ENGLISH VERSION: PERSONALIZED DESENSITIZATION WITH ACETYLSALICYLIC ACID IN PATIENTS WITH HYPERSENSITIVITY TO NON-STEROIDAL ANTI-INFLAMMATORY DRUGS

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Aims: Hypersensitivity to nonsteroidal anti-inflammatory drugs (NSAIDs) has various mechanisms and represents different clinical syndromes from anaphylaxis to severe bronchospasm. The prevalence of aspirin hypersensitivity among patients with asthma and nasal polyps reaches 25.6%. Respiratory reactions associated with aspirin or other NSAIDs are not immunological. The basis of these reactions is non-allergic hypersensitivity of the cross-reactive type. Desensitization followed by long-term aspirin therapy is an effective method of treating hypersensitivity to aspirin or other NSAIDs. Using aspirin 600-1200 mg/day can significantly alleviate the symptoms of asthma, allergic rhinitis. Methods: We successfully applied aspirin desensitization for method of patients with hypersensitivity to NSAIDs. According to the method, an hour before the desensitization, daily montelukast 10 mg was taken orally, then aspirin every 3 hours. Results: Three patients underwent desensitization of aspirin. The dose was selected individually depending on the clinical manifestations of drug-induced adverse reactions (AR). ARs during desensitization were treated by iv dexamethasone administration. Subsequent doses did not cause AR. Doses of aspirin were increased to a maximum of 1250 mg daily, and were continued for the long-term use. Conclusion: It is possible to conclude that the initial dose of aspirin should be 16–40mg; it is possible to increase the dose if the initial dosage is well tolerated; symptoms of moderate intolerance are treated by 4-8 mg iv dexamethasone; prior to desensitization, we recommended to use montelukast 10 mg, it is safe to practice desensitization of aspirin according to a personalized technique by a specialist in an intensive care unit. Keywords: aspirin desensitization, asthma, personalized treatment.

Hypersensitivity reactions caused by ASA or other NSAIDs (synonyms, used previously, are “aspirin triad” and “asthmatic triad”) are manifested by bronchial obstruction, shortness of breath, nasal congestion or rhinitis, and are observed in patients with chronic respiratory diseases (asthma, rhinosinusitis and nasal polyps). The prevalence of this group of reactions ranges from 4.3 to 20%, depending on the population of the examined subjects and diagnostic methods. The presence of chronic rhinosinusitis with nasal polyps, severe asthma and female sex are associated with a high incidence of hypersensitivity to ASA and other NSAIDs.

Respiratory system responses, caused by ASA or other NSAIDs, are not immunological, i.e. they are not based on an antigen-antibody interaction. The non-allergic hypersensitivity of the cross-reactive type underlies these reactions. The main mechanism of these reactions consists in the inhibition of COX-1 by most NSAIDs – COX-1 inhibitors. Such a change in COX-1 activity is clearly manifested under conditions of impaired arachidonic acid metabolism. The disorders of arachidonate metabolism are as follows: 1) increased levels of leukotriene E4 in urine, expired breath condensate, bronchoalveolar lavage, induced phlegm and saliva due to increased 5-lipoxygenase activity; 2) decreased production of prostaglandin E2 by the epithelium of the upper and lower respiratory tract, accompanied by the inhibition of COX-2 and reduced basal production of lipoxin A4 by peripheral blood leukocytes; 3) increased expression of type 1 leukotriene receptors by the nasal mucosa cells [3].
Inhibition of COX-1 leads to activation of mast cells and eosinophils with a release of inflammatory mediators. Therefore, such reactions are characterized by manifestations of chronic eosinophilic inflammation. COX-2 inhibitors and low doses of COX-1 inhibitors are usually well tolerated by patients. Cysteinyl-leukotrienes (LT) are one of the main mediators, causing NSAID intolerance reactions. Consequently, blockade of type 1 leukotriene receptors by pharmacological inhibitors can alleviate clinical symptoms.

Viral infections may be an important factor in the persistence of respiratory tract inflammation in such patients.

Genetic polymorphism may play a certain role in the development of hypersensitivity reactions to NSAIDs. Associations of respiratory diseases, exacerbated by NSAIDs intake, with dipeptidyl peptidase-10 polymorphism, ALOX-15, IL-5 receptor, COX-1, etc. have been described [4, 5].

Clinical manifestations of hypersensitivity as a rule are observed in 30-180 min after administration of ASA / NSAIDs: patients develop non-bronchial symptoms (rhinorrhea, nasal congestion), ocular, skin (urticaria and / or angioedema) and gastrointestinal symptoms. Usually, a patient with hypersensitivity to NSAIDs presents with past history of chronic rhinosinusitis (sometimes complicated by mucosal hypertrophy and nasal polyps) and / or asthma. However, in some patients the intake of NSAIDs may cause the first bronchial asthma attack. Anosmia is a characteristic clinical feature in such patients.

Desensitization, followed by a long-term ASA therapy, is an effective treatment for patients with respiratory diseases, exacerbated by NSAIDs. The use of ASA at a dose of 600-1200 mg per day can significantly alleviate the symptoms of asthma and affected upper respiratory tract, as well as reduce the dose of intranasal corticosteroids (recommendation level B). In some patients, desensitization with ASA leads to a decreased frequency of polyps recurrence and repeated surgeries. Along with improving the management of bronchial asthma, desensitization with ASA also reduces the need for oral glucocorticoids [6].

We have tested and successfully applied the technique of desensitization with acetylsalicylic acid for patients with hypersensitivity to NSAIDs. One hour prior to desensitization, patients were prescribed montelukast 10 mg orally, 1 tablet, and ASA every 3 hours.

We present several typical cases that demonstrate an individual approach to desensitization.

**Clinical case No. 1**

Patient N., 25 years old, presented with complaints of low-productive cough with white sputum; cough persisted throughout the day; nasal congestion, hypernasal voice, lack of sense of smell. Case history: these symptoms had persisted since August 2017, starting with cough and nasal congestion. The patient had been treated by otolaryngologist for sinusitis. In November 2017, the patient had sustained an injury (back muscles strain) in a gym; she had been prescribed diclofenac sodium. After injection of the drug, the patient had developed bronchospasm for the first time. The attack had been arrested by a team of emergency medical service with dexamethasone 4 mg iv by stream infusion.

For the second time, bronchospasm attack had developed on taking 2 tablets of ibuprofen for menstrual pain. The patient had arrested the attack by oral administration of cetirizine hydrochloride 10 mg and dexamethasone 4 mg im. The patient referred to allergist who prescribed a checkup, during which an increase in IgE (172 IU / ml) and changes in the spirogram were detected. Before desensitization, the patient was examined:

- Blood test for total IgE (31.01.2018) – 172 IU / ml (norm is up to 100.0).
- 07.02.2018: Allergic tests. Pollen screening panel. Allergens from the pollen of birch, willow, black alder, hazel, poplar, dandelion, plantain, timothy grass, cockfoot, couch grass, common ragweed, common wormwood, meadow fescue, rye, grass, trees – the results were negative. Household panel: allergen from cat and dog dander, pillow feathers, the mixture of D. pteronyssius and farinae ticks – the results were negative. Conclusion: No sensitization to pollen, household, epidemic allergens was detected.
- 08.07.2018: X-ray of thoracic organs. Conclusion: No radiographically visible focal and infiltrative changes in the lungs were detected.
- 09.07.2018: Allergic tests were repeated. Pollen screening panel. Allergen from the pollen of birch, willow, black alder, hazel, poplar, dandelion, plantain, timothy grass, cockfoot, couch grass, common ragweed, common wormwood, meadow fescue, rye, grass, trees – the results were negative.
- 09.07.2018: Complete blood count from the vein: increased eosinophils – 9% (norm is up to 0.5-5.5%), no other deviations from the norm.
- 17.10.2018: Multispiral computed tomography of the paranasal sinuses. Conclusion: Spiral computed tomography pattern of hyperplastic pansinusitis.
- Nasocystogram (22.10.2018): increased neutrophil granulocytes – 87% (norm is up to 65-75%); eosinophilic granulocytes – 8% (norm is up to 0-5%), no other deviations from the norm.
- 24.10.18: Clinical diagnosis: Polyposic rhinosinusitis, intolerance to non-steroidal anti-inflammatory drugs (NSAIDs). Compensated BA.

The following treatment was prescribed: cetirizine hydrochloride 10 mg 1 tab once a day, budesonide / formoterol 160 / 4.5 μg 1 dose once a day, montelukast sodium 10 mg 1 tab once a day; consultation with otolaryngologist.

After examination, the patient was recommended to undergo desensitization to NSAIDs.

**Conduction of desensitization:**

The 1st dose of ASA was taken orally on 13.11.18 – 16 mg, the 2nd dose was 37.5 mg, 30 minutes after receiving the second dose, the patient developed nasal congestion, dry wheezing, shortness of breath, bronchospasm, which were arrested with dexamethasone 4mg iv by stream infusion; after arresting bronchospasm, the 3rd dose of ASA continued – 75 mg; in 30 minutes the patient again developed bronchospasm with rhinorrhea, dry wheezing, shortness of breath, which were arrested with dexamethasone 8 mg iv by stream infusion and inhalation of salbutamol 100 μg, 1 dose. On this day, ASA intake at a dose of 75 mg was discontinued.

The next day, on 14.11.18, one hour prior to administration of ASA, the patient took montelukast orally 10 mg, 1 tab, then the 1st dose of ASA (37.5 mg) was administered; the 2nd dose was 75 mg; the patient did not present any complaints, the 3rd dose was 150 mg; the
Clinical case No. 2

Patient M., 39 years old, presented with complaints of nasal congestion, rhinorrhea, hypernasal voice, lack of sense of smell, constant intake of vasconstrictor drugs. Case history: the patient had been ill since 2011, when she had noticed the annoying cough and nasal congestion. She referred to allergist; assays for worm infestation and allergy tests had been prescribed. The results had been normal. In 2011, during menstruation, the patient had taken ketorolac trometamol 30 mg for the first time. In 30 minutes, the patient had developed bronchospasm (wheezing, bursting chest pain, facial hyperremia). The patient had arrested the attack with dexamethasone and chloropyramine hydrochloride 25 mg.

The second episode of bronchospasm had occurred in the patient upon taking 1 tablet of ASA during SARS; the clinical presentation of bronchospasm was the same as when taking ketorolac. The attack had been arrested by the patient with dexamethasone 4 mg i/m and chloropyramine 25 mg 1 tab. The patient was examined: 09.11.2011: Analysis for total IgE – 86.7 IU / ml (norm is up to 87.0); Ab to Ascaris IgG – 0.17 (norm is up to 0.17); Ab to Trichinella IgG – 0.16 (norm is up to 0.16); IgE to allergens of dust Dermatophagoides pteronyssinus, Dermatophagoides farinae, German cockroach <0.1 kU / l were absent, IgE to allergens of microorganisms of Penicillium notatum, Cladospor herbarum, Asper fumigatus, Can albicans, Alternaria tenuis group <0.1 kU/l were absent.

15.12.2011: Spirogram: Moderate bronchial asthma by obstructive type. Test with bronchial spasmolytic (salbutamol 300 μg) – the result was positive. FEV1 Act1 – 2.19; % (A1 / P) – 67.3%; Act2 – 2.58; % (A2 / P) – 79.5%; % (A2 / A1) – 118.0%.

13.01.16: MRI (the paranasal sinuses). Conclusion: CT signs of polysinous rhinosinusitis.

Diagnosis: Polysinos of the nasal sinuses. The patient refused to undergo the recommended surgical treatment. Hormonal treatment with Avamis nasally and desensitization with acetylsalicylic acid were recommended.

Conduction of desensitization:

The 1st dose was taken orally on 23.10.18 – 18 mg; the patient did not present any complaints; the 2nd dose was 37 mg; the patient did not present any complaints; the 3rd dose – 75mg; 30 minutes after receiving the 3rd dose, the patient developed nasal congestion, rhinorrhea, sneezing, difficulty breathing; these symptoms were arrested with 4 mg of dexamethasone iv by stream infusion; the 4th dose was 150 mg; the patient did not present any complaints.

The next day, 24.10.18, the 1st dose of ASA was 225 mg, the 2nd dose was 300 mg, the 3rd dose was 375 mg, and the 4th dose was 450 mg. 25.10.18: on the last day of desensitization, ASA at a dose of 500 mg was administered: the 1st dose of ASA was 625 mg; the 2nd dose – 625 mg; the patient did not present any complaints. The patient was discharged from the department with a maximum dose of ASA (1250 mg).

Clinical case No. 3

Patient I., 28 years old, presented with complaints of nasal congestion, hypernasal voice, rhinorrhea, lack of sense of smell. Case history: these symptoms had appeared since 2006. The patient had applied self-treatment with vasoconstrictor drops. In 2012, the condition had been aggravated with cough, shortness of breath, and the following diagnosis had been made: Bronchial asthma. Polysinus rhinosinusitis. Deflection of the nasal septum. In 2014, the patient underwent surgery for polyps and deflected nasal septum.

In 2015, during the menstrual cycle, the patient had taken a pill of ketorolac. She had developed bronchospasm, which had been manifested by dry cough, sore throat, suffocation. The patient was examined: 02.03.2016: Allergic tests, ISAC package – the result was negative.

08.07.2014: Complete blood count: RBC – 4.58 * 10^{12} / l; Hb – 140 g / l; WBC – 5.97 * 10^{3} / l; ESR – 4 mm / h; PL – 322 * 10^{3} / l; leucogram: e. – 23.1%; bas. – 2.3%; s. – 37.9%; lymph. – 29.8%; mon. – 6.9%. Blood glucose – 4.96 mmol / l.

08.07.2014: Markers of viral hepatitis B and C were negative.

After examination, clinical diagnosis was made: Polysinous rhinosinusitis, intolerance to non-steroidal anti-inflammatory drugs (NSAIDs). Compensated BA.

After examination, the patient was recommended to undergo desensitization to NSAIDs.

Conduction of desensitization:

The 1st dose was taken orally on 19.06.18 – 40 mg; moderate rhinorrhea appeared; the 2nd dose – 40 mg; 30 minutes after the 2nd dose, the patient developed nasal congestion, rhinorrhea, sneezing, sore throat and dry rales, which were arrested with 4 mg of dexamethasone iv by stream infusion; after arresting bronchospasm, the 3rd dose continued – 100 mg; the patient did not present any complaints.

The next day, on 20.06.18, the 1st dose of ASA was 125 mg, the patient did not present any complaints; the 2nd dose was 250 mg, the 3rd dose – 375 mg; the patient did not present any complaints on that day.

21.06.18: One hour prior to administration of ASA, the patient took montelukast orally 10 mg (1 tab); on the last day of desensitization, ASA at a dose of 500 mg was administered: the 1st dose of ASA was 625 mg; the 2nd dose – 625 mg. The patient did not present any complaints. The patient was discharged from the department with a maximum dose of ASA (1250 mg).

After desensitization and further administration of ASA, the patient’s status was satisfactory. In August 2018, the patient developed rhinorrhea, nasal congestion. The patient was prescribed a check-up:

29.08.2018: Blood test for IgE – 258 IU / ml (norm is up to 20).

04.09.2018: Allergic tests. Pollen screening panel. There were increased concentrations of immunoglobulin E to goldenrod – 0.894 kU / l (class II 0.70-3.49); to dandelion – 2.54 kU / l (class II 0.70-3.49); to wormwood – 11.2 kU / l (class III 3.50 – 17.49).
Following the examination, it was found that the patient had sensitization with pollen antibodies. The patient was recommended to undergo specific immunotherapy with wormwood pollen.

**Discussion**

Three patients were desensitized with acetylsalicylic acid, and desensitization was personalized by nature. A dose of ASA was selected individually for each patient, depending on the anamnestic clinical manifestations of anaphylactic reactions. Two patients had an allergic reaction in the form of bronchospasm after ASA intake: one dose of 40 mg of ASA and the other dose of 37.5 mg of ASA. The 3rd patient developed rhinorrhea and sneezing on administering the 3rd dose of ASA (75 mg). In all three patients, allergic reactions were arrested by i/v administration of dexamethasone. Later on, subsequent doses did not cause allergic reactions. Doses of ASA were raised to a maximum of 1250 mg per day in 2 doses for a long-term administration.

One patient subsequently presented with clinically significant sensitization to wormwood pollen. In our opinion, this sensitization was masked by severe intolerance to ASA and revealed a seasonal allergy after desensitization.

Thus:

1) The initial dose of ASA should be 16-40 mg, depending on the presence of severe reactions in case history;
2) An increase in the dose of ASA is performed in case of satisfactory dose tolerance (mild rhinorrhea symptoms are allowed);
3) The appearance of moderate intolerance symptoms to ASA is effectively arrested by i/v administration of 4-8 mg of dexamethasone;
4) Most often, the symptoms of intolerance in desensitization with ASA are manifested when receiving doses of 37-75 mg;
5) It is advisable to administer 10 mg of monteleukast before desensitization with ASA;
6) Manifestation of desensitization with ASA is a safe technology when conducted using a personalized technique by trained personnel under conditions of intensive care wards;
7) Successful desensitization with ASA can contribute to the detection of seasonal allergies, previously hidden by year-round symptoms.

**References**