutational explosion can lead to the emergence of persisting mutations disrupting synthesis of lipopolysaccharides, the primary target of colistin.

Genomes of the Е62 and Е74 isolates carried mutations that can explain their resistance to meropenem/imipenem and reduced susceptibility to colistin. The missense mutation in the oprD gene reported in this study may have caused a structural change in the OprD porin, which transports meropenem and imipenem inside the bacterial cell [19]. The search of GeneBank (https://www.ncbi.nlm.nih.gov/genbank) identified only one clinical isolate with a similar amino acid sequence of the OprD porin (GCA_003194245.1). Similar to our mutant, this isolate obtained in 2013 was also resistant to meropenem and imipenem (MIC > 32 μg/ml). Another mutation that could have reduced susceptibility to carbapenems was the missense mutation in the mexD gene encoding the subunit of the MexCD-OprJ efflux pump. The MexCD-OprJ system is involved in the efflux of β-lactams; its hyperexpression is correlated with carbapenem resistance in P. aeruginosa [20]. The phoQ gene codes for the sensor histidine kinase, which is part of the two-component regulatory PhoPQ system. Mutations in phoQ were reported to cause resistance to polymyxins in P. aeruginosa, including specimens isolated from patients with cystic fibrosis [21, 22].

Thus, all phenotypic characteristics of carbapenem-resistant isolates of P. aeruginosa with reduced susceptibility to colistin observed in our study were associated with mutations.

**CONCLUSIONS**

The phenomenon of cross-resistance described in this paper may be due to the fact that the rate of point mutations in P. aeruginosa, specifically in the genes implicated in antimicrobials resistance, increases under stress conditions. Our findings prove that exposure to meropenem can lead to resistance not only to other β-lactams but also to colistin used as a last resort drug for P. aeruginosa infections, which seriously complicates the treatment strategy and limits its options.
Литература


DNA mismatch repair is a unique biological mechanism for repairing DNA damage occurring during cell division [1]. Proteins involved in DNA mismatch repair are encoded by 6 key genes: MSH2, MLH1, PMS2, MSH3, MSH6, and MLH3. Inherited or sporadic mutations, as well as other epigenetic events like MLH1 promoter hypermethylation, can inactivate any of these genes and disrupt the normal functioning of the entire mechanism [2]. This results in the accumulation of multiple unrepaired mutations in the genome and changes to the length of microsatellites, which are 2–9 bp-long sequences in the euchromatic portion of the genome [3]. Consequently, if the microsatellite is located within an intron region, a reading frame shift may occur in the coding sequence, followed by the inactivation of various genes. Thus, defects in the DNA mismatch repair mechanism give rise to a genomic phenotype known as microsatellite instability (MSI) [1].

MSI-positive tumors are less aggressive than those that do not have the MSI phenotype, presumably due to a high
The aim of our study was to evaluate the MSI status in colorectal NENs, which, in our opinion, may hold promise for future research.

Associated with a specific set of its molecular characteristics, these findings show that the MSI-status of the tumor is correlated to the presence of mutations in the BRAF gene (MGMT tumors (40.6% vs 20.2%); the most frequently methylated tumors (11/89), including NECs of the colon, stomach and adenoneuroendocrine carcinomas (NECs; 33%).

So far, there have been no randomized clinical trials evaluating the effectiveness of different immunotherapy regimens for colorectal neuroendocrine neoplasms (NENs). Besides, the prevalence of MSI-positive colorectal and gastrointestinal NENs remains understudied, as is the impact of the MSI status on the clinical outcome.

We hypothesize that MSI-positive colorectal NENs are a separate group of tumors with a different clinical presentation and prognosis. Colorectal NENs are relatively rare, so enrolling a large number of patients in the clinical trial may pose a problem. In the largest molecular genetic studies of colorectal NENs conducted so far, the average number of patients did not exceed 100. In earlier publications estimating colorectal NENs conducted so far, the average number of patients did not exceed 100. In earlier publications estimating colorectal NENs, the average number of patients did not exceed 100.

Due to the versatility of this marker, which expands indications for CPI therapy, testing for the patient’s MSI status is becoming an essential diagnostic procedure approved unanimously by the leading cancer research communities, including ASCO, ESMO, NCCN, and RUSSCO [2]. The MSI status affects the choice of treatment strategy for patients with early-stage CRC, which is being reflected in contemporary clinical guidelines [5].

Today, MSI is recognized as an important predictor of tumor response to immunotherapy regardless of the primary tumor site [1].

The study included 29 patients undergoing surgical treatment for colorectal NENs at the National Medical Research Centre for Oncology (Rostov-on-Don) between 2015 and 2018. Of them, 15 were men and 14 were woman. The mean age at diagnosis was 62.5 years. The mean follow-up period was 3.8 years. The following inclusion criteria were applied: expression of neuroendocrine differentiation markers (chromogranin A, synaptophysin) confirmed by immunohistochemistry; informed consent to participate in the study. All tissue specimens were analyzed using the 2019 WHO classification criteria.

DNA was isolated from the paraffinized tissue samples obtained during surgery. Briefly, 10 slices were prepared from the paraffin-embedded tumor or seemingly healthy tissue using a microtome; then, they were deparaffinized in xylol. The samples were incubated with a lysis buffer in the presence of proteinase K at 58°C for 6–12 h until complete tissue lysis. After that, total DNA was isolated and purified using a DNA-sorb-B kit (AmpliSens; Russia) following the manufacturer’s protocol. DNA concentrations were measured with a Qubit 2.0 fluorometer (LifeTechnologies; USA).

Five monomorphic microsatellite loci (NR21, NR24, NR27, BAT25 and BAT26) were tested for their MSI status by means of fragment analysis. Each of the obtained total DNA templates were PCR-amplified in 5 reactions. Each 20 μl PCR reaction contained 10–20 ng of DNA, 0.175 μM of each primer, 2 mM of dNTP, 15 mM of MgCl₂, and 0.5 un. of Taq-polymerase. The following PCR protocol was applied: initial denaturation at 94°C for 5 min, followed by 40 cycles of denaturation at 94°C for 30 s, primer annealing at 59°C, and final extension at 72°C for 45 s. The size of the PCR product ranged from 50 to 350 pn.

Detection of the fluorescently tagged PCR products was performed by means of fragment analysis. Briefly, 1 μl of the obtained PCR products was combined with 19μl of Hi-Di formamide and 0.5 μl of GeneScan™ 600 LIZ Size Standard (Thermo Fisher; USA). The samples were incubated in a CH-100 heating/cooling dry block thermostat (Biosan; USA) at 95°C for 5 min and processed in an ABI PRISM 3500 genetic analyzer (Applied Biosystems; USA) following the manufacturer’s protocol. The obtained data were analyzed in GeneMapper Software (Thermo Fisher; USA). The peak detection value was set to 50 relative fluorescence units (RFU). MSI was concluded

![Fig. 1. Distribution of colorectal NENs by primary tumor location](image-url)
if 2 or more of the studied loci were polymorphic. The level of MLH1 methylation was determined by bisulfate-converted DNA pyrosequencing using a PyroMark Q24 CpG MLH1 assay according to the manufacturer’s protocol (QIAGEN; Germany).

Statistical analysis was carried out in Statsoft Statistica 10.0 (StatSoft; USA) for Windows 10. Primary data analysis was performed using descriptive statistics (central tendencies and measures of variability).

RESULTS

Colorectal NENs were grouped by primary location. Of them, 55% (n = 16) were cecal, 17% (n = 5) were NENs of the rectum, 17% (n = 5) were tumors of the ascending colon, 7% (n = 2) were localized to the right hepatic flexure, and 4% (n = 1) were appendiceal (Fig. 1).

The majority of the patients included in the study had stage III cancer (52%, n = 15) (Fig. 2).

In our cohort of patients, NENs of the colon occurred more frequently than other histological subtypes (55%, n = 16), and their frequency was directly correlated with the stage of the disease. Patient distribution by the histological subtype and stage of cancer is shown in Fig. 3.

Microsatellite stability (MSS) was confirmed in 83% of cases (n = 24); 17% (n = 5) of the tumors were MSI-positive. All MSI-positive tumors were well or moderately differentiated stage I cancers: two of them were G1 and 3 were G2 neuroendocrine neoplasms of the rectum. All of those 5 tumors had microsatellite instability in all of the 5 studied STR loci.

MLH1 was hypermethylated in all MSI-positive specimens (Me = 20%, range: 14–42%) and hypomethylated in all samples with the MSS phenotype (Me = 4%, range: 4–14%). Based on the obtained data, we concluded that the leading cause of the MSI-status in colorectal NENs was inhibition of transcription of the key mismatch repair system genes caused by hypermethylation of their promoter. In the literature, this mechanism was previously described for sporadic colorectal adenocarcinomas.

The distribution of colonic NENs with the confirmed MSS phenotype by primary location and stage is shown in Fig. 4 and 5.

For our patients with MSS-positive NENs of the colon, the three-year survival rate was 50% for stage II, 33% for stage III, and 0% for stage IV. For the patients with stage I MSI-H NENs of the rectum, the three-year survival rate was 100%.

DISCUSSION

Our findings on the prevalence of microsatellite instability in colorectal NENs were compared to the published data on colorectal cancer. In a recent study of Russian researchers conducted in a Russian cohort of patients with CRC (n = 359), the MSI phenotype was observed in 6.4% of cases (23/359) and was correlated with younger age (p = 0.023), the presence
of multiple primary lesions \(p = 0.0299\), mucinous component \(p < 0.0001\), high grade \(p = 0.0025\) and right-sided location of the tumor in the colon \(p < 0.0001\) [7]. In our study, all MSI-positive NENs \(n = 5\) were low-grade and localized to the rectum.

According to the contemporary literature, there is a clear trend that the rate of MSI detection is inversely correlated with the stage of the disease [1, 3–5]. In our study, all MSI-positive NENs of the colon were stage I cancers. A larger patient sample with adequate representation of every stage of the disease is needed to check this trend for NENs of the colon.

The analysis of microsatellite instability patterns and frequency of occurrence in well-differentiated (G1/G2) neuroendocrine pancreatic tumors studied in our previous publication and NENs of the colon reveals certain differences. The MSI-positive phenotype was observed in 14% of pancreatic NENs [8] vs 17% of NENs of the colon. Besides, in pancreatic NENs, the MSI-positive phenotype was not associated with MLH1 promotor methylation. By contrast, in NENs of the colon the MSI-positive phenotype was associated only with MLH1 hypermethylation. Perhaps, this epigenetic mechanism typical for adenocarcinomas and NENs of the colon is related to their colorectal origin.

Cellular differentiation is a very important criteria in the context of gastrointestinal NENs, which is reflected in the 2019 updated WHO classification. High- and low-grade NENs are different groups of tumors completely heterogenous in terms of their genetic characteristics. Besides, NENs originating in different organs differ in their basic molecular markers. This raises the question of whether there is a link between the degree of cellular differentiation in NENs (NETs and NECs), primary tumor location and MSI pattern distribution. The data provided in the literature is controversial. For example, of 239 studied gastrointestinal and pulmonary NENs, only 4 specimens (1 G3 NETs of the pancreas and 3 NECs of the colon) were MSI-positive [9]. In another study the prevalent primary NEC location were the stomach \(n = 21\) and the pancreas \(n = 6\), and MSI was detected in none of the total 33 lesions [10]. At the same time, according to the meta-analysis of 33 retrospective studies and 8 case reports, MSI was observed in approximately 10% of gastric and colonic NECs [11]. The MSI phenotype was not confirmed in any of 56 well-differentiated NETs of the rectum \(n = 56\), small bowel \(n = 14\), colon \(n = 38\), and pancreas \(n = 16\) [9, 12–15]. According to other reports, 10–33% of pancreatic NETs are MSI-positive [9]. In our study the majority of the specimens \(55\%, n = 16\) were represented by NECs of the colon, but their MSI status was negative. However, there are reports confirming the MSI phenotype in 16%, 7%, 10%, and 14% of NECs of the colon, respectively [6, 9, 17–21]. The analysis of the relevant literature suggests that the frequency of MSI could be higher among poorly differentiated NENs. Still, our findings demonstrate the opposite: in our study, MSI was confirmed for well-differentiated grade G1 and G2 NENs of the rectum. That said, only a larger patient sample will drawing reliable conclusions about the pattern of MSI distribution in NENs of the colon.

CONCLUSIONS

The analysis of frequency of microsatellite instability in colorectal NENs depends on the tumor grade, primary location and stage
revealed that 17% of the samples were MSI-positive. All of them were stage I well-differentiated G1 and G2 NENs located in the rectum. The calculated three-year survival rates demonstrate a direct correlation between the frequency of MSI occurrence in colorectal NENs and the stage of the disease. According to the currently held research, the prevalence of MSI among NENs is similar to the prevalence of MSI among adenocarcinomas of the same organ. Comparison of our findings to the frequency of MSI in CRC reveals that well-differentiated NENs of the colon may be characterized by a higher rate of the MSI-positive phenotype (17% vs. 10–15%). Notably, testing patients with NENs for MSI is not part of the standard diagnostic protocol, which we believe is wrong. We are cautiously optimistic in suggesting that novel immunotherapies may be effective against this class of tumors.

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Литература


Development of highly sensitive methods for drug analysis is an ongoing challenge posed by modern biomedical and pharmaceutical chemistry. Drug analysis is essential to monitor the quality and purity of pharmaceuticals, study the delivery vehicles for therapeutic agents, to assess the effectiveness of the substance incorporation into the drug delivery system, to estimate the kinetic parameters of reactions, catalyzed by enzymes involved in xenobiotic metabolism, and to study the mechanisms of the drug-DNA interactions from the perspective of pharmacogenomics. The study was aimed to develop an electrochemical technique for detection of a number of drugs. The method is based on electrochemical oxidation of organic molecules at positive potentials between +0.0–1.6 V. The commercially available three-contact electrodes obtained by screen printing with unmodified graphite working electrode were used for analysis. It is shown that electrochemical technique allows for simultaneous detection of several compounds at various working electrode potentials, and for detection of drugs over a wide range of the clinically meaningful drug concentrations (50 μM–10 mM), which could be used when working with biological fluids (blood plasma, blood serum, blood, urine), as well as when performing drug monitoring and drug–drug interaction analysis.

Keywords: electroanalysis, drugs, voltammetry, unmodified screen-printed graphite electrodes

Funding: the study was carried out within the framework of the Russian Federation fundamental research program for the long-term period for 2021–2030.

Author contribution: Agafonova LE — experimental procedure, data processing, manuscript writing, building graphs; Bulko TV — sample preparation, experimental procedure; Kuzikov AV — statistical data processing, manuscript writing; Masamrekh RA — sample preparation, experimental procedure; Shumyan’tseva VV — concept, manuscript writing, data analysis.

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SENSORS FOR ANALYSIS OF DRUGS, DRUG-DRUG INTERACTIONS, AND CATALYTIC ACTIVITY OF ENZYMES

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Development of highly sensitive methods for drug analysis is an ongoing challenge posed by modern biomedical and pharmaceutical chemistry. Drug analysis is essential to monitor the quality and purity of pharmaceuticals, study the delivery vehicles for therapeutic agents, to assess the effectiveness of the substance incorporation into the drug delivery system, to estimate the kinetic parameters of reactions, catalyzed by enzymes involved in xenobiotic metabolism, and to study the mechanisms of the drug-DNA interactions from the perspective of pharmacogenomics. The study was aimed to develop an electrochemical technique for detection of a number of drugs. The method is based on electrochemical oxidation of organic molecules at positive potentials between +0.0–1.6 V. The commercially available three-contact electrodes obtained by screen printing with unmodified graphite working electrode were used for analysis. It is shown that electrochemical technique allows for simultaneous detection of several compounds at various working electrode potentials, and for detection of drugs over a wide range of the clinically meaningful drug concentrations (50 μM–10 mM), which could be used when working with biological fluids (blood plasma, blood serum, blood, urine), as well as when performing drug monitoring and drug–drug interaction analysis.

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CENSORS ДЛЯ АНАЛИЗА ЛЕКАРСТВЕННЫХ ПРЕПАРАТОВ, МЕЖЛЕКАРСТВЕННЫХ ВЗАИМОДЕЙСТВИЙ И КАТАЛИТИЧЕСКОЙ АКТИВНОСТИ ФЕРМЕНТОВ

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Разработка высокочувствительных методов анализа лекарственных препаратов является актуальной задачей современной биоаналитической и фармакологической химии. Анализ лекарственных препаратов необходим для мониторинга качества и чистоты, для исследования средств доставки терапевтических средств и определения эффективности включения субстанций в системы доставки, для исследования кинетических параметров реакций, катализируемых ферментами метаболизма ксенобиотиков, для исследования механизма взаимодействия лекарств с ДНК с позиции фармакогеномики. Цель работы было разработать электрохимический метод регистрации ряда лекарственных препаратов. Метод основан на реакции электроокисления органических молекул при положительных значениях потенциалов в диапазоне +0.0–1.6 V. Для анализа использовали коммерчески доступные трехконтактные электроды, получаемые методом трафаретной печати с немодифицированным графитовым рабочим электродом. Показано, что электрохимический метод позволяет одновременно детектировать несколько соединений при разных значениях рабочих потенциалов и для работы с биологическими жидкостями (плазмой, сывороткой, кровью, мочой), для лекарственного мониторинга и анализа межлекарственных взаимодействий.

Ключевые слова: электроанализ, лекарственные препараты, вольтарен, нурофен, парацетамол, немодифицированные печатные графитовые электроды

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Electrochemical method of analysis is a powerful tool to estimate the drug content and purity, as well as the drug concentration both in pharmaceutical fluids and in biological fluids or tissues (urine, blood serum, blood plasma, whole blood, cell lysates). Despite the use of various drug evaluation methods (such as spectrophotometry, colorimetry, fluorescence spectroscopy, gas chromatography–mass spectrometry, high-performance liquid chromatography (HPLC), thin-layer chromatography, titrimetry, capillary electrophoresis, high-performance liquid chromatography–tandem mass spectrometry and thermogravimetric analysis, radiometry, immunoassay) [1], electrochemical techniques are in demand as well due to high sensitivity, unique electrochemical signatures of the relevant compounds, reasonable cost, fast electrochemical analysis speed, low sample volumes (2–60 μL), and portable equipment. Electroanalysis allows for simultaneous detection of several compounds at various working electrode potentials, and for detection of drugs over a wide range of clinically meaningful drug concentrations (50 μM–10 mM), which could be used when working with biological fluids (blood plasma, blood serum, blood, urine), as well as when performing drug monitoring and drug–drug interaction analysis.

Keywords: electroanalysis, drugs, voltammetry, unmodified screen-printed graphite electrodes

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of several pharmaceuticals and makes it possible to assess these drugs by various electrochemical methods in order to improve the test sensitivity (cyclic and stripping voltammetry, square-wave voltammetry, differential pulse voltammetry, chronocoulometry, and electrochemical impedance spectroscopy) [2–6].

Cytochrome P450 is a superfamily of heme-thiolate monoxygenases involved in metabolism of xenobiotics and endogenous compounds [7]. The method for measuring catalytic activity of this hemoprotein class by electrochemical oxidation of substrate drugs performed well in the previously published study [4], aimed at measuring the catalytic activity of cytochrome P450 3A4. Cytochrome P450 19A1 (CYP19A1, aromatase) is a key enzyme for estrogen biosynthesis [8]. Electrochemical methods for determination of estrone or β-estradiol using the electrodes, modified with various nanocomposites, had been previously developed in order to quantify the products of the CYP19A1-dependent electrocatalytic reaction [9–12]. Electrochemical oxidation of estrone or β-estradiol, being the aromatase metabolites, could be also detected on the commercially available three-prong printed graphite electrodes (PGE) [13]. Electrooxidation of (S)-7-hydroxywarfarin, being the cytochrome P450 2C9 metabolite, was used to evaluate the kinetic parameters of this hemoprotein [5].

The study was aimed to develop the reagentless electrochemical method for drug identification and quantification under physiological conditions in order to provide the possibility of assessing drugs in blood serum using the commercially available screen-printed three-prong electrodes with the graphite working electrode.

METHODS

Electrochemical measurements were performed with the PGSTAT 12 Autolab and PGSTAT 312N Autolab potentiostats (Metrohm Autolab Ins.; Netherlands) and the GEPES and NOVA software, versions 4.9.7 and 2.0, respectively (Netherlands). The three-prong printed graphite electrodes (PGE, ColorElectronics; Russia) were used, together with the graphite working and auxiliary electrodes, and the silver chloride reference electrode. The diameter of working electrode was 0.2 cm (the area was 0.0314 cm²). All potentials were referred to the silver/silver chloride reference electrode (Ag/AgCl).

The following reagents were used in the study: monobasic potassium phosphate (Reachem; Russia), sodium chloride (Reachem; Russia), diclofenac sodium (substance, Sigma-Aldrich; India), ibuprofen sodium (substance, Sigma-Aldrich; India), acetaminophen (pharmaceutical dosage form, Pharmstandard; Russia), mexidol (pharmaceutical dosage form, Pharmasoft; Russia), serum (S 1005-14, UsBiological; USA).

The 10 mM stock solutions were prepared in the 0.1 M potassium phosphate buffer (pH 7.4), containing 0.05 M NaCl, the desired concentrations were obtained by diluting with buffer and stored at +4 °C.

The real-time measurements were performed by differential pulse voltammetry (DPV) over the potential range of (0–1.8) V with the 0.01 V potential step and 25 Hz frequency, and by cyclic voltammetry (CV). Experiments were performed under aerobic conditions at room temperature. A total of 60 μL (volume essential for uniform distribution of the drop across the electrodes) of the drug solution to be analyzed were applied to the surface of the disposable PGE, covering the working electrode, auxiliary electrode, and reference electrode [14]. At least three electrodes were used to assess the results repeatability for each concentration.

Peak current of analyte oxidation was plotted against analyte concentration in order to calculate sensitivity and the detection limit. The resulting calibration dependencies were used to calculate sensitivity (equation 1) and the detection limit (equation 2) [15]:

\[ S = \frac{\Delta I}{\Delta C} \]  
\[ Cl_{lim} = \frac{3.3}{S} \]

where S — sensitivity, I — current, C — drug concentration, Cl — detection limit, σ — residual standard deviation (standard deviation of the regression coefficient b).

To remove protein components from blood serum, 2.5 mL of blood serum were collected in the 10 mL glass tube and
added 2.5 mL of the 15% (w/v) acetonitrile solution of zinc sulfate (50/40, v/v) or 10% trichloroacetic acid (1:10). The test tube was shaken for 20 min and equilibrated at 4 °C for 15 min, then the solution was centrifuged at 13,500 rpm for 5 min. Subsequently, supernatant was discarded, and the solution was used for further analysis [16]. After protein precipitation, blood serum was diluted 10 times with the 0.1 M potassium phosphate buffer, containing 50 mM NaCl, pH 7.4.

RESULTS

Rational selection of the electrode type is the key point of electrochemical analysis, which is essential for the most effective electron transfer and detection of molecules, biochemical events, and catalytic current, being the indicator of electrocatalysis [17]. It is shown, that modification of the electrode working surface with nanomaterials (carbon nanotubes, graphene, graphene oxide, metal nanoparticles) contributes to the increased sensor analytical sensitivity [6]. However, modified PGE may acquire background characteristics that impede direct registration of electrooxidation/electroreduction [18]. In unmodified PGE, background characteristics in the electrolyte buffer show no unintended additional signals, and the current values registered are rather low [5]. Furthermore, unmodified electrodes are commercially available, reproducibility of electrochemical analysis is high, which constitutes an essential element of the further sensors’ practical utilization for drug analysis and purity evaluation, particularly in the clinical diagnostics laboratories. In this regard, we have developed methods for drug detection using precisely the unmodified graphite working electrode surfaces.

Table. Electroanalytical characteristics of drugs, obtained by differential pulse voltammetry on PGE in the 0.1 M potassium phosphate buffer containing 50 mM NaCl, pH 7.4 (n = 4–5; p = 0.95)

<table>
<thead>
<tr>
<th>Pharmaceutical (manufacturer)</th>
<th>Linear concentration range, M</th>
<th>Eα, V</th>
<th>Regression equation</th>
<th>R²</th>
<th>Sensitivity, A/M</th>
<th>Detection limit, M</th>
</tr>
</thead>
<tbody>
<tr>
<td>Drug A (Pharmstandard)</td>
<td>5 × 10⁻⁵ – 1 × 10⁻³</td>
<td>0.50 ± 0.03</td>
<td>(8.6 ± 0.5) × 10⁻⁵</td>
<td>0.985</td>
<td>8.6 × 10⁻⁴</td>
<td>2.9 × 10⁻⁵</td>
</tr>
<tr>
<td>Drug I (Sigma-Aldrich)</td>
<td>5 × 10⁻⁵ – 1 × 10⁻³</td>
<td>1.29 ± 0.02</td>
<td>(1.2 ± 0.1) × 10⁻⁴</td>
<td>0.983</td>
<td>1.2 × 10⁻⁴</td>
<td>4.1 × 10⁻⁵</td>
</tr>
<tr>
<td>Drug D (Sigma-Aldrich)</td>
<td>5 × 10⁻⁴ – 5 × 10⁻⁴</td>
<td>0.57 ± 0.05</td>
<td>(4.4 ± 0.5) × 10⁻⁴</td>
<td>0.966</td>
<td>4.4 × 10⁻⁴</td>
<td>4.6 × 10⁻⁵</td>
</tr>
</tbody>
</table>

Electroanalytical characteristics of pharmaceuticals

N-acetyl-para-aminophenol, or drug A is an antipyretic and analgesic agent commonly used in patients with mild to moderate pain or to lower the body temperature, including in viral and bacterial infections [3]. It has been shown that N-acetyl-p-benzoquinone imine (NAPQI) is the main product of the drug A oxidation [3]. The mechanism, underlying the analgesic effect, is associated with prostaglandin synthesis inhibition in the central nervous system [2, 3, 19].

Fig. 1 provides the differential pulse voltammogram of the 1 mM drug A acquired on the unmodified PGE in the potential range of (0–1.2) V. Under aerobic conditions, drug A oxidation occurs at the Eox potential of (0.50 ± 0.03) V (vs. Ag/AgCl). The figure insert shows the linear growth of the drug A oxidation peak current with the concentration increase from 0.05 mM to 1.00 mM. The drug A oxidation potential is stable within the margin of error.

Table provides electroanalytical characteristics (concentration range, oxidation potential), equation for the relationship between the oxidation peak current and the concentration, R² coefficient of determination, as well as the electrochemical system sensitivity, calculated using equation 1, and the detection limit of drug A, calculated using equation 2.

(RS)-2-(4-isobutylphenyl)-propionic acid, or drug I is a non-steroidal anti-inflammatory drug from the group of the propionic acid derivatives, which possesses analgesic and antipyretic activity [12, 13]. Fig. 2 provides the drug I differential pulse voltammograms acquired on PGE in the potential range of (0.6—1.6) V and the concentration range of (0.5–10) mM. Electrochemical oxidation of the 1 mM drug I occurs at high potential values (1.29 ± 0.02) V (see Table).
(2-(2,6-dichloroanilino)-phenylacetic acid), or drug D is a non-steroidal anti-inflammatory drug from the group of the phenylacetic acid derivatives. The sodium salt is used in the pharmaceutical dosage forms. The drug has multiple trading names, it is prescribed to patients with many disorders, such as rheumatoid arthritis, osteoarthritis, various inflammation conditions [20, 21]. The drug has analgesic, anti-inflammatory, and anticancer effects [21]. Drug D undergoes extensive metabolism, mediated by glucuronosyltransferase, to form diclofenac acyl glucuronide. On exposure to cytochrome P450, drug D undergoes oxidative metabolism to form 4'-hydroxydiclofenac (catalyzed by cytochrome P450 2C9) and 5-hydroxydiclofenac (catalyzed by cytochrome P450 3A4) [22, 23].

In some cases, the drug can cause unwanted side effects: stomach hemorrhage, high blood pressure in patients with Shy–Drager syndrome and diabetes mellitus. The long-term use may result in infarction or stroke [20]. That is why the drug D analysis remains an ongoing challenge posed by bioanalytical and pharmaceutical chemistry.

Fig. 3 provides the drug D differential pulse voltammograms acquired on PGE in the potential range of (0.2 – 1.0) V and the concentration range of (50 – 500) μM. The oxidation potential of (0.57 ± 0.05) V was stable within the margin of error in the studied concentration range. The linear relationship was found between the oxidation peak current and the drug D concentration (see Table).

**DISCUSSION**

CV in the scan rate range of (0.05–0.18) V/s was used to characterize the drug electrochemical oxidation processes. The CVV results showed the linear relationship between the oxidation peak currents of the 1 mM drug A, 100 mM drug D and 5 mM drug I, and the scan rate square root ν₁/2 (Fig. 2 (a), 4 (a), 6 (a), Appendix), reflecting the diffusion-controlled electrochemical oxidation of the drugs on the unmodified PGE in the studied range of the potential scan rates [24]. There was also a linear relationship between the oxidation peak potentials and the logarithm of scan rate, log ν (Fig. 2 (b), 4(b), 6(b), Appendix), typical for irreversible electrochemical processes. This is confirmed by the CV technique: only the oxidation peaks of the 1 mM drug A, 100 mM drug D and 5 mM drug I are observed on the unmodified PGE (Fig. 1, 3 and 5, Appendix), suggesting the irreversible nature of electrochemical reactions. These results are in line with the mechanisms of the studied drugs electrochemical oxidation [2, 3, 6, 19–22, 25–30].

Polypharmacy, i.e. prescribing several medications to one patient, is widely used in clinical pharmacology and therapeutics. Pharmacokinetic and pharmacodynamic parameters should be monitored in multimorbid patients (prescribed two or more medications). Electroanalysis based on the drug electrochemical oxidation registration allows for simultaneous detection of several compounds. This approach can be used to determine the drugs’ mutual influence. Thus, antioxidant metabolic drug methyl 2-ethyl-6-methyl-3-hydroxyprydâne succinate, drug M, which is prescribed to improve cerebral circulation, is commonly used in various groups of patients with comorbidities. PGE were used to analyze the drug combinations D + M (Fig. 4), D + I (Fig. 5). Electrochemical oxidation of drug M is detected at the potential of +0.67 ± 0.06 V (Fig. 4). However, drug M (98 μM) has virtually no effect on the drug D (100 μM) electrochemical oxidation. There is a slight shift of the drug D oxidation potential by (5 ± 2) mV toward the more positive (anodic) area, which is indicative of the more complicated electrochemical oxidation process. The oxidation peak current is slightly reduced (± 6%), which allows for the drug D quantification in the presence of drug M (Fig. 4). Fig. 5 provides DPV for the combination of the 25 μM drug D and 200 μM drug I acquired on the PGE. The measurements have been performed over the potential range of (0–1.6) V. Electrooxidation peaks of the drugs D (E_{ox} = +0.54 ± 0.02 V) and I (E_{ox} = +1.29 ± 0.02 V) are clearly distinguished, however, the concentration dependence remains.

Drug D was also determined by electrochemical oxidation in the blood serum samples. The concentration of drug D added to blood serum was 100 μM, and the oxidation peak current recorded was 7.5 × 10^{-8} A (Fig. 3). According to the insert in the Fig. 3 and the calibration dependence (see Table), the drug D concentration calculated using the regression equation was (95 ± 5) μM.

**CONCLUSIONS**

The method for drug analysis using the commercially available disposable unmodified screen-printed electrodes has been developed. Sensitivity of the method corresponds to the clinically relevant concentration ranges. Electrooxidation of pharmaceuticals using electrodes as measuring tools is an effective analytical approach to the drug purity assessment and drug concentration determination, particularly in biological fluids used for therapeutic drug monitoring. The methods for drug quantification based on electrochemical oxidation may be used to assess catalytic activity of enzymes involved in the phase I metabolism, such as cytochrome P450, flavin-containing monooxygenases, aldehyde oxidase, aldehyde dehydrogenase, alcohol dehydrogenase, carboxylesterase, and to assess activity of enzymes involved in the phase II metabolism, such as UDP-glucuronosyltransferases, sulfotransferase, glutathione transferase. The method can be also used for registration of the comparative kinetics of the polymorphic variants and genetically engineered mutant enzymes in order to develop the biocatalyst isoforms important in technology.
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Parkinson’s disease (PD) and essential tremor (ET) are common movement disorders that predominantly affect the elderly [1, 2]. Both diagnoses are clinical and rely on the sum of their typical neurological manifestations. According to the criteria for PD published by the International Parkinson and Movement Disorder Society in 2015, bradykinesia combined with resting tremor and/or rigidity in the presence of supportive criteria and the absence of absolute exclusion criteria indicates clinically definite or clinically probable PD [3]. Importantly, apart from motor manifestations, the clinical picture of PD can include non-motor symptoms that predate motor impairment and progress gradually as the disease advances [4].

According to the updated criteria proposed by the International Parkinson and Movement Disorder Society in 2017, ET is defined as “an isolated tremor syndrome of bilateral upper limb action tremor with at least 3 years’ duration, with or without tremor in other locations” [5]. In practice, patients with ET often present with additional neurological symptoms that go beyond the definition of ET, including resting tremor, impaired tandem gait, etc. Such cases are classified as ET plus.

VISUAL ANALYSIS OF NIGROSOME-1 IN THE DIFFERENTIAL DIAGNOSIS OF PARKINSON’S DISEASE AND ESSENTIAL TREMOR

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Differentiation between Parkinson’s disease, especially in its early stages, and essential tremor, which is a phenotypically similar movement disorder, still remains an unsolved challenge for neurology. The aim of this study was to assess the diagnostic significance of nigrosome imaging (nigrosomes are dopaminergic neuron clusters in the substantia nigra of the midbrain) using 3T high-resolution SWI-MRI. The study was conducted in 20 patients with Parkinson’s disease and 10 patients with essential tremor. Visual analysis of the acquired nigrosome-1 images was performed using a 4-point ordinal rating scale. Differences in sex, age and duration of the disease were calculated using the Fisher exact test and the Mann–Whitney U test. The diagnostic value of the method was assessed using Pearson’s chi-squared test. Nigrosome-1 was bilaterally or unilaterally absent in 70% of parkinsonian patients. By contrast, nigrosome-1 was bilaterally intact in all patients (100%) with essential tremor (p < 0.001). Our preliminary findings demonstrate the high potential of noninvasive nigrosome-1 imaging in the differential diagnosis of Parkinson’s disease and essential tremor.

Keywords: Parkinson’s disease, essential tremor, nigrosome-1, magnetic resonance imaging, SWI

Author contribution: Moksalenko AN — clinical assessment, data acquisition and interpretation, literature analysis, manuscript preparation; Filatov AS — data analysis and interpretation, manuscript preparation; Konovolov RN — data analysis and interpretation, study planning and supervision; Fedotova EYu, Illarioshkin SN — study planning and supervision.

Compliance with ethical standards: The study was approved by the Ethics Committee of the Research Center of Neurology (Protocol № 2–5/20 dated March 18, 2020). Informed consent was obtained from all study participants.

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BILOGICAL ANALYSIS OF NIGROSOMES-1 IN THE DIFFERENTIAL DIAGNOSIS OF PARKINSON’S DISEASE AND ESSENTIAL TREMOR

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Дифференциальная диагностика болезни Паркинсона и фенотипически схожего двигательного расстройства — эссенциального тремора, особенно в дебюте заболевания, остается одной из нерешенных задач современной неврологии. Целью исследования было оценить диагностическую значимость визуализации нигросом — кластеров дофаминергических нейронов в черной субстанции (ЧС) среднего мозга, выявляемых при использовании SWI-режима высокоразрешающей магнитно-резонансной томографии (3 Тесла). У 20 пациентов с болезнью Паркинсона и у 10 пациентов с эссенциальным тремором. Визуальный анализ изображений нигросомы-1 проводили с использованием четырехчленной порядковой шкалы. Различия по гендерному, возрастному составу и продолжительности заболевания рассчитывали с помощью точного критерия Фишера, U-критерия Манна–Уитни. Для расчета диагностической ценности данной методики использовали критерий χ² Пирсона. У пациентов с болезнью Паркинсона в 70% случаев наблюдали одно- или двустороннее исчезновение нигросомы-1. Еще у двух пациентов с болезнью Паркинсона (10%) выявили менее специфические изменения черной субстанции — уменьшение объема нигросомы-1. Напротив, у всех пациентов с эссенциальным тремором (100%) нигросома-1 оставалась сохранной с двух сторон (p < 0.001). Полученные предварительные результаты демонстрируют высокий потенциал методики визуального анализа нигросомы-1 в дифференциальной диагностике болезни Паркинсона и эссенциального тремора.

Ключевые слова: болезнь Паркинсона, эссенциальный тремор, нигросома-1, магнитно-резонансная томография, SWI-режим

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manifestations, many patients with ET have various non-motor symptoms [2] that usually do not have any particular clinical significance but complicate differentiation between ET and PD.

Radionuclide imaging, e.g. positron-emission tomography (PET), single photon emission computed tomography (SPECT) and transcranial sonography (TCS), can be used to differentiate between ET and PD by assessing damage to the substantia nigra (SN), the primary target of neurodegeneration in PD, which remains intact in ET [6, 7]. However, radionuclide imaging has objective limitations impeding its exploitation in clinical neurological practice.

The use of magnetic resonance imaging (MRI) for diagnosing PD and differentiating it from nondeenerative forms of parkinsonism became possible with the spread of high-field MR scanners and the introduction of additional MRI sequences into the standard MRI protocol.

Dopaminergic neurons of SN are arranged into cell clusters called nigrosomes [8]. Nigrosome-1, the largest of 5 known nigrosomes, appears on high-resolution susceptibility weighted images (SWI) as an oval slightly hyperintense region in the dorsal SN. Nigrosome-1 divides SN into 2 parts, bearing resemblance to a swallow tail, hence its name “the swallow tail sign” [9]. Recent research has shown that location of the hyperintense nigrosome-1 region in the surrounding hypointense SN structures can be quite variable and does not always fit the “swallow tail” profile [10]. Patients with PD demonstrate a loss of dorsolateral nigral hyperintensity due to the involvement of nigrosome-1 in neurodegeneration [9, 11]. In ET, structural and functional changes have been reported in the cerebellum and the brain stem (predominantly in the locus coeruleus) [12]. Despite the lack of consistency between the results of pathomorphological studies and the understudied pathogenesis of ET, so far there has been no reliable evidence about the presence of pronounced SN degeneration in patients with ET comparable to that in patients with PD. Consequently, attempts have been made to determine the diagnostic significance of visual assessment of nigrosome-1 images in discriminating between PD and ET. The method has demonstrated high sensitivity and high specificity; besides, it does not require image post-processing and therefore is effective and suitable for clinical practice [13, 14].

To our knowledge, there are no publications analyzing the described neuroimaging pattern of SN changes in the Russian cohort of patients with movement disorders. The aim of this study was to assess the biomarker role of dorsolateral nigral hyperintensity loss in differentiating between PD and ET, which is a phenotypically similar disorder.

METHODS

Participants

Participants were recruited from in- and outpatients undergoing treatment at the Research Center of Neurology from January to October 2020. The study included 20 patients with tremor-inducing metabolic disorders; structural damage to the brain (neoplasms, infarction, brain injury sequelae); MRI artifacts precluding the analysis of MR images; age under 18 and above 80 years.

MRI protocol and analysis of MR images

MRI protocol

All MR images were acquired using a 3T Siemens MAGNETOM Verio scanner equipped with an 8-channel head coil. SWI sequences were acquired to assess nigrosome-1 appearance (TR = 27 ms, TE = 20 ms, slice thickness = 1.5 mm, dist. factor = 20%, FoV = 172 × 230 mm², scan time = 2 min 59 s). Besides, T2, T1 MPR, T2 FLAIR and DWI images were acquired to exclude other causes of parkinsonism. The axial plane was parallel to the line connecting the anterior and posterior commissures across all brain structures.

Qualitative analysis of acquired images

On the acquired SW images, nigrosome-1 appeared as an oval slightly hyperintense region in the hypointense area of the dorsal midbrain (SN). Visual analysis of the images was performed using the following 4-point ordinal scale: 0 points — the norm (nigrosome-1 is visualized bilaterally); 1 point — the image has no diagnostic value (nigrosome-1 is poorly visualized on one or both sides or is diminished in size, i.e. partially lost); 2 points — abnormality (nigrosome-1 is absent unilaterally); 3 points — abnormality (nigrosome-1 is absent bilaterally). For illustrative purposes, MR images of 4 patients with different nigrosome-1 appearance are provided in Fig. 1. Qualitative analysis was conducted by 2 radiologists who had no access to the patients’ medical records and were working independently. If their conclusions were conflicting, preference was given to the opinion of the more experienced radiologists.

Statistical analysis

The results of the study are presented below as medians and lower and upper quartiles (Med, lq, uq). Demographic characteristics of the patients (age, sex, duration of the disease) were compared using the Fisher exact test and the Mann–Whitney U-test. Nigrosome-1 scores were compared between the groups using Pearson’s chi squared test. In all statistical tests, the significance threshold was assumed to be $p < 0.05$. The data were analyzed in StatTech v1.1.0, SPSS Statistics.

RESULTS

Demographic characteristics

The PD and ET groups did not differ significantly in terms of sex and age ($p = 0.246, p = 0.082$, respectively). The duration of the disease was significantly longer in the patients with ET than in those with PD ($p < 0.003$). The analysis of associations between the disease and sex was performed using Fisher’s exact test; the associations between age and disease duration were tested using the Mann–Whitney U test. Demographic characteristics of the patients are provided in Table.

Neuroimaging data

Nigrosome-1 was clearly visible bilaterally in all patients with ET ($n = 10$), so all patients from group 2 scored 0 points on the rating scale (100%).
Fig. 1. Susceptibility-weighted MR images in the axial plane; slices pass through the cerebral peduncles. The substantia nigra is hypointense on SWI sequences, whereas nigrosomes-1 (yellow circles) are hyperintense. The figure shows different patterns of nigrosome-1 appearance in patients with PD (A — 0 points, B — 1 point (nigrosome-1 is reduced in volume on the left), C — 2 points, D — 3 points).

However, it was absent in 70% of patients with PD (n = 14); the ratio of unilateral and bilateral loss of dorsolateral nigral intensity was 1:1. Accordingly, 7 patients with PD scored 2 points (35%) and 7 other patients with PD scored 3 points (35%).

Nigrosome-1 was intact (0 points) in 4 patients with PD (20%); 2 more patients with PD (10%) scored 1 point: their MRI scans showed a reduction in nigrosome-1 size on one side, which was interpreted as having no diagnostic value. Comparison of the PD and ET groups demonstrated a significant difference in the results expressed as percentage (p < 0.001, Pearson’s χ²).

Thus, the study demonstrates a high diagnostic value of non-invasive visual nigrosome-1 assessment in differentiating between PD and ET: the sensitivity and specificity of the method were 70% and 100%, respectively. The results are provided in Fig. 2.

DISCUSSION

Oftentimes, discrimination between early-stage PD and phenotypically similar disorders poses a certain difficulty to a neurologist. The aim of this study was to assess the diagnostic significance of non-invasive nigrosome-1 assessment in differentiating PD from ET.

It has been over 20 years since heterogeneity of the SN pars compacta (i.e. identification of nigrosomes and the nigral matrix by immunohistochemical staining) was discovered and the staging of nigrosome damage due to PD-related neurodegeneration was pathomorphologically confirmed [8, 15]. Non-invasive imaging of nigrosome-1 became possible with the spread of high-field MR scanners and the introduction of SWI sequences into the standard brain MRI protocol [9, 16]. SWI is a technique that utilizes 3D pulse MRI sequences sensitive to magnetic field inhomogeneities. It is based on the following phenomenon: iron, calcium and deoxyhemoglobin can enhance a local magnetic field and induce a positive phase shift, in comparison with the surrounding cerebral tissues. Tissues containing these paramagnetic agents appear on SW images as regions of hypointense MR signal [17, 18].

In healthy subjects, SN appears on MR images as a hypointense midbrain region dorsally divided into 2 segments by an oval hypointense area. Histopathological studies have confirmed that this dorsolateral nigral hyperintensity corresponds to nigrosome-1 and that signal enhancement may be associated with low iron content in this region in comparison with the surrounding SN [19].

Nigrosome-1 is not visualized in patients with PD. Apart from the loss of dopaminergic neurons, this may be associated with iron accumulation occurring in parallel [20, 21]. The loss of dorsolateral nigral hyperintensity is currently regarded as one of the most promising biomarkers of PD. For example, a recent meta-analysis reports that the diagnostic accuracy of nigrosome-1 imaging for the differentiation between patients with idiopathic PD and healthy individuals demonstrates high sensitivity and high specificity [22].

<table>
<thead>
<tr>
<th>Table. Demographic characteristics of the patients</th>
<th>PD</th>
<th>ET</th>
<th>p</th>
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<tr>
<td><strong>Sex, abs. (%)</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Women</td>
<td>11 (55)</td>
<td>8 (80)</td>
<td>0.246</td>
</tr>
<tr>
<td>Men</td>
<td>9 (45)</td>
<td>2 (20)</td>
<td></td>
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<tr>
<td><strong>Age, Me [Q₁–Q₃]</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Full years</td>
<td>60 [52.25–66.5]</td>
<td>73.5 [58.5–77.25]</td>
<td>0.082</td>
</tr>
<tr>
<td><strong>Disease duration, Me [Q₁–Q₃]</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Full years</td>
<td>3 [2–8.5]</td>
<td>10 [8.5–15.75]</td>
<td>0.003</td>
</tr>
</tbody>
</table>
The diagnostic value of this neuroimaging marker in differentiating between PD and ET was assessed in two studies published in 2019. Jin L. et al. analyzed MR images of 68 patients with PD, 25 patients with ET and 34 control subjects. The method demonstrated high sensitivity (79.4%) and high specificity (92.0%) [13]. M. S. Perez Akly et al. studied dorsolateral nigral hyperintensity in 16 patients with PD and 16 patients with ET. The results were comparable to the results of the study by Jin L. et al. According to one of 2 involved radiologists, the sensitivity and specificity of the method were 93.75% and 87.5%, respectively. The second radiologist reported 93.75% sensitivity and 75% specificity [14]. Thus, the method was shown to be effective in differentiating between PD and ET by 2 independent research teams.

Our study also confirms the diagnostic value of noninvasive nigrosome-1 imaging. In contrast with ET patients, the absence of dorsolateral nigral hyperintensity in SN was observed in the majority of our PD patients. The sensitivity and specificity of the method tested on the small cohort of patients were 70% and 100%, respectively.

Artifacts from motion and metal dental implants were a significant limitation of our study. From initially examined 39 patients (26 patients with PD, 13 patients with ET), only 30 whose MR images were suitable for the analysis were included in the study. To reduce the number of artifacts from motion, the patient’s head can be stabilized with sand sacks or foam pillow support, and mild medical sedation can be applied [16].

CONCLUSIONS

Our findings supported by the results of foreign studies lead us to conclude that noninvasive neuroimaging has a potential to become a useful tool in the differential diagnosis of diseases accompanied by tremor and other movement disorders, including differentiation between PD and ET, especially in the early stages of the disease.

References

Литература


MEDIUM-TERM OUTCOMES OF EXTRAARTICULAR CORRECTIVE OSTEOTOMY FOR SLIPPED CAPITAL FEMORAL EPIPHYSIS

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Despite the diversity of surgical options for slipped capital femoral epiphysis (SCFE), there is an ongoing search for the technique that would ensure a satisfactory outcome, stable fixation of bone fragments and a low rate of complications. The aim of this study was to improve the surgical technique for SCFE in patients with moderate and severe SCFE. The study included 52 children (16 girls and 36 boys) aged 10–15 years (the mean age was 13.2 years) with chronic severe (Krechmar’s stage III) stable (according to Loder’s classification) SCFE. The control group (n = 16) underwent a classic Imhauser procedure; the main group (n = 36) underwent a tritrapezoidal osteotomy proposed by the authors of the study. The patients were examined prior to surgery and in the late follow-up period (the mean follow-up time was 4.7 years, ranging from 1 to 10 years). The procedure included a clinical examination, history taking, radiography to measure the slip angle and the severity of the slip, and the Harris hip score to assess hip function. After 4.7 years, both groups demonstrated an increase in the range of motion, in comparison with their preoperative results (p < 0.05), good Harris hip scores (94 points in the main group and 81 points in the control group. Postoperative radiographs showed consolidation of the bone, recovery of the proximal femur anatomy. Leg length discrepancy improved significantly in both groups. The proposed technique for extraarticular osteotomy allows recovering the length of the affected leg, the anatomy and physiology of the hip joint, is simple and less traumatic.

Keywords: slipped capital femoral epiphysis, corrective extraarticular femoral osteotomy, hip joint, Imhauser procedure

Author contribution: all authors contributed equally to the study and the manuscript, all read and approved the final version of the manuscript.

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Slipped capital femoral epiphysis (SCFE) is a relatively rare, predominantly juvenile disorder [1]. Due to a variety of causes, including endocrine, the osseous tissue of the metaphysis at the epiphyseal-metaphyseal junction undergoes structural transformation resulting in the disruption of the osteoclast/osteoblast balance and accompanied by the spatial arrangement of the extracellular elements of connective tissue. The bone resorbs, and the epiphysis slips out of its normal position [2, 3]. The typical underlying mechanism of SCFE is associated with high axial load and is characterized by the posterior-inferior displacement of the epiphysis and its retroversion [4].

СРЕДНЕСРОЧНЫЕ РЕЗУЛЬТАТЫ ВНЕСУСТАВНОЙ КОРРИГИРУЮЩЕЙ ОСТЕОТОМИИ БЕДРА ПРИ ЮНОШЕСКОМ ЭПИФИЗЕОЛИЗЕ ГОЛОВКИ БЕДРЕННОЙ КОСТИ

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Несмотря на множество предложенных методов хирургического лечения юношеского эпифизеолиза головки бедренной кости ЮЭГБК, продолжаются поиски вариантов операций, обеспечивающих устойчивое положение, стабильную фиксацию костных фрагментов и нормальный уровень осложнений. Поэтому была разработана техника лечения пациентов с ЮЭГБК среднетяжелой степени. В исследование вошли 52 ребенка в возрасте 10–15 лет (средний возраст 13 лет), из них 16 девочек и 36 мальчиков, страдающие ЮЭГБК тяжелой степени (3-я стадия по классификации Кричмара, хроническое течение, стабильного типа (классификация Лодера). На контрольной группе (n = 16) выполнена стандартная операция по Имхаузеру, на исследуемой (n = 36) — авторская трехплоскостная остеотомия. Пациенты были обследованы до операции и в отдаленные сроки (средний срок наблюдения составил 4.7 года (от 1 до 10 лет) с помощью клинического метода (сбор анамнеза, объективное исследование), рентгенологического метода (определение степени соскальзывания и угла соскальзывания), а также опросника функционального состояния (Harris hip score). В среднем через 4.7 года в обеих группах отмечено увеличение объема движений в сравнении с дооперационными показателями (p ≤ 0.05), хорошие функциональные показатели HHS (в исследуемой группе — 94 балла, в контрольной — 81 балл); на контрольных рентгенограммах отмечена консолидация костных фрагментов с сохранением коррекции проксимального отдела бедра, длина конечности также восстановилась в обеих группах. Предложенная клиническая остеотомия позволяет восстановить длину конечности, анатомо-физиологические взаимоотношения в тазобедренном суставе, проста в исполнении и менее травматична.

Ключевые слова: юношеский эпифизеолиз головки бедренной кости, внесуставная корригирующая остеотомия бедра, тазобедренный сустав, некоррекция бедра, операция Имхуазер

Вклад авторов: все авторы внесли существенный вклад в проведение исследования и подготовку статьи, прочли и одобрили финальную версию перед публикацией.

Соблюдение этических стандартов: исследование одобрено этическим комитетом РНИМУ им. Н. И. Пирогова (протокол № 213 от 13 декабря 2021 г.) выполнено в соответствии с этическими стандартами Хельсинской декларации; родители пациентов дали согласие на обработку и публикацию их персональных данных.

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The disease occurs in 4–5 individuals per 100,000 population. It is more likely to affect pubescent boys aged 12–13 years (the male to female ratio is 3 : 2). Clinically, it presents as outer thigh pain or pain in the hip and knee joints (as a rule, pain in the knee joint is more common), which often urges the doctor to search for a possible knee joint pathology and thus misdiagnose the patient, because the true cause of pain, i.e. slipped proximal femoral epiphysis, remains overlooked.

There are a few classifications of SCFE used in clinical practice. The Southwick Slip Angle Classification is based on the epiphysis–metaphyseal angle and categorizes the degree of epiphyseal displacement as mild (0–30°), moderate (30°–50°) and severe (> 50°). The Loder Classification evaluates epiphyseal stability and the patient’s ability to bear weight. A stable slip means that the patient is able to bear weight with or without crutches; an unstable slip means that the patient is unable to bear weight even with crutches.

Another classification was proposed by Krechmar in 1982:

Stage I: predisplacement; no signs of epiphyseal displacement, pronounced changes in the proximal physis (growth plate) and the metaphysis (femoral neck);
Stage II: the epiphysis is displaced posteriorly (< 30°) and inferiorly (< 15°); there are structural changes in the metaphysis; the proximal physis is open;
Stage III: the epiphysis is displaced posteriorly (> 30°) and inferiorly (> 15°); there are structural changes in the metaphysis; the physis is open;
Stage IV: acute postero Inferior displacement of the epiphysis; the physis is open;
Stage V: residual proximal femoral deformity with various degrees of epiphyseal displacement and the closed proximal physis.

This grading system integrates some of the abovementioned classifications and is, in our opinion, the most convenient.

On examination, patients with SCFE have a limp, the affected leg appears shorter, there is excessive external rotation of the hip, progressing over time; internal rotation of the hip is limited; reaching the full range of motion is painful. Krechmar’s stage III is characterized by the positive Hofmeister-Drehmann’s sign. The diagnosis is confirmed by anteroposterior and frog-leg lateral radiographs. In 20% of cases, SCFE is bilateral [5].

Because the condition is rare, it is often diagnosed in the advanced stage. In the majority of cases, patients with SCFE are hospitalized when the proximal femur deformity becomes very pronounced [1]. SCFE has a serious social impact, so it is important to ensure prophylaxis of early hip joint osteoarthritis, hip impingement syndrome and avascular necrosis of the epiphysis, which has been correlated with SCFE in a number of studies [6, 7].

The main goal of SCFE treatment is to prevent further femoral deformation, stabilize the proximal epiphysis and preserve blood supply to the epiphysis [8].

A diversity of surgical interventions for chronic SCFE have been proposed, including in situ fixation with pins, screws or plates, epiphysiodesis and various types of proximal femur osteotomy [9].

At present, the preferred treatment option for moderate and severe epiphyseal displacement (> 30°) is osteotomy [10]. Depending on its site, osteotomy can be classified into subcapital (the Dunn procedure, Fish cuneiform osteotomy), osteotomy conducted at the femoral neck base (Kramer intraarticular osteotomy Badama extraarticular osteotomy) and intertrochanteric (the Southwick and Imhauser procedures). The outcome is measured by assessing hip joint function, the presence of residual displacement, the adverse sequelae of the surgical intervention (impaired blood supply to the femoral head), and the simplicity of the surgical technique [11].

In theory, proximal osteotomies of the femoral neck (Dunn procedures) are the ideal tool for restoring the anatomy of the proximal femur because SCFE-related deformities arise at this particular anatomical site [12]. However, there are reports that these procedures impede blood supply to the femoral head and cause avascular necrosis of the femoral head in 10–26% of patients. Due to the high risk of avascular necrosis, it was proposed to perform osteotomy in the intertrochanteric area of the femur [13].

Southwick osteotomy is a classic surgical intervention for SCFE. It corrects the metaphyseal-diaphyseal angle and eliminates excessive external rotation of the hip; however, it does not significantly affect the position of the proximal epiphysis in the acetabulum. In the past, neither surgeons, nor patients were fully satisfied with the outcome, which gave rise to multiplane osteotomies [14].

A triplane corrective osteotomy was proposed by Krasnov AI (RU 2364365, C2). It is more pathogenetically reasonable because it simultaneously corrects deformities in the frontal, horizontal and sagittal planes; fixation is performed with an angled blade plate [15].

Among the main drawbacks of the Krasnov osteotomy are its technical difficulty and the need to cut off the greater trochanter, which results in prolonged operative time. Besides, during this type of surgery, rotation is performed according to the position of the blade, i.e. around the longitudinal axis of the metaphysis. If the epiphysis is significantly displaced posteriorly (>40°) and only slightly inferiorly, which is a common occurrence, rotation of the proximal femur around the metaphyseal axis pushes the epiphysis and the metaphysis into a valgus and varus position, respectively. This results in a hip subluxation and/or the varus deformity of the metaphysis, with the high position of the greater trochanter, leading to the dysfunction of gluteal muscles and limping [16].

There is another variant of triplane corrective osteotomy with an angled blade plate (RU 2604039, C1). Advantageously, by changing the rotational axis of the proximal femur, correction can be performed in the frontal, horizontal and sagittal planes, preventing angular deformity of the femoral diaphysis, hip subluxation and varus deformity of the metaphysis [17].

In 1966, Imhauser described an intertrochanteric osteotomy that eliminated femoral varus and metaphyseal extension and rotation. The surgery is essentially a cuneiform osteotomy involving resection of the anterior or anterolateral fragment of the bone, followed by blade plate fixation [18]. There is a wealth of publications about this procedure, indicative of its popularity. It should be noted that femoral head subluxation in the setting of excessive valgization and a deformity of the proximal femur are common complications of this intervention [19–21].

Considering all currently existing surgical treatment options, their complications, stability of epiphyseal fixation, duration of postoperative immobilization and patient outcomes, we developed an original technique for SCFE correction based on the analysis of the aforementioned procedures. The aim of this study was to improve the outcomes of SCFE treatment in children and to assess the effectiveness of the proposed technique.

METHODS

Our retrospective study included 52 children with SCFE undergoing treatment at the Children’s hospital of the Department of Traumatology, Orthopedics, and Disaster Surgery.
from 2010 to 2020. All patients underwent a clinical examination (medical history and complaints, demographic characteristics, symptoms, range of motion assessment) and radiography; pain and hip function were assessed using the Harris hip score [22].

The patients were divided into 2 groups. The control group included patients undergoing a classic intertrochanteric Imhauser osteotomy, the main group comprised patients undergoing the original variant of osteotomy proposed by the authors of this study [23].

The following inclusion criteria were applied: posterior (30°) and/or inferior (>15°) displacement of the proximal femoral epiphysis, the growth plate being open; no past history of hip surgery; the absence of technical errors during surgery.

Exclusion criteria: posterior displacement of the epiphysis by < 30° or > 75°; the closed growth plate; a past history of hip surgery; complete separation of the epiphysis from the metaphysis (acute slip).

The clinical examination included identification of complaints and assessment of gait, i.e. limping, the ability to ambulate, pain during ambulation, pain during movements, fixed external rotation of the affected hip; limited flexion, internal rotation and adduction of the leg.

Anteroposterior and frog-leg lateral radiographs of the hip were acquired before and after surgery and in the late follow-up period. Radiographic staging was done in accordance with Krechmar and Loder classifications [24, 25].

The main outcome measure was hip function assessment with the Harris hip score. This tool was developed to evaluate the outcomes of hip surgery. It includes 4 domains: pain, function, deformity, and range of motion. For each domain, the total score is calculated (the maximum total score is 100 points). The higher the score, the better the quality of life. The score over 90 points one year after surgery was interpreted as an excellent outcome, 80–90 points as good, 65–79 points as satisfactory, and below 65 points as unsatisfactory.

All patients were examined prior to surgery and in the late follow-up period. All patients underwent an osteotomy and were advised to unload the operated leg for 4–12 months. After 4–12 months, the plate was removed and the follow-up observation continued.

Statistical analysis was conducted in SPSS (IBM SPSS Statistics 22; USA), and Excel (Microsoft; USA). The significance of differences between the groups was assessed using the non-parametric Kruskal-Wallis test; correlations between two quantitative variables were measured using the Spearman rank correlation coefficient. Differences were considered significant at $p < 0.05$.

Surgical technique

In the proposed technique, the spatial position of the axis around which the proximal femur is rotated during femoral
osteotomy is different, as is the osteotomy type (Fig. 1). Fixation is performed with a Trotsenko–Nuzhdin plate. The points of entry for the jaws were planned 0.3–0.5 cm superiorto the greater trochanter growth plate and at the posterior epiphyseal displacement distance from the midline of the lateral face of the greater trochanter. The jaws of the plate must be inserted so that the angle between the line parallel to the axis of the diaphyseal part of the plate and the line parallel to the femoral diaphysis equals the angle of epiphyseal retroversion. Blade insertion channels were formed in the proximal femur. Then, high intertrochanteric osteotomy was performed and the jaws were introduced in the prepared channels. The diaphyseal part of the plate was pulled posteriorly, keeping some space between the diaphysis and the diaphyseal part of the plate, until the midlines going through the central axis of the plate and the femoral diaphysis coincided. Thus, the femoral head was recovered from its retroverted position and derotated. Then the diaphyseal part of the plate was pressed to the femoral diaphysis and the inferior displacement of the epiphysis was eliminated. The locking part of the plate was anchored to the greater trochanter; the screws were inserted in such a way that they passed outside the femoral neck. The diaphyseal part of the plate was anchored to the femur. Importantly, the maximum angle of forward rotation of the proximal femoral fragment should be 45°; rotation of over 45° is prohibited due to the risk of ischemic complications. If the epiphysis was displaced posteriorly by over 45°, residual displacement was corrected by derotating the proximal fragment, using the formula: MEA — 45° (where MEA is a metaphyseal-epiphyseal angle before surgery). If the epiphysis was displaced inferiorly, its position was corrected by valgization of the proximal fragment. To fix the Trotsenko-Nuzhdin plate, the jaws were introduced into the greater trochanter; this saves the femoral neck from injury and helps to preserve blood supply to the proximal femur.

**Postoperative rehabilitation**

Postoperative rehabilitation was different in the main and control groups. In the main group, the patients remained on bed rest for 6 months and wore an antirotation foot support. The control group remained on bed rest for 3 months after surgery and wore a spica cast. Verticalization was encouraged.
The follow-up examination performed 4.7 years (on average) after the surgical procedure revealed an improvement in internal hip rotation by an average of 16.7° (20.1° in the main group vs 9.1° in the control group; \( p = 0.0024 \)). The leg length discrepancy was compensated and was now 0.5 cm on average (0 cm in the main group vs 1 cm in the control group; \( p = 0.5 \)) (see Table).

Hip function assessment was performed only in the late follow-up period (4.7 years after the osteotomy), so it is impossible to track how hip function had been changing over that period, but the end result can be measured in both groups. The average Harris score in the late follow-up period was 89 points (94 points in the main group vs 81 points in the control group; \( p = 0.001 \) (Fig. 5).

Chondrolysis is a serious complication of surgery for SCFE and the disorder itself. Chondrolysis was observed in one patient in the main group (2.8%) and two patients in the control group (5.6%; \( p = 0.0013 \)).

**DISCUSSION**

The literature offers a diversity of surgical techniques for treating chronic SCFE. The choice of the technique depends on the stage of the disorder and the experience and skills of the operating surgeon.

Today, SCFE is usually managed with intertrochanteric and subtrochanteric osteotomies of the femur involving fixation with an angled blade plate and screws or by an osteotomy of the femoral neck with screw fixation.

The original technique for metaphyseal osteotomy proposed by the authors of this study has demonstrated good outcomes and a low rate of complications.

Demographic parameters (age and sex), the average range of motion for external rotation of the hip and leg length discrepancy were comparable between the groups (\( p > 0.05 \)). There was a significant difference in the range of motion for internal rotation: 4.8° in the main vs 6.9° in the control group (\( p = 0.006 \)). The length of hospital stay was comparable, but the average operative times differed significantly, being shorter in the main group.

The follow-up examination conducted 4.7 years after the osteotomy revealed an increase in internal rotation, which was
significant in the main group (20.1°), compared to the control group (9.1°; \( p = 0.0024 \)).

Hip function was assessed 4.7 after the osteotomy using the Harris hip score; the average score was 94 points for the main group and 81 points for the control group (\( p = 0.001 \)).

The analysis of the obtained data showed that the main group subjected to a triplane osteotomy and fixation with the Trotsenko-Nuzhdin blade plate tolerated the procedure better (the intervention was shorter due to its technical simplicity). The duration of postoperative immobilization with antitrotation support was minimal in the main group, which facilitated the early commencement of rehabilitation. Consequently, most patients in the main group were able to improve internal hip rotation and restore the length of the affected leg to the maximum possible extent. Owing to the design of the Trotsenko-Nuzhdin plate (the bifurcated blade is introduced into the greater trochanter), the femoral neck was not traumatized intraoperatively, which had a positive effect on blood supply to the epiphysis.

Corrective femoral osteotomy involving correction of the proximal femur rotation axis allows recovering the proper centration of the femoral epiphysis, prevents angular deformation of the femoral diaphysis, hip subluxation and varus deformity of the metaphysis, reduces the risk of avascular necrosis of the epiphysis and articular cartilage chondrolysis, delays progression of hip arthropathy, and puts off the need for hip replacement until much later.

**CONCLUSIONS**

The proposed technique for corrective osteotomy of the femur in patients with stage 3 chronic SCFE prevents subluxation of the affected hip, deformity of the proximal femur and shortens rehabilitation time. The simplicity of the technique and stability of fixation result in shorter operative time, less intraoperative blood loss and make postoperative patient management less complicated.

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ADHERENCE TO TREATMENT IN VISUALLY IMPAIRED INDIVIDUALS

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Adherence to treatment is one of the major challenges posed by modern medicine. Today, cataract is the leading cause of reversible blindness and visual disability. The study was aimed to assess adherence to timely surgical treatment in individuals with cataract. The data of the cross-sectional, population-based Ural Eye and Medical Study were assessed. Among 546 participants, there were 46.3% men and 53.7% women, 59.6% urban residents, 40.4% rural residents. Their average age was 66.36 ± 9.47 years (40–88 years). Statistical data analysis was performed using the IBM SPSS Statistic software package. The findings showed that the lack of awareness of the disorder and low interest in surgery were the main factors, affecting the patients’ motivation for cataract treatment. At the same time, the frequency of ophthalmology visits was inversely related to the patients’ age (OR 1.24; 95% CI 1.04–1.49) and the duration of vision loss (OR 1.08; 95% CI 0.81–1.43), and directly related to the cataract diagnosis age (OR 1.20; 95% CI 1.04–1.38), the presence of ophthalmologist in the community clinic (OR 1.71; 95% CI 1.29–2.26), trust in the doctor (OR 3.62; 95% CI 3.02–4.35), ophthalmologist’s explanation of the cataract complications and advanced treatment methods (OR 1.62; 95% CI 1.34–1.97). Understanding the main factors, contributing to low treatment adherence in patients with cataracts, would make it possible to optimize the measures to improve healthcare delivery to such patients, associated with the increased coverage of surgical treatment.

Keywords: adherence to surgical treatment, cataract, awareness, quality of life

Author contribution: Bikbov MM — study concept and design, advising, editing; Israfilova GZ, Gilmanshin TR — data acquisition and processing, manuscript writing and editing.

Compliance with ethical standards: the study was approved by the Ethics Committee of the Ufa Eye Research Institute and conducted in accordance with the fundamental ethical principles of the Declaration of Helsinki, GCP (Good Clinical Practice) principles, and current regulatory requirements; the informed consent was submitted by all study participants.

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The term “adherence to treatment” emerged in the domestic scientific literature about 20 years ago, but it has become widely used recently, in the last 5–6 years. According to the World Health Organization (WHO), adherence to treatment is the degree to which a patient’s behavior matches the doctor’s prescriptions for taking medications, following dietary recommendations and/or lifestyle changes. It is believed that the patients’ non-compliance with medical recommendations results from the lack of effective doctor–patient communication. The patients’ personal and psychological characteristics, clinical features of the disease, treatment type, socio-economic factors, and the features of healthcare delivery in different countries are the other reasons for low treatment adherence [1]. This issue is patricularly relevant for eye diseases, since eye lesions are not potentially fatal, especially when the lesions could be removed. Many patients pay little attention to cataracts, since they usually have no prominent subjective symptoms for a long time. However, even the treatable eye lesions constitute a major medical and social challenge. The patient’s psychology in terms of the cataract treatment and prevention is one of the

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leading causes of blindness and low vision in the world [2]. High medical and social significance of cataracts is determined by a number of factors: high levels of blindness (reversible) and vision disability, significant direct (expenses for surgery, conservative treatment, and management of postoperative complications) and consequential (loss of labor potential of the society, traffic accidents, injuries, and falls due to visual impairment) economic losses, high prevalence of the disorder, clear upward trend in the number of individuals with reduced lens transparency of both retirement and working age [3–7].

It should be noted that the treatment success depends not only on the properly constructed healthcare system, but also on the patient’s compliance with the recommendations on the prevention and treatment of the disease, given by healthcare professionals [8, 9]. Poor adherence is a proven risk factor of any disorder, which reduces the effectiveness and increases the cost of treatment, increases the risk of various complications, worsens the outcome, and reduces the patients’ quality of life [10, 11]. Experts from the WHO have identified a large number of factors that affect the patient’s adherence to treatment. According to the WHO classification (2003), the factors are divided into five interconnected groups: 1) patient-related factors; 2) factors related to medical personnel and healthcare system organization; 3) factors associated with ongoing therapy; 4) factors associated with the patient’s condition (disease); 5) socio-economic factors [12]. In this regard, the papers, which propose structured approach to describing the causes of low treatment adherence, are of great interest.

The causes are divided into five major categories: socio-demographic; psychological; resulting from the disorder or the disease treatment; economic; related to the healthcare system [1]. Filling the gap in studying various factors, negatively affecting the capability of overcoming the treatment adherence barriers in patients with cataracts, would result in the significantly decreased incidence of this socially significant disorder. This is especially important in terms of improving the cooperation between the doctor and the patient, which contributes to faster recovery, improved quality of life and healthy lifestyle creation, and makes it possible to develop the measures to improve the quality of care provided to patients with cataracts.

For that reason, identification and analysis of the existing factors, including the psychological factors, which affect treatment adherence in patients with cataract, seem to be relevant.

The study was aimed to assess the attitude towards factors, affecting adherence to timely surgical treatment, in individuals with cataracts.

METHODS

Data of the cross-sectional, population-based Ural Eye and Medical Study (UEMS), conducted in 2015–2017 in the Ufa Eye Research Institute, were assessed. Currently, UEMS is the largest ophthalmological population-based study in the Russian Federation, which is aimed at studying the prevalence of socially significant eye diseases and associated risk factors.

The stages of the study were in line with the generally accepted standards. The study included the following: protocol development and the research instrument selection, data acquisition, scaling and database creation, statistical processing, analysis, and interpretation of the results [13, 14].

Inclusion criteria: voluntary consent to participate in the project; age over 40; permanent residence in the studied urban and rural areas.

To assess alterations in lens transparency, we used the LOCS III grading scale (Lens Opacities Classification System, 1993), according to which three types of lens opacification were distinguished: nuclear, cortical, and subcapsular [15]. According to this classification, nuclear lens opacities were divided into six grades. It should be noted that the most important are the changes in the lens nucleus, corresponding to grade 3 and above. For that reason, in the paper we link the nuclear cataract to these changes.

To study the respondents’ awareness of the lens disorder and their adherence to timely treatment, we conducted a questionnaire survey among people diagnosed with cataracts. The study was carried out by the written survey using the specially compiled questionnaire, which contained 25 questions concerning demographic information (gender, age), socio-economic data (level of education), the frequency and the possibility of ophthalmology visits, individual’s awareness of the disease, risk factors, and treatment methods, as well as the questions to assess the reasons, preventing the timely cataract surgery.

The study involved 546 individuals: 253 men (46.3%) and 293 women (53.7%), 325 urban residents (59.6%), 221 rural residents (40.4%). The average age was 66.36 ± 9.47 years (40–88 years). The number of observations required was justified using the method, developed by Otdelnova KA [16]. Statistical analysis was carried out using the IBM SPSS Statistic software package, version 23.0 (SPSS: An IBM Company; USA). The methods of descriptive and comparative statistics for quantitative characteristics were selected based on the distribution type assessment using the Shapiro–Wilk test. The groups were compared based on the qualitative characteristics using the Pearson’s chi-squared ($\chi^2$) test or Fisher’s exact test (in case of a small sample size or when expected frequencies were less than five in at least one cell of the contingency table). To characterize the univariate regression models and assess the degree of influence of each predictor on the disease development (outcome), the following parameters were calculated: regression coefficient ($\beta$), standard error of the coefficient (SE), Wald chi-square statistics (W). Odds ratios (OR), 95% confidence intervals (CI) for OR, and predictor significance were calculated based on these data. When performing statistical analysis, the achieved significance level ($p$) was calculated; the differences were considered significant at $p < 0.05$.

RESULTS

The analysis revealed that the majority of the respondents (60.85%) noted they had a secondary specialized education. A total of 21.70% respondents had higher education, 13.62% had secondary general education, and only 3.83% had incomplete school education. A survey of people over 40 years of age on their health-related behaviour showed that 39.2% of respondents got their ophthalmic checkups annually, 33.7% visited a doctor twice a year or more frequently, and 27.1% visited a specialist less than once a year. It should be noted that women were much more likely to seek ophthalmological care within a year (78.9 vs. 66.4%, $\chi^2 = 8.34; p = 0.01$). Furthermore, almost all patients (68.9% of men and 71.9% of women, $\chi^2 = 6.51; p = 0.04$) noted that they needed constant medical supervision. Among rural residents, characterized by the lack of healthcare access, the proportion of those who contacted a healthcare institution throughout the year was slightly less than among urban residents (71.6 and 73.6%, respectively, $\chi^2 = 1.13; p = 0.2$). Thus, rural residents remain active in applying to healthcare institutions.
It was found that the lack of knowledge about the need to see a doctor (19.6%) because of reduced vision was the main reason, preventing the respondents with cataracts from visiting an ophthalmologist on the annual basis. According to the respondents, the equally important reasons were as follows: being busy (18.2%), and no specialist physician in the community clinic (13.4%) (Table 1).

Patients with cataracts are characterized by low anxiety, and often by the condition severity underestimation. They do not consider their disorder to be the condition, significantly limiting their daily living activities, and assume that surgical treatment is fully capable to restore visual functions.

To investigate the respondents’ knowledge about cataract, the questions were included in the questionnaire, allowing us to assess the respondent’s awareness of the features of his/her eye disorder. The survey revealed a good knowledge on the disorder in the majority of respondents (60.2%); 17.8% of individuals had some ideas about the cataract, 5.9% were completely unaware, and 16.1% found it difficult to answer the question.

At the same time it should be noted that a large proportion of patients took an active stand on the matter. Thus, 60.4% of patients would like to know as much as possible about their disorder; 25.1% of individuals thought it was enough that the doctor was aware of the disease. The majority of patients (65.6%) would like to receive additional information about the disease from an ophthalmologist, as from a competent specialist. Furthermore, 19.3% of respondents would obtain such information from the online resources, 11.1% would get information in healthcare institutions (stands, brochures), 2.6% would read popular science magazines, and 1.4% would discuss the matter with their friends.

The doctor’s high professional level and positive personal qualities are essential for the patients’ compliance with recommendations, and the lack of this component increases the likelihood of self-treatment [10, 20]. The survey showed that the majority of the respondents (63.6 %) fully trusted their attending physicians and highly appreciated the physicians’ professional level. One third of the respondents (27.9%) noted the doctor’s high professional qualification and good personal qualities. The minority of respondents (1.4%) seemed not to fully trust their doctors and did not consider them to be the highly qualified professionals. Only 7.1% of respondents found it difficult to express their attitude. The following factors were among the reasons for “dissatisfaction” with the doctor: insufficient attention paid by the doctor (52.7%), the doctor’s inability to win over the patient (14.9%), and the doctor’s low level of competence (12.4%). However, the findings might indicate the high level of confidence in the doctors’ professional qualification and personality.

When a patient comes to see an ophthalmologist, it is extremely important for a doctor to explain the clinical features of cataract, such as functional vision impairment, which reduces the quality of life, and to provide information about the modern personalized treatment methods in case of reversible vision loss. Thus, 70.3% of individuals in the studied group indicated

![Fig. 1. Age distribution of respondents taking into account the source of information about cataracts](image-url)
Table 2. Factors associated with the frequency of ophthalmology visits in patients with cataracts

<table>
<thead>
<tr>
<th>Indicator (factor)</th>
<th>β</th>
<th>W</th>
<th>p</th>
<th>OR</th>
<th>95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years)</td>
<td>-1.54</td>
<td>22.54</td>
<td>&lt; 0.001</td>
<td>1.24</td>
<td>1.04–1.49</td>
</tr>
<tr>
<td>Duration of vision loss (years)</td>
<td>-0.39</td>
<td>8.26</td>
<td>0.004</td>
<td>1.08</td>
<td>0.81–1.43</td>
</tr>
<tr>
<td>Time since the diagnosis of cataract (years)</td>
<td>0.59</td>
<td>6.39</td>
<td>0.01</td>
<td>1.2</td>
<td>1.04–1.38</td>
</tr>
<tr>
<td>Presence of ophtalmologist in the community clinic (1 — present, 2 — absent)</td>
<td>0.55</td>
<td>13.54</td>
<td>&lt; 0.001</td>
<td>1.71</td>
<td>1.29–2.26</td>
</tr>
<tr>
<td>Confidence in the doctor (1 — yes, 2 — no)</td>
<td>0.33</td>
<td>19.248</td>
<td>&lt; 0.001</td>
<td>3.62</td>
<td>3.02–4.35</td>
</tr>
<tr>
<td>Have the ophtalmologist explained about cataracts and treatment methods (1 — yes, 2 — no)</td>
<td>0.48</td>
<td>23.63</td>
<td>&lt; 0.001</td>
<td>1.62</td>
<td>1.34–1.97</td>
</tr>
</tbody>
</table>

Note: β — regression coefficient; SE — standard error of the coefficient, W — Wald chi-square statistics, OR — odds ratio, 95% CI — two-sided 95% CI for OR.

that they learned about their eye disease when talking with the doctor; 15.2% of individuals got information from their friends and relatives, and 14.5% of respondents obtained information from mass media, such as TV, Internet, and press. Younger patients more often received information from the online resources, friends or relatives, while the respondents of the older age groups received information from ophthalmologists (Fig. 1).

We have defined the frequency of ophthalmology visits depending on the duration of visual impairment, the patients’ age, and the presence of ophtalmologist in the community clinic (Table 2).

The analysis revealed the impact of the patients’ age on the frequency of ophthalmology visits. Thus, every one-year increase in age reduces the probability of visiting a specialist physician by 1.24 times. Longer time since the diagnosis of cataract in the surveyed patients results in the 1.2 higher frequency of visits to ophthalmologists. Among individuals, who have ophtalmologists in their community clinics, the probability of annual ophthalmology visits is 1.71 times higher compared to individuals, having no ophtalmologists in their community clinics.

Studying the trust towards doctors as a factor affecting the patients’ adherence to regular medical visits, and their willingness to comply with the specialists’ recommendations, revealed the direct relationship. The calculated odds ratio (OR — 3.6) indicates that the chances to undergo ophtalmic examination are higher in individuals, who trust their doctors. A significant relationship has been also found between the frequency of ophthalmology visits and explaining the existing eye disorder. Thus, individuals, to whom the specialists explain the course and complications of cataract, as well as the advanced treatment methods, have a 1.6 times higher probability of the ophthalmology visits.

We have ranked the factors that, according to the respondents, prevent the timely cataract surgery (Fig. 2). It was found that 36.1% of respondents were unaware of the need for cataract surgery. In every fourth respondent, his/her professional or household activity was one of the key factors, constraining the decision on the timely cataract surgery. A total of 16.8% individuals reported they were afraid of the forthcoming treatment; 10.3% of individuals postponed surgical treatment due to the concomitant somatic disease.

These findings indicate the lack of the patients’ awareness of the disease and modern treatment methods. In this regard, informing the population, especially people over the age of 40, about cataract as a serious disease, which contributes to reversible blindness, is an urgent task.

DISCUSSION

The study has shown that insufficient awareness of the disease and the lack of interest in surgical treatment are the major factors, which reduce the patient’s motivation for cataract treatment. At the same time, ophthalmology visit frequency is inversely related to the patients’ age and the duration of vision loss, and is directly related to the presence of ophtalmologist in the community clinic, trust in the doctor, and ophtalmologist’s explanation of the cataract complications and advanced treatment methods. It should be noted that the lack of knowledge about the need for cataract surgery is a significant problem, especially among older age groups.

Fig. 2. Distribution of reasons for refusal of the cataract surgery
for cataract surgery has been the main obstacle to timely cataract surgery according to 36.1% of respondents, being busy due to employment or household activities was identified by 25.3%, being afraid of the forthcoming treatment was reported by 16.8%, and concomitant somatic disorders were mentioned by 10.3%. These findings are in line with the results of studying the patients’ treatment adherence in certain areas of medicine. Thus, a survey of patients with cardiovascular diseases revealed that the key factors contributing to low treatment motivation were as follows: misunderstanding of the doctor’s instructions (33.7%), fear of side effects or building up a tolerance (40.2%), comorbidity (35.9%) [17], no symptoms of the disease [18], patient’s refusal to depend on drugs or medical personnel, denial of the disease [19].

When studying dental patients, the key factors, negatively affecting the treatment motivation, were as follows: patients’ age (the vast majority of individuals were 35–49 years old); gender (women sought medical attention more often than men) [20]; quality of dental care [21]; underestimation of the condition severity by the patient; treatment costs [22]; painful procedure-related fear. In most cases the need for visiting a dentist is associated with acute pain (58.9%), 31.1% of patients schedule their dental visits, and only 22.8% of respondents would have routine check ups [23].

Thus, the study has shown that the factors associated with the patient’s personality (lack of awareness, being busy, fear, etc.), including psychosocial factors (beliefs, perceptions, lack of motivation), as well as the lack of trust in the doctor–patient communication, are the most significant barriers for the cataract treatment adherence. The patient’s limited knowledge about cataract and its sequelae together with the lack of motivation and positive preoperative expectations, as well as limited awareness of the advanced treatment methods, result in the delayed application for effective surgical intervention, and reduced quality of life.

CONCLUSIONS

To date, strengthening the prevention and treatment of sociomedical conditions, being the main cause of blindness and low vision, is a major challenge posed by the healthcare system. At the current stage, increasing the patients’ adherence to treatment is one of the main factors of improving the health status and the quality of life of the population. When analyzing attitudes towards factors that affected adherence to timely treatment in individuals with cataracts, it was found that the patient-related factors were the most significant (lack of awareness of the disorder, low patient’s interest). The findings make it possible to optimize the measures to improve healthcare delivery to patients with cataracts in order to increase the coverage of surgical treatment in such patients.

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