NSAID intolerance in chronic idiopathic urticaria: A study of its relationship with histamine-releasing activity of patients’ sera


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SUMMARY

Background: about one fourth of patients with chronic idiopathic urticaria (CIU) experience flares of hives after taking chemically unrelated nonsteroidal anti-inflammatory drugs (NSAID). The reasons for such intolerance are still elusive.

Objective: this study aimed to investigate NSAID intolerance in patients with CIU in view of the in vivo and in vitro histamine releasing activity of their sera.

Methods: 117 adults (M/F 41/76) with CIU underwent intradermal test with autologous serum, and the ability of their sera to induce histamine release from normal blood donors was evaluated. NSAID intolerance was ascertained by careful interview.

Results: overall, 32/117 (27 %) patients reported NSAID intolerance. The prevalence of NSAID intolerance did not differ in the three subgroups: negative on both in vivo and in vitro tests (9/36; 25 %), positive or intradermal test but negative on basophil histamine release assay (16/58; 28 %), or positive on both in vivo and in vitro tests (7/23; 30 %).

Conclusion: in patients with CIU intolerance to NSAID does not depend on the mechanism of histamine release.

Key words: Urticaria. Nonsteroidal anti-inflammatory drugs. Drug allergy. Cross-reactivity.
Patients

117 adults (M/F 41/76; mean age 37 years) with chronic urticaria, defined as recurrence of hives with or without angioedema for more than 6 weeks, were studied. A careful interview was carried out in order to detect patients who experienced exacerbations after taking NSAID.

Skin tests

Patients underwent intradermal test with 0.05 ml of sterile, fresh autologous serum and with saline as negative control. A skin prick test with histamine 10 mg/ml was used as positive control. Readings were taken at 15 and 40 min. Only an unequivocal wheal formation in response to serum injection was considered as a positive test. All intradermal tests were performed at least 5 days after the ingestion of the last antihistamine tablet (cetirizine 10 mg, fexofenadine 180 mg, loratadine 10 mg in all cases) and during a phase of moderate to strong clinical activity of the disease (9). After the skin tests, patients’ sera were stored at –20 °C for subsequent histamine release assays.

Basophil histamine release assay (HRA)

Leukocyte suspensions from normal donors were prepared by dextran sedimentation of peripheral venous blood, anticoagulated with 0.01 M EDTA, and mixed with 6% dextran in saline solution (Solplex 70, Sifra, Verona, Italy) and 30 mM dextrose (Sigma Chemical, St Louis, MO, USA). The cells were allowed to settle for 60-90 min at room temperature, according to Lichtenstein et al (13). The leukocyte-rich plasma was aspirated and centrifuged at 300 g for 15 min at 4 °C, and the cell button was washed twice in Hepes-buffered saline solution, pH 7.4, containing (mM): NaCl 140, dextrose 5.5, KCl 5, Hepes 5. Leukocytes (with about 7 x 10⁶ basophils) were then re-suspended in 100 μl volumes of Hepes-buffered saline solution with 1.8 mM CaCl₂ and 0.5 mM MgCl₂, and incubated with sera at the dilution of 1:2, making a final volume of 200 μl. After incubation for 40 min at 37 °C, the reaction was stopped by addition of 800 μl of ice-cold buffer solution and centrifugation at 1,000 g for 10 min at 4 °C. After centrifugation, the supernatants were aspirated, mixed with an equal volume of 6% HClO₄ and centrifuged at 1,000 g for 10 min at 4 °C. Histamine concentration in the supernatants was measured by an automated fluorometric method, according to Ruff et al (14). Spontaneous histamine release was evaluated by measuring histamine concentration in the supernatant of unstimulated cells, incubated for 40 min at 37 °C. Total histamine content was obtained by adding 100 μl of 6% HClO₄ to 100 μl of cell suspension. Net histamine release was calculated as percentage of total histamine content, after subtraction of spontaneous release. A 5% release cut-off value was used. Sera were tested with basophils from three normal donors, showing 30% net histamine release on challenge with an optimal dose of anti-IgE (10 μg/ml; Sigma Chemicals, St Louis, MO, USA).

RESULTS

32/117 (27%) patients reported NSAID intolerance (i.e. exacerbation of their disease after taking NSAID). On the basis of intradermal tests and HRA results, CIU patients were divided into 3 subgroups (table I):

1. Negative both on skin and serological tests (n = 36). 9 (25%) of these patients did not tolerate NSAID.
2. Positive on intradermal test but negative on HRA (n = 58). Such condition has been associated with a circulating autoantibodies to FcεRI or IgE (8-11). 16 (28%) of these subjects had a history of NSAID intolerance.
3. Positive on both intradermal test and HRA (n = 23). This condition has been associated with circulating autoantibodies to FcεRI or IgE (8-11). 7 (30%) of these subjects has a history of NSAID intolerance.

The prevalence of NSAID intolerance was almost identical in the three subgroups.

Table I

| Prevalence of NSAID intolerance in different subgroups of patients with CIU |
|-----------------|-----------------|-----------------|
|                  | ID test | BHR | NSAID intolerance (%) |
| I group (n = 36) | Negative | Negative | 25 |
| II group (n = 58) | Positive | Negative | 28 |
| III group (n = 23) | Positive | Positive | 30 |

BHR: basophil histamine release assay; ID test: intradermal test with autologous serum.
DISCUSSION

The overall prevalence of intolerance to NSAID among our patients was very similar to that observed by others authors (20-30 %) (1, 4-6), which shows that our population was representative of patients with CIU. The prevalence of NSAID intolerance was nearly identical in three subgroups of urticaria patients distinguished on the basis of different in vivo and in vitro histamine releasing properties of their sera; this shows that in patients with CIU, intolerance to NSAID does not depend on the mechanism of histamine release (whether induced by autoantibodies to FcεRI or IgE or by other hitherto not defined factors). Recently, an overexpression of LTC4 synthase, a key enzyme in cysteinyi leukotrienes production (16, 17), was observed in about 70 % patients with aspirin-induced asthma (AIA), and this has led to hypothesize that COX-inhibition may be responsible for NSAID-induced asthma attacks in AIA patients. Good results on both skin disorder and NSAID intolerance have been reported in some patients with CIU by the use of leukotriene receptor antagonists (18-20). Moreover, CIU patients show a chronic activation of the basophil/mast cell system (21, 22), and their sera may induce histamine release as well as de novo LTC4 production by basophils of normal subjects (23). Altogether, these observations suggest that LTC4 overexpression could underline NSAID-induced exacerbations of chronic urticaria as well. A study aiming to investigate this aspect is still in progress; unfortunately, preliminary (unpublished) results do not seem to support this hypothesis. At present, the reasons why one fourth of patients with chronic urticaria experience flares of hives after taking chemically unrelated NSAID remain undefined.

RESUMEN

Fundamento: cerca de un cuarto de pacientes con urticaria crónica idiopática (CIU) experimenta un enrojecimiento de las pápulas después de la suministración de fármacos antiinflamatorios no esteroideos (AINES). Alergia a medicamentos. Las razones de dicha intolerancia son todavía incomprensibles.

Objetivos: este estudio se propone investigar la intolerancia a los NSAID en pacientes con CIU examinando la capacidad de liberación de histamina in vivo e in vitro del suero de estos pacientes.

Métodos: a 117 adultos (M/F 41/76) con CIU se les practicó un test intradérmico con suero autólogo y se evaluó la capacidad del suero de inducir la liberación de histamina en la sangre de donadores sanos. La intolerancia a NSAID fue indagada a través de un cuidadoso interrogatorio.

Resultados: 32/117 (27 %) pacientes reportaron intolerancia a NSAID. No se encontró diferencia en cuanto a la prevalencia de la intolerancia a NSAID en los tres subgrupos: negativos en ambos tests in vivo e in vitro (9/36; 25 %), positivo en el test intradérmico pero negativo en el test de liberación de histamina (16/58; 28 %), o positivos en ambos tests in vivo e in vitro (7/23, 30 %).

Conclusiones: en pacientes con CIU la intolerancia a NSAID no depende del mecanismo de liberación de histamina.


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