Russian guidelines for the management of COPD: algorithm of pharmacologic treatment

Abstract: The high prevalence of COPD together with its high level of misdiagnosis and late diagnosis dictate the necessity for the development and implementation of clinical practice guidelines (CPGs) in order to improve the management of this disease. High-quality, evidence-based international CPGs need to be adapted to the particular situation of each country or region. A new version of the Russian Respiratory Society guidelines released at the end of 2016 was based on the proposal by Global Initiative for Obstructive Lung Disease but adapted to the characteristics of the Russian health system and included an algorithm of pharmacologic treatment of COPD. The proposed algorithm had to comply with the requirements of the Russian Ministry of Health to be included into the unified electronic rubricator, which required a balance between the level of information and the simplicity of the graphic design. This was achieved by: exclusion of the initial diagnostic process, grouping together the common pharmacologic and nonpharmacologic measures for all patients, and the decision not to use the letters A–D for simplicity and clarity. At all stages of the treatment algorithm, efficacy and safety have to be carefully assessed. Escalation and de-escalation is possible in the case of lack of or insufficient efficacy or safety issues. Bronchodilators should not be discontinued except in the case of significant side effects. At the same time, inhaled corticosteroid (ICS) withdrawal is not represented in the algorithm, because it was agreed that there is insufficient evidence to establish clear criteria for ICSs discontinuation. Finally, based on the Global Initiative for Obstructive Lung Disease statement, the proposed algorithm reflects and summarizes different approaches to the pharmacological treatment of COPD taking into account the reality of health care in the Russian Federation.

Keywords: COPD, clinical guidelines, Russia, treatment algorithm

Introduction
COPD is an important health care problem due to its high prevalence and impact in terms of morbidity and mortality. A recent study conducted in 12 regions of the Russian Federation observed a prevalence of COPD of 15.3% related to the high rates of smoking in the Russian population. In addition, other studies have demonstrated the magnitude of the misdiagnosis of COPD in Russia and the particular characteristics of the COPD patients in this country. In a study of more than 1,500 Russian COPD patients, 53% had a forced expiratory volume in one second (% predicted) below 50% and up to 51% were frequent exacerbators. In a large survey in middle and Eastern Europe, the Russian patients with COPD were among those with the highest prevalence of frequent exacerbators. The high prevalence of the disease, the high rates of misdiagnosis, and the level of severity of the patients attended in Russian centers indicate the need for the development and implementation of a national clinical practice guideline (CPG) to improve the management of this disease.
The Global Initiative for Chronic Obstructive Lung Disease (GOLD) has published statements about the diagnosis and treatment of COPD; similarly, international scientific societies such as the European Respiratory Society and the American Thoracic Society have produced high-quality, evidence-based CPG. However, these GOLD recommendations need to be adapted to the particular situation of each country or region, taking into account the characteristics of the patients, the health care system, and the availability of treatments.

**Russian guidelines for the management of COPD**

The first national Russian guidelines on the diagnosis and management of COPD were developed and published in 1999 and have been updated several times thereafter. The most important update was published in 2014, when the Ministry of Health (MOH) of the Russian Federation proposed a standardized model for the development of CPG for all specialties and diseases according to the “International Statistical Classification of Diseases and Related Health Problems.” It was intended that CPGs had to become the key documents to determine different aspects of disease management, including legal and disability issues. The definition of a CPG by the Russian MOH was an “… evidence-based document representing physician activity in terms of diagnosis, treatment, rehabilitation and prophylaxis of the disease which helps to choose the right clinical decision.”

Russian guidelines may serve as an example of revolutionary approach to guidelines, closing the gap between professional societies and regulatory authorities.

The new attitude of the Russian MOH to guidelines was based on the understanding that CPGs could not effectively function as an independent isolated document of any professional society with no connection to the reality and complexity of real clinical practice and health care (which is regulated by the number of laws, by-laws, and subordinate acts) and not complying with other regulating documents.

Initially, the MOH built the conception that a CPG should be a major document regulating not only clinical practice but also expert evaluation of medical service quality and occupational disability. The electronic rubricator proposed by the MOH makes it possible to standardize guidelines in several terms (size, parts and sections, algorithms, etc.) and integrate these with all regulatory documents and other guidelines by cross-links and cross-references.

According to MOH definition, the unified electronic rubricator is a universally hierarchically structured integrated database for all CPG which is systematically updated. The unified electronic rubricator initially was designed to include all available approved guidelines in a single format and structure. According to MOH policy, before final placement of newly developed CPG into the unified electronic rubricator they need two-stage approval: by national professional society and MOH itself.

All parts and chapters are typed in by completing special prespecified forms which allow to keep the size and structure of the final document. Only documents complying with the structure, style, and other features can be placed into the rubricator. The whole database is equipped by search engine and cross-references.

The electronic rubricator also meant that the CPG must have a unified structure with the list of necessarily required and recommended sections. Among mandatory parts which seem to be important for COPD implementation in real life but which are not present in other available COPD guidelines are “COPD patient follow-up” and “criteria for evaluation of medical care quality.” In the first section, follow-up is specified according to severity and the level of the medical institution or health care provider (eg, 1st – local or 2nd – intermunicipal center) responsible for the follow-up of COPD patients with the number of visits per year and the list of examinations necessarily provided at any level in these medical institutions. In the medical care quality control section, evaluation criteria are listed for diagnostic and management procedures according to the severity of the disease and phase (stable condition or exacerbation).

Taking into account all these requirements and in concordance with them, an expert committee of the Russian Respiratory Society released a new version of the national COPD guidelines at the end of 2016. This update was based on the proposal by GOLD but adapted to the characteristics of the Russian health system and included an algorithm of pharmacologic treatment following recently published algorithms for the treatment of COPD. We believe that this proposal will contribute to the debate of the optimal treatment algorithm that combines strong scientific evidence with the reality of clinical practice for the treatment of COPD.

**Algorithm of pharmacologic treatment**

The algorithm proposed had to comply with the requirements of the Russian MOH to be included in the electronic rubricator. Therefore, a balance was sought between the level of
information and the simplicity of the graphic design. First, the diagnostic process was excluded from the algorithm and it was assumed that patients already fulfilled the internationally accepted diagnostic criteria for COPD. Second, although the decision tree included the characteristics proposed by GOLD—i.e., evaluation of symptoms and risks—it was decided not to use the letters A–D for simplicity and clarity. Third, an initial group of common pharmacologic and nonpharmacologic measures was excluded from the algorithm and placed before the decision tree, because these measures apply to all patients, irrespective of the level of symptoms or risk (Figure 1).

The treatment algorithm initiates with the evaluation of symptoms with the modified Medical Research Council dyspnea scale or COPD Assessment Test questionnaire, as suggested by GOLD, and using the same thresholds. However, long-acting bronchodilators (instead of short-acting) are suggested as the preferred therapeutic option. This decision was based on the higher efficacy of long-acting drugs and the possibility of underrecognition or underreporting of symptoms in patients with COPD.

Although it is recognized that initial therapy with a long-acting antimuscarinic agent (LAMA) or long-acting β2 agonist (LABA) is indicated in less symptomatic patients, the majority of COPD patients in Russia initially visit a doctor with a high level of symptoms and exercise intolerance. In the case of high symptom level (modified Medical Research Council ≥2 or COPD Assessment Test >10 units), a combined LABA/LAMA is recommended after establishing the diagnosis of COPD. The LABA/LAMA combination is more effective than monotherapies in dyspnea relief, increasing exercise tolerance and improving patients’ quality of life.

LABA/LAMA is also the preferred option when patients on bronchodilator monotherapy have recurrent exacerbations due to the superiority of the LABA/LAMA versus long-acting bronchodilators in monotherapy and especially versus the LABA/inhaled corticosteroid (ICS) combination in the prevention of exacerbations.

Combinations including ICS should not be used as first-line therapy in COPD. ICS as a component of a double (LABA/ICS) or triple (LAMA/LABA/ICS) therapy can be

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**Figure 1** Pharmacologic treatment of COPD.
**Notes:** Proposed algorithm of the Russian National guidelines for the management of COPD. *Predominantly non-infectious exacerbations (ACO or eosinophilic inflammation type).

**Abbreviations:** LTOT, long-term oxygen therapy; NIV, noninvasive ventilation; mMRC, modified Medical Research Council; CAT, COPD Assessment Test; LABA, long-acting beta-adrenergic; LAMA, long-acting antimuscarinic; ICS, inhaled corticosteroid; ACO, asthma–COPD overlap.
administered in situations when recurrent exacerbations occur during first-line therapy with long-acting bronchodilators and especially in patients who have concomitant asthma, based on internationally accepted criteria, or eosinophilic inflammation. However, there is no consensus yet on the threshold level of blood eosinophils, which predicts response to ICS in COPD. In patients presenting recurrent exacerbations in spite of receiving LABA/LAMA or triple therapy, it is necessary to investigate the phenotype and consider the possibility of administering phenotype-specific therapy. This therapy may include roflumilast in the case of severe airflow limitation, chronic bronchitis, and previous hospitalization; mucolytic agents particularly in chronic bronchitic patients with frequent exacerbations; and/or macrolides in the case of frequent infective exacerbations, especially in the presence of associated bronchiectasis.

At all stages of treatment, the efficacy and safety have to be carefully assessed, and escalation and de-escalation is possible in the case of lack of or insufficient efficacy of safety issues. Since COPD is a progressive disease and bronchodilators are only effective during administration, they should not be discontinued except in the case of significant side effects. In contrast, there is evidence of excessive and inadequate use of ICS in COPD, which may be associated with short- and long-term side effects. In fact, in a recent study conducted in Russia up to 19% of GOLD B patients were on LABA/ICS and 12% on triple therapy. However, ICS withdrawal is not represented in the algorithm, because it was agreed by the panel that there is insufficient evidence to establish clear criteria as to when and how to discontinue ICS in patients with COPD.

Conclusion
Although based on the GOLD statement, the algorithm proposed reflects and summarizes different approaches to the pharmacological treatment of COPD. It is well recognized that long-acting bronchodilators are the bases of treatment, but the most severe patients require a more personalized approach taking into account their particular phenotype in order to select the most appropriate treatment that achieves the best balance between high efficacy and low risk of side effects. In addition, the treatment algorithm must be simple to permit implementation in the electronic clinical records, and it should be easy to remember and use by clinicians in order to reduce the gaps in the application of COPD-treatment recommendations.

Disclosure
The authors report no conflicts of interest in this work.

References


