In Vitro Diagnosis of Hypersensitivity to Nonsteroidal Anti-Inflammatory Drugs (NSAID) Comparison of Two Methods

Baló-Banga JM1* and Schweitzer K2

1Department of Dermatology, Medical Center of Hungarian Defense Forces, Budapest, Hungary
2Department of Pathophysiology, Medical Center of Hungarian Defense Forces, Budapest, Hungary

*Corresponding author: Baló-Banga JM, Department of Dermatology, Medical Center of Hungarian Defense Forces, H-1062 Budapest, Podmaniczky u. 109-111, Hungary, Tel: +36-1-475-2628; Fax: +36-1-302-0347; E-mail: balmat01@freemail.hu

Received date: April 12, 2017; Accepted date: May 05, 2017; Published date: May 13, 2017

Copyright: © 2017 Banga BJM, et al. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

Abstract

Objective: Current concept distinguishes between cross intolerance (non-immune) and single or multiple hypersensitivity based (immune) adverse reactions of non-steroidal anti-inflammatory drugs (NSAID) due to their potential to inhibit cyclooxygenase (COX) isoenzymes (COX-1, COX-2). Recently we described a rapid IL-6 release assay using blood mononuclear cells of patients with various clinical forms of drug hypersensitivity. Here we present data of a comprehensive analysis of the IL-6 release test and the classical IgE immuno-assay for their sensitivity in cases with adverse reactions to NSAIDs grouped according to the new clinical classification.

Methods: Total and specific serum IgE against 9 different HSA coupled-NSAIDs were determined by manual ELISA tests (55 cases) and compared to drug-specific release from preformed IL-6 pool of PBMCs of patients sensitized to the same NSAIDs after short (20') incubation of 4 standardized concentrations (51 cases and 9 controls) and IL-6 measurement from their cell free supernatants including positive and negative intraassay controls.

Results: The ratio of cross intolerant to specific hypersensitive (HS) cases was higher in the IgE group (and total IgE too,) than in the IL-6 release tested ones. There was no difference, however, in the overall ratio of early and accelerated plus late onset adverse events based on individual histories. Nine NSAIDs were tested in both groups which represented all major COX-1 inhibitors. The positivity of validated test results was double within the IL-6 tested group (65.4% vs. 36.9%). In some cases non-drug components of NSAID formulations were responsible for the observed (mainly) anaphylactic reactions. Positive results in both groups were scattered amongst cross intolerant and single to multiple hypersensitive (HS) subgroups. To our knowledge no comprehensive analysis had been performed before either on clinical phenotypes dependent IgE immunoassays or on NSAID-induced 'early' T-cell activation after those specified adverse events.

Conclusion: Specific HS and multiple non cross-reactive NSAID sensitizations exceeded non-immune reactions in both in vitro tested groups. Some intolerant patients revealed detectable ASA antibodies of IgE type. Preformed IL-6 release by PBMC was more sensitive than specific IgE immunoassays as an in vivo diagnostic tool. The results indicate that checking of non-drug components should be considered in allergy workups. ASA in vivo provocations need further standardization.

Keywords: NSAID cross intolerance; NSAID hypersensitivity; COX-1; COX-2; Drug-specific serum IgE; IL-6 release; Provocation tests; Tablet additives

Abbreviations: Acet: Paracetamol; AGEP: Acute Generalized Erythematous Pustulosis; ANO: Angio Neurotic Oedema; ASA: Acetyl Salicylic Acid (Aspirin); bid: Twice A Day; CIU: Chronic Idiopathic Erythematous Pustulosis; Con A: Concanavalin A; Df: Diclofenac; DRESS: Drug Rash with Eosinophilia and Systemic Symptoms; DSRIFE: Symmetrical Drug-Related Intertriginous and Flexural Exanthema; sNIUA: Single NSAID-Induced Urticaria/Angioedema; SNIUA: Single NSAID-Induced Urticaria/Angioedema or Anaphylaxis; SNIDr: Single NSAID-Induced Delayed Reactions; SNIDr: Single NSAID-Induced Delayed Reactions; TEN: Toxic Epidermal Necrolysis; UA: Urticaria Angioedema.

Introduction

Various NSAID inhibit cyclooxygenase isoenzymes (COX)-1 and -2 in plasma membranes to different extents. Because of their antipyretic and pain killing effect they are the mostly used drugs worldwide. Except for gastrointestinal and cardiovascular side effects they are responsible for the majority of drug hypersensitivity reactions [1] of both immunological and non-immunological types. Some years ago a unified classification by an expert panel has been published [2] in which there are two main groups: the “cross intolerance reactions”...
without allergic sensitization and the “classic drug allergic” ones. The former cause provocation by chemically non-related compounds, exclusively due to their relative affinity to COX-1 and COX-2 receptors in sensitive subjects. These can trigger either respiratory reactions like aspirin induced rhinosinusitis and asthma [3] or provoke cutaneous reactions like urticaria-angioedema (UA) progressing even to anaphylaxis. Chronic idiopathic urticaria (CIU) worsened by salicylates is a typical example and occurs in about 24% in hives patients [4]. Recently, basophil activation test and not serum IgE was proposed to indicate in vitro diagnosis in this subgroup of patients [5].

Salicylic acid are performed in frames of “early urticaria test series” including food additives with reading times different from those for contact allergy (i.e. 20’-70’-24 hrs). A large series of photo patch testing has revealed 9.2% positivity with NSAIDs [11].

Many blood tests have been described in the past 60 years to diagnose adverse drug reactions. ASA specific IgE could be detected in a (most likely) sNIUA case [12]. Recent publications: fail to confirm the applicability of drug specific IgE determination in the differential diagnosis of NSAID hypersensitivity and raise the possibility of metabolites as culprit substances [13]. In earlier studies our group has compared drug-specific IgE levels in serum, binding to drug-HSA discs using 125I coupled anti IgE and found high specificity but only 18.2% sensitivity against single dose non-blinded oral challenges in a retrospective study. Except for ASA, pyrazolones were also tested along with 5 different antibiotics [14] in hypersensitive and control cases but no detailed data for the various single drugs were evaluated. Cellular tests are difficult to perform, take days and are expensive. Pichler and Nyfeler found 78% sensitivity and 85% specificity of lymphocyte transformation test ( LTT) in 100 and 102 patients and stated that “pseudo allergies” to NSAID (i.e. Cross intolerance) were responsible for false positive results [15]. Basophil activation tests resulted in conflicting outcomes [16].

Table 1: Groups of NSAID hypersensitivity reactions.

<table>
<thead>
<tr>
<th>Classification</th>
<th>Clinical Expression</th>
</tr>
</thead>
<tbody>
<tr>
<td>Former NSAIADs**-Induced urticaria/angioedema</td>
<td>These reactions occur in about 24% of patients.</td>
</tr>
</tbody>
</table>
| Former NSAIADs**-Induced delayed reactions           | These reactions are delayed – i.e. sNIDRs. Tests are thus mandatory. Current concept prefers in vitro methods that expose the patients, like prick-tests for screening, intradermal tests [7], of both early and late-reading, and provocation tests [8]-the gold standard. There are serious limitations, however. One of them is history of anaphylaxis albeit no validated data comparing its severity grades with systemic adverse effects of skin testing have been reported. It was stated, that except for pyrazolones, none of other NSAID chemical classes would be recommended for routine skin testing due to lacking standardization [7]. We attempted to fill in this gap and proposed uniformly 10-3M test solutions for a wide variety of drugs, including the NSAIDs; pyrazolones (enolic acid derivatives) DF (acetic acid derivative) and ibu (propionic acid derivative) [9]. Oral provocation tests for aspirin are validated and are recommended to be performed even if another NSAID is suspected as a culprit in order to confirm or exclude COX-1 dependent cross-reactivity (NECD or NIUA-Table 1). These diagnoses arise if positive challenges even with minor symptoms occur after intake of both ASA and another chemically unrelated NSAID substance. The incremental doses of ASA starting from 10 mg up to 500 mg (cumulative dose) within one day are given according to established protocol [8]. No detailed descriptions for other NSAIDs oral tests could be found in the literature. Positive skin or general symptoms may arise within 4 hours after the last intake [10]. Patch test with ASA in particular and

Table 2: Phenotypic expression of drug hypersensitivity symptoms.

<table>
<thead>
<tr>
<th>Clinical Manifestation</th>
<th>Classic Drug Reaction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anaphylaxis</td>
<td>- Exacerbated respiratory disease</td>
</tr>
<tr>
<td>Urticaria/angioedema</td>
<td>- Exacerbated cutaneous disease</td>
</tr>
<tr>
<td>Single NSAID**-Induced urticaria/angioedema</td>
<td>- Induced urticaria/angioedema</td>
</tr>
<tr>
<td>Single NSAID**-Induced delayed reactions</td>
<td>- Anaphylaxis</td>
</tr>
<tr>
<td>Cross intolerance reactions</td>
<td>- Cross reactivity</td>
</tr>
<tr>
<td>*Cross reactivity with chemically related drugs.</td>
<td></td>
</tr>
</tbody>
</table>


In our previous recent study we found that preformed IL-6 released within 20 minutes at any or more of 4 different micro molar standard drug dilutions, tested on mononuclear cells above 50% over their diluents/ levels, significantly correlated with the patient’s history of drug–induced skin symptoms and with in vivo tests. Sensitivity of 85.4% and specificity of 82.4% of the IL-6 release assay was found [1]. Out of 98 patients 58 had a positive history of one or more NSAID. Thirty per cent of all tests in the patients and 27% in the control group had been performed with these drugs [1].

Our aim was to compare 2 independent in vitro diagnostic tests; drug specific IgE ELISA determination from sera and the 20 min. “early” T-cell activation detected by IL-6 release by standard μmolar concentrations of various NSAIDs based on self-reported history or the symptoms seen. Cross-intolerance phenotypes were scrutinized for...
results were accepted at >0.7 O.D. [Class 2] and >3.5 O.D. (Class used.

cases approved by the local Ethical Committee and patients gave their written consent to participate. Their symptoms were either seen by our staff or reported by the patients. The time elapsing between symptoms and tests was less than one year.

In vitro Methods

Two groups were formed with some overlapping cases. Group A consisted of 56 patients (36 women, mean age 46.6 yrs; 20 men, mean age 59 yrs.). Their sera taken after the adverse event was tested for specific IgE against various human serum albumin coupled NSAIDs: ASA, Ibuprofen, Acet, DF, Indomethacin, Pyrazolones (Met) and Oxicams by Hycore® (Great Britain) manufactured EIA (122 tests). The manual method was used. Threshold for positivity was 0.35 O.D. units. “True” positive results were accepted at >0.7 O.D. [Class 2] and >3.5 O.D. (Class 3=strongly positive). In 14 cases the total IgE was co-determined. In 13 cases in vitro tests have been carried out along with the in vivo tests.

Group B consisted of 51 patients (45 women, mean age 44.3 yrs; 6 men, mean age 50.3 yrs.). In addition 9 control subjects (5 men and 4 women) with proven tolerance were tested as well. Their mean age was 50.3 years.

Isolation of PBMC on Ficoll-Paque™ (Amersham Bioscience UK) gradients followed by two washes with PBS and resuspension in Dulbecco’s MEM containing 10 mM CaCl2 and MgCl2 each and 7 mM of glucose (energy source) was performed as described earlier [1]. The incubation of 1.1 x 10⁶/ml of PBMC (≥ 85% lymphocytes, viability >95%) without any plasma or serum was carried out in 450 µl aliquots with PHA-P (168 µg/ml) or ConA (5 µg/ml) as positive and PBS as negative controls and 4 standard (µmolar) dilutions (1.5; 2.5; 3.5; 5.0) for each NSAID added in 50 µl volumes. The final concentration for Asa, Ibuprofen, Acet, Met, Oxicams and additives were thus 10 times less (Figure 3). The incubation at 37°C for 20 min. was finished by cooling and centrifugation at 30-50 x g for 6 minutes. Altogether 91 series=546 single tests were performed.

In some cases tablet additives: sodium lauryl sulfate (SDS) and ferric oxides (yellow, red, brown; molecular mass: 159.7 M) were used. After incubation cell free supernatants obtained were frozen at-70°C. In the second step released IL-6 was measured by ELISA method using enzyme labeled monoclonal antibodies by Diagnosticum Ltd. Hungary [1]. Threshold for positivity was +50% IL-6 increases above negative (PBS) level at any of the 4 standard test concentrations provided that PHA-P or ConA samples were reactive as well. In 16 cases the total IgE was measured and in 18 cases in vivo tests have been performed within this group.

In vivo Methods

Patch tests have been carried out with ASA, Ibuprofen, Acet and Melox using 5-10% (w/w) of pure substances dissolved in petrolatum. Readings at 20-40, 70 and 24 hrs for ASA +Salicylic acid (Briel Co., Germany) and 20-40, 48-72 hrs for other drugs, tested. Positive readings included contact urticaria and/or dermatitis (1+4+ local strength). Intradermal tests have been performed using 10⁻³M solutions of pure substances in sterile saline compared to histamine (10⁻⁴M) positive and (saline) negative controls. Threshold positivity was noted if urtica >3 mm of diameter (d) and/or erythema >25 mm² developed within 20-40’ or papule >3 mm (d) has occurred at 24 hrs [9]. Oral challenges have been performed using single blind administration of a fraction of a tablet (1/4 or ½) given between 8-9 am. followed by close observation for 4 hrs and telephone contact for additional 24 hrs. Positive results were accepted if skin or respiratory symptoms and/or >20% deviation in vital parameters (blood pressure, pulse rate) have developed.

Statistical Analysis

Summary statistics and two tailed t-tests have been used

Results

1. Patients were (re) grouped according to the new classification as shown in Table 1. Both test cohorts revealed the majority of cases within the single NSAID induced sNIUA as well as in sNIUAA fractions and less in the delayed type reactions (sNIUAA). For this reason the two test series can be compared. In the present study Group A=67%; Group B=73% of immediate-early reactions could be noted. Regarding overall distribution of tested NSAIDs within the two groups a similar pattern can be recognized with some differences: There were 33% more tests with ASA in the IgE than in the IL-6 group, whereas 43% less tests of DF in the spec. IgE cohort as compared to the IL-6 group. The testing frequency was tailored according to each patient’s individual history (Figure 1).

Figure 1: Distribution of tests due to various drugs within the two groups “n” means the total tests in each group; in Group B only active cases (no controls) have been evaluated. Algopyrin=Metamisol (met).

The categories of “cross intolerance”, respiratory symptoms dominance (NERD) and multivalent NSAID induced NIUA appeared mostly in the specific IgE group (9 and 7 cases). In NERD only ASA was positive; in 4/9 Class 2-3 and in 1/9 borderline positive cases. Two cases in this subgroup had suffered from typical symptom triad of Samter [15]. In suspected NIUA (NECD) cases of the specific IgE group only one reacted positively (Class 2 units) to ASA and DF. The remaining 6 revealed 13 negative or weakly positive (Class 1) results to
various drugs; ASA, Df, Ibu, Met, Acet. The clinical phenotype in all cases was urticaria and ANO. Three positive provocation tests occurred. The sNIUA cases of Group A revealed 14/30 (47%) positive specific IgE results with highest prevalence of Melox (1/1), Ibu (2/3) ASA (5/9), followed by Met (4/8), Acet and Df (both 1/3); NERD was suspected in the IL-6 release group in one case. ASA was not tested but Df was weakly positive. In the NIUA suspect fraction of Group B, 3 cases were included. All had urticaria and ANO and multiple NSAID induced reactions with preferential localization in the periorbital area. All of 7 tests with various drugs were positive. Among them one patient (23 years man) has suffered from polyvalent allergies including various foods, pollens and house dust mite. In the sNIUA cases of Group B there were 24/32 (75%) positive IL-6 release results. The highest prevalence occurred with Nap (2/2), followed by Met (8/10), ASA (5/7), and Df (4/8), Melox (1/2), Ibu (1/2). It has to be mentioned that one of 2 tests with Ibu was false negative, proven by intradermal test positivity later. The least positive results (1/4) were encountered by testing Acet and Tramoxenic acid (0/1). The sNI'Dr fraction of Group A contained 18 patients and 9 positive at test rates 3/13 (23%). There were equally 4/13 positive tests for ASA and Met; 3/13 for Acet and 2/13 for Df. Ibu was negatively tested. Within the sNI'Dr fraction of Group B 13 patients and 19 tests were considered. Out of them 10 had positive results, one of whom had tolerated the drug tested. Thirteen tests were positive and 7/13 (54%) single drug positivities had occurred, while 3 patients showed reactions to 2 unrelated drugs. Three patients have revealed 6 (only) negative reactions. Df, Acet and Met were positively tested at equal higher rates while ASA and Ibu at equal low rates. The most frequent diagnoses within the sNI'Dr groups were maculo-papular rashes, fixed drug exanthems, vasculitis and purpurae, DRESS, late onset urticaria, prurigo and SDRIFE'.

2. Total IgE levels: In Group A significantly elevated total IgE (363 kU/l ± 83 s.e.m.) over Group B (89.7 kU/l ± 28.3 s.e.m.) was noted. Two-tailed probability gave p=0.0067 value. In spite of the significant difference, both groups revealed many positive specific IgE results against pollens, food antigens and to other drugs, mostly to antibiotics.

3. Comparison of humoral (spec IgE) and cellular (PBMC IL-6 release) tests (Table 3.): Criteria upon which the 2 tests could be compared regarding negativity and positivity taking into account the grading as well are demonstrated in Table 3. There were nearly the same rates of negative results in the two groups. The corresponding definitions are explained in column 1 for the specific IgE determinations and in column 4 for the IL-6 release assay.

Positive test rates are also similar except for strong positivity occurring in IL-6 tested cases with more than double frequency. There were 8 cases (all women) and 9 tests in this group.

<table>
<thead>
<tr>
<th>Definition 1</th>
<th>Result 1</th>
<th>Spec. IgE (%) n=119</th>
<th>Definition 2</th>
<th>Result 2</th>
<th>IL-6 release (%) n=87</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;0.35 O.D.</td>
<td>Negative</td>
<td>34.4</td>
<td>No signif. release</td>
<td>Negative</td>
<td>34.3</td>
</tr>
<tr>
<td>Class 1</td>
<td>weakly + uncertain</td>
<td>28.6</td>
<td>single peak, except at 0.15 µM</td>
<td>weakly +</td>
<td>16.1</td>
</tr>
<tr>
<td>Class 2</td>
<td>positive</td>
<td>29.4</td>
<td>&gt;1 peaks or one at 0.15 µM</td>
<td>positive</td>
<td>35.8</td>
</tr>
<tr>
<td>Class 3</td>
<td>strong pos.</td>
<td>5.0</td>
<td>&gt;1 peaks incl. one at 0.15 µM</td>
<td>strong pos.</td>
<td>12.7</td>
</tr>
<tr>
<td>&lt;0.35 O.D.</td>
<td>undefined, negative</td>
<td>2.5</td>
<td>Any release, not &gt; (+) 50% backgr.</td>
<td>Undefined negative</td>
<td>0</td>
</tr>
</tbody>
</table>

Definition 1 is the widely accepted class grading (25) except for the fluorescent enzyme immunoassay (FEIA-ref 29); Definition 2 was created from the previous work of our group (ref 1). Left side columns /Group A/, right side columns /Group B/. Column 6 reflects the results without the 9 control person's 16 negative tests (validated by in vivo tolerance) because in Group A there were no negative controls included.

Table 3: Evaluation of relative frequencies of negative and positive results obtained by the 2 tests.

Upon comparison with Group A in Group B 4 of eight cases with strong positivity belonged to late reactors (sNI'Dr), 3 cases to sNIUA and one to sNIUA-Anaphylaxis (Table 1). Seven of 9 tests with various NSAIDs and 2/9 with SDS gave strong positive results. The total IgE values measured in this group fell into negative range (20-30 kU/l). The four strong reactors (5 tests) of Group A were women, 3 of them belonged to sNIUA and one to sNI'Dr. All had very high specific IgE values.

4. The negative, weakly positive and undefined results amounted altogether to 65.5% of specific IgE tests (lines 2+3+6, column 3). Within the Group B even slightly positive results (IL-6 > +50% at any of specified drug dilutions-see definition) could be clinically validated as true ones. This has resulted in 65.5% overall positivity rate of this test (lines 3+4+5, column 6) against 34.5% of group A. While ASA was tested most frequently in Group A resulting in 55.2% positivity (Classes 2-3) all other drugs yielded only from 25% (DF, Oxicams) to 33.3% (Ibu) "true" positive results in all subgroups together. Acet and Met were at 30.4 and 30.3%, respectively.

In the IL-6 release assay the most frequently tested drug was Df with 41% positive results, followed by Met and ASA revealing 83.3% and 76.5% positivity. Nap (Propionic acid derivative) was positive in all tested cases. Melox tests were equally positive and negative (50%).

5. Non-NSAID ingredients of tablets as culprit substances could be detected by the IL-6 release assay. In some unexpectedly occurring negative results, other non-drug components of the formulation e.g. SDS in some ASA tablets or ferric oxides in DF tablets were tested positively. Figure 3 demonstrates the results obtained by IL-6 release from sensitized mononuclear cells due to the suspected culprit drug ASA and tablet ingredient SDS. The results of two independent experiments (and times) are shown. While rapid onset purpura on the legs persisted, the 64 y old woman revealed negative IL-6 test to aspirin which she took while the rash has appeared. When the drug was decided to test SDS (unknown** amount in the stomach protective ASA tablet). Strongly positive result has occurred. In vivo proof has been obtained later as the patch test was highly positive as well. The
patient continued to take another ASA formulation without SDS and has remained symptom free. Ferric oxides as tablet colorants have the same molecular weight but their different color depends on shape and size of particles. All four patients tested were women and the suspected drug was Df. Three patients had negative in vitro results and only one was positive with Df. Two of the 4 showed anaphylaxis and emergency unit care was necessary. The other 2 had suffered from widespread UA rash. “Yellow” Fe₂O₃ was positive in all of them as well as in one of 4 tests the “brown” iron oxide stain. Two patients were tested with the red colorant as well but were negative.

Figure 2: Percentual distribution of patients tested by different methods due to internationally agreed phenotypic classification.

Figure 3: The effect of allergy eliciting hapten on the release of IL-6 from sensitized PBMCs. SDS is an additive within some ASA tablets. Test with ASA: negative, patient has tolerated ASA tablets without SDS. Test with SDS positive!

Figure 4: Patch test performed to prove SDS hypersensitivity in a 64 year old women who took gastric mucosa protective ASA tablets (reading after 48 hrs). Her in vitro results are shown in Figure 3. Arrow indicates methyl-isothiazolidin (0.1%); PPD=paraphenylene diamine free base ; Colofon=colofony; Detergens sulfuratum FoNo (Formulæ Normales)=medicinal hair shampoo in Hungary, containing small amount of SDS.

6. In vivo - in vitro comparisons: In 16 patients of the IgE test group seven skin tests (intradermal, patch) and 9 provocations have been performed. Out of 11 positive reactions, one patch, 2 intradermal and 8 provocations were observed. The most positive results were obtained with ASA (7) followed by Df (2) and Ibu (2). In patients of the IL-6 group 21 in vivo parallel test were carried out. Out of 8 positive reactions one patch and 7 provocations have been performed. ASA, Df and Ibu were positive in 2 cases (each) Acet and Met in one case (each). In addition, 13 tests in 8 tolerant control subjects of this group were negative. In one control case however, 125 mg of oral Met intake was tolerated while 250 mg caused mild symptoms within the observation period. Figure 5 shows positive patch tests with two chemically different NSAIDs. Figure 6 demonstrates disseminated red spots on the neck due to positive patch testing with ASA (20-70) on the back and subsequent spreading.

Figure 5: Multiple positive patch tests (reading after 48 hrs) by Df (acetic acid derivative) and Ibu (arylpropionic acid derivative) after non-immediate but accelerated (sNIUA) symptoms in a 64 y old woman.
Discussion

In an earlier series of our group the ratio of immediate-early to delayed reactions was 6:4 [10]. Since that time a new clinical classification has emerged (Table 1). Our results pointed to the fact that hypersensitivity related phenotypes (sNIUA, sNIUAA, sNIDr—see Table 1) to various NSAIDs clearly exceeded those of presumed “cross intolerance” cases without true sensitization. The higher representation of NECD (9/56) and of NIUA (7/56) in Group A against those (1/51) and (3/51), respectively, in Group B reflected the first line selection of IgE-based diagnostics for rapid onset respiratory and urticarial reactions. This is also expressed by significant difference in total IgE values. The specific IgE tests showed within the NERD subgroup 5/9 negative results and 4/9 true (≥Class 2) positive readings with ASA. Out of them one of two with Samter triad [16] and IgE positivity has tolerated even 200 mg of aspirin given in gradually increasing amounts under ward conditions. The other patient with negative IgE result to ASA has developed respiratory symptoms to 100 mg ASA given orally without known previous sensitization but turned to tolerate this drug if 50mg bid was administered. Both decided to continue ASA treatment under ward conditions.

The key event is the “flare-ups” depending on the high level of T-cell reaction elicited by the first event but disappearance later with elapsing time. Moreover, recent work has stated multiple NSAID hypersensitivity without allergy to aspirin [23]. From our sNIUA cases of Group A 2/3 were “true” positive to ASA and from sNIUA of Group B 4/8 tested for ASA due to history-reacted all positively. IL-6 release assay results therefore, were of no help separating NIUA and sNIUA subgroups as distinct clinical entities. The higher incidence of negative tests by Df and Melox might reflect their lower potency of COX1/COX2 inhibition [3] well established by now for each NSAIDs in use [21]. The sNIDr phenotypes showed the most single drug induced positive tests in both Groups (A&B). In Group A 4/18 patients had multiple hypersensitivities with chemically non-related drugs. In Group B 2/13 patients reacted positively with chemically distinct drugs (2 each). The most frequent associations in both test cohorts were Met (Pyrosoleno-enic acid derivative) and Acet (p-aminophenol derivative). The mechanism of “early” release of preformed IL-6 from mononuclear cells has been studied immediately from its preformed pools and binds to both its soluble and cell membrane attached receptors. Furthermore, it also binds to the ubiquitous membrane receptor gp 130 and through this “trans-signaling” and forming of a “functional receptor complex” an immune response could start. Except for “turning on” antigen presenting monocytes, lymphocyte activation starts as well and T-cell proliferation would occur [24]. We found that multiple test concentrations were needed to detect total sensitivity related to clinical phenotypes [1].

More severe or widespread rashes tended to release significant (>50% over background) IL-6 at the lowest or at multiple drug concentrations. The grading of test positivity was based on these observations (Table 3) and supported also by our previous results on a large group of various drugs [1]. The serum specific IgE levels were divided into classes which indicate tenfold increases of analyte in each step (Classes 1-4). It is generally agreed that only class 2 or greater could be interpreted as clinically significant or positive. Below this, class 1 is dubious or negative and 0 is definitely negative [25]. This was the basis of our comparison as detailed in Table 3. The good correlation of the percentages of negative and of undefined negative
results within the Groups A and B could argue for the comparability of the two groups. The approximately two fold increase of positivity rates in favour of IL-6 release tests stresses its applicability in the differential diagnosis of NSAID-induced adverse reactions. There is no need for sterile cell cultures and much less time is necessary to obtain results. The lack of sensitivity of available NSAID–specific IgE assays does not exclude the role of IgE-mediated clinical presentations or skin test results [26]. Recently, Steiner et al., listed a number of publications considering data for NSAID specific IgE tested in “early onset” hypersensitivity reactions as “not available” [27]. Single drug tests to the pyrazolone derivate propyphenazon were of high diagnostic value though [28].

Our results could at least partly fill in this gap, regarding other NSAIDs. It is important to mention that modern automated analyzers would be much less sensitive in detecting pathologically elevated NSAID-specific IgE levels than the “older” manual ELISA test systems [29]. The finding of HS related to non-drug components of some NSAID formulations was established by using the more sensitive IL-6 release test and this could widen our horizon looking into adverse reactions occurring after taking “pain-killers” or anti-thrombotic drugs. The weakness of the above studies is the lack of ASA tests for all patients in all cross-intolerant subgroups to match with the provocation tests. One patient in the sNIUA subgroup A who reacted positively to oral Nimesulid (COX-2 antagonist) had positive skin tests to various sulphonamides, a basic structure to all COX-2 antagonists in use.

The aim of future clinical research is to differentiate between cross intolerance reactions of selective COX-1 inhibitors and multiple allergy syndrome involving both COX-1 and COX-2 inhibitors.

Conclusion

T-cell IL-6 release measured after short incubation from supernatants can be recommended to supplement to or substitute for in vivo testing after hypersensitivity events to NSAIDs. Specific IgE determinations are of limited value because of high false negativity. The intolerance reactions with no HS need a different allergy workup starting with in vivo tests with ASA. The majority of our skin patients did not fall into these categories. Multiple non cross-reactive HS was common among early accelerated phenotypes.

Funding Source

This work was supported by the Hungarian Research Grants OMFB-00284/04 and OMFB-00285/04.

Conflicts of Interest

Authors declare no conflict of interest.

Acknowledgements

Authors are indebted to L.A. Réthy, MD, K. Rásky and Prof. K. Balogh for critically reviewing the manuscript.

References


