Penicillin and cephalosporin cross-reactivity: role of side chain and synthetic cefadroxil epitopes

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Abstract

Background: Analysis of cross-reactivity is necessary for prescribing safe cephalosporins for penicillin allergic patients. Amoxicillin (AX) is the betalactam most often involved in immediate hypersensitivity reactions (IHRs), and cefadroxil (CX) the most likely betalactam to cross-react with AX, since they share the same R1 side chain, unlike cefuroxime (CO), with a structurally different R1. We aimed to analyse cross-reactivity with CX and CO in patients with confirmed IHRs to AX, including sIgE recognition to AX, CX, CO, and novel synthetic determinants of CX.

Methods: Fifty-four patients with confirmed IHRs to AX based on skin test (ST) and/or drug provocation test (DPT) were included. Serum sIgE to AX and benzylpenicillin was determined by Radioallergosorbent test (RAST). Two potential determinants of CX, involving intact or modified R1 structure, with open betalactam ring, were synthesised and sIgE evaluated by RAST inhibition assay.

Results: Tolerance to CX (Group A) was observed in 64.8% cases and cross-reactivity in 35.2% cases (Group B). Cross-reactivity with CO was only found in 1.8% cases from Group B. ST to CX showed a negative predictive value of 94.6%. RAST inhibition assays showed higher recognition to CX as well as to both synthetic determinants (66% of positive cases) in Group B.

Conclusions: Cross-reactivity with CX in AX allergic patients is 35%, being ST not enough for prediction. R1, although critical for recognition, is not the unique factor. The synthetic determinants of CX, 1-(HOPhG-Ser-Bu) and 2-(pyrazinone) are promising tools for determining in vitro cross-reactivity to CX in AX allergic patients.

Keywords: Amoxicillin, Betalactam, Cephalosporin, Cross-reactivity, Drug allergy, Antigenic determinant, Specific IgE

Background

Betalactams (BLs) are the drugs most frequently involved in immediate (IgE-mediated) hypersensitivity reactions (IHRs) [1–3], which could be explained by their ability to act as haptens due to their high chemical reactivity against proteins [4, 5]. BL chemical structure is formed by a 4-membered ring (the so-called BL ring) that in penicillins is fused to a 5-membered thiazolidine ring, and in cephalosporins to a 6-membered dihydrothiazine ring (Fig. 1). These drugs have a side chain (R1) bound to the BL ring; besides, cephalosporins have a second side chain (R2) bound to the dihydrothiazine ring, whose chemical structures distinguish the different compounds [6, 7].

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Penicillins are the most consumed antibiotics in Europe, representing 37% of total consumption, followed by cephalosporins with a 15% of total antibiotic consumption [8]. Among them, amoxicillin (AX) is the most consumed and the most often involved in IHRs to BLs followed by cephalosporins [3, 9] which include the following: cefuroxime (CO), ceftriaxone, cefazolin, cefaclor, and cefadroxil (CX) [10, 11], with different percentage of cross-reactivity between them [6], highly related to their chemical structure [12–14]. Cross-reactivity rate with cephalosporins in penicillin-allergic patients with IgE-mediated reactions ranges from 0% to almost 40% depending on the chemical structure of the BL involved [15–22], specifically on similarity in the R1 side chain [23, 24]. In this context, AX, which shares the same amino R1 side chain with CX (Fig. 1), could have a high cross-reactivity [19–21]. Conversely, CO, with a different R1 side chain, has shown tolerance in patients with IHRs to penicillins [16–19] and, more recently, similar results have been found with cefazolin and cefditoren [22, 25].

Cross-reactivity has important clinical implications, especially for searching safe alternative for further treatments, and an accurate diagnosis based on skin testing (ST) is recommended, being the role of drug provocation tests (DPT) controversial [3, 9, 26]. In vitro evaluation of cross-reactivity to BLs, mainly based on immunoassays, is limited by the difficulty for studying the structure of cephalosporin-protein conjugates [27]. Although several reports have addressed this issue [28–30], the antigenic determinants of cephalosporins are currently not well-known [31].

To our knowledge, structure–activity relationship (SAR) studies have been the unique successful approach for investigating cephalosporin epitopes [28–30, 32]. In this context, we have elucidated precise epitope structures through synthesis and immunologic evaluation of well-defined structures proposed as antigenic determinants for cephalosporins with different R1, bearing different functionalities at the C-6 of the cephalosporin (methyl, hydroxymethyl, aldehyde, mercaptomethyl) and without involvement of the remaining dihydrothiazine ring [29, 30]. Moreover, we have identified a novel synthetic pyrazinone structure as an antigenic determinant of cefaclor [28], formed after reaction of the amino group in the R1 with the likely aldehyde functionality at C-6 of the original cephalosporin [28, 32]. CX is another aminocephalosporin that could follow the same fragmentation and reactivity pathways as cefaclor [32].

In this study we have evaluated the in vivo degree of cross-reactivity with CX and CO in patients with confirmed IHR to AX and the immunological recognition of AX and these cephalosporins by serum specific IgE (sIgE). The ultimate aim of this study was to evaluate if synthetic structures, proposed as potential antigenic determinants mimicking the fragment of CX, which would remain coupled to the protein, can help get insight into the structure responsible for CX allergies and, therefore, study cross-reactivities between AX and CX.

### Methods

#### Patients

The studied group was obtained from the Regional University Hospital of Málaga Drug Allergy Database. This prospective cohort includes all patients with confirmed drug allergy from 1984 to 2019 after an allergological workup including clinical history, ST, and DPT.

Patients with IHR to AX (allergic to the whole penicillin group or AX selective reactors with good tolerance to penicillin V (PV)) were diagnosed following the European Academy of Allergy and Clinical Immunology (EAACI) recommendations [9, 33]. Tolerance to CX and CO was evaluated and, based on CX tolerance, patients were classified into: Group A with tolerance (demonstrated by negative ST and DPT) and Group B with cross-reactivity (demonstrated by positive ST or DPT) (Fig. 2).

#### Skin test

Skin prick (SPT) and, if negative, intradermal tests (IDT) were performed as described [9, 33], using benzylpenicilloyl-poly-L-lysine (PPL, DAP, Diater, Leganés, Spain) at 1.07·10^{-2} M, minor determinant mixture (MDM: benzylpenicillin, benzylpenicilloate,
and benzylpenilloate) at 1.5 M and AX (Diater laboratories, Madrid, Spain); CX (Lilly SA, Madrid) and CO (GlaxoSmithKline S.A, Madrid) all at 20 mg/mL. Since May 2011 DAP composition has changed and includes the major determinant benzylpenicilloylocta-L-lysine (BP-OL) at 0.04 mg/mL, equivalent to $8.64 \times 10^{-5}$ M concentration of the benzylpenicilloyl (BPO) moiety, and the minor determinant (MD) at 0.5 mg/mL, equivalent to $1.5 \times 10^{-3}$ M concentration of sodium benzylpenilliole. Cephalosporin reagents were prepared according to Romano [19, 34].

Readings were done after 20 min and considered positive: (i) In SPT, if a wheal larger than 3 mm surrounded by erythema appeared, with a negative response to the control saline; (ii) In IDT, if the increase in diameter of the wheal area marked initially was greater than 3 mm surrounded by erythema. Positive data expressed as the mean diameter recorded by measuring the largest and the smallest diameters at right angles to each other [35].

**Drug provocation test**

In subjects with negative ST to PPL/BP-OL and MDM/MD, oral DPT with PV was performed at incremental dose (50, 100, 150 mg) each 40-min until reaching the total cumulative dose (TCD) of 400 mg, followed by a 2 day therapeutic course of PV of 400 mg/8-h at home [33]. If DPT with PV and ST to AX was negative, oral DPT with AX was performed (50, 100, 150, 200 mg) until TCD of 500 mg, followed by a 2 day therapeutic course of AX 500 mg/8-h at home. For cross-reactivity analysis, if ST was negative, CX was orally administered (50, 100,
150, 200 mg) until TCD of 500 mg, followed by a 2 day therapeutic course of CX 500 mg/8-h. Finally, CO was administered following this procedure. Patients were carefully monitored during DPT and for 2 h after the last dose, complete equipment for cardiopulmonary resuscitation was available [36].

**In vitro sIgE determination by radioallergosorbent test (RAST)**
It was done using BP and AX conjugated to Poly-L-Lysine (PLL) (Sigma, St. Louis, MO) resulting in BPO-PLL and AXO-PLL in the solid phase, as described [37, 38], and radiolabeled anti-IgE antibody (kindly provided by Thermo Fisher Scientific and radiolabelled in our laboratory) [28]. Samples were considered positive if they were higher than 2.5% of label uptake, which was the mean + 2SD of a negative control group.

**Synthesis of chemical structures**
The molecule 1 (HOPhG-Ser-Bu) (Fig. 3a) was synthesised as described [30].

The molecule 2 (pyrazinone) (Fig. 3a) was synthesised following the Ugi/Desprotect/Cyclize strategy (Fig. 3b) [39], adapting protocols from cefaclor pyrazinone synthesis [28]. The synthetic methodology and characterisation are shown in Fig. 3 below.

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**Fig. 3** a Degradation hypothesis of cefadroxil (CX) after nucleophilic opening of betalactam ring by protein amino groups, leading to the cephalosporoyl intermediate, which degrades through dihydrotiazine fragmentation, and leading different functionality at carbon 6, hydroxyl and aldehyde respectively, and eventually resulting in the proposed antigenic determinants. Those equivalent synthetic structures for further immunological evaluation are represented in the square. b Synthesis of pyrazinone (molecule 2), pyrazin-2(1H)-one, proposed as CX determinant, through Ugi/Deprotect/Cyclize strategy.
of the pure compound can be found in this article’s Additional file 1.

**RAST inhibition assay**

This was done as described [38], incubating sera from patients with RAST values higher than 7% with different BLs (AX, CX, and CO) and the synthetic determinants of CX (1 and 2) in two ten-fold decreasing concentrations (100 mM and 10 mM) for 18 h at room temperature. After this, the AXO-PLL disc was added, followed by the previous described RAST procedure. The results were expressed as percentage inhibition with respect to the non-inhibited serum. Comparison of the inhibition capacity of the different inhibitors was made at 50% inhibition.

**Statistical analysis**

Description of quantitative variable included the median, mean, standard deviation (SD), and interquartile range (IR). Differences in percentage between the groups were compared by Chi-square analysis, numeric demographic data by Student t test. Comparisons for variables without a normal distribution were performed by the Mann–Whitney test for non-related samples and by Friedman test for related samples. All statistical analyses were done using the software package GraphPad PRISM v7. A P < 0.05 was considered statistically significant.

**Results**

From 1393 patients with confirmed BL hypersensitivity evaluated from 1984 to 2019, 994 subjects were confirmed with IHRS to AX, from which we randomly selected 54 patients, whose cross-reactivity to CX and CO was evaluated and flow-charts analysed (Fig. 2). The mean age was 41.7 ± 12.04 years; 35 (64.8%) were males; 51 (94.4%) had 1 episode and 3 (5.6%) two; in 32 (56.1%) episodes AX-CLV was the culprit and in 25 (43.9%) AX. Allergological work-up was confirmed with IHRS to AX, from which we randomly selected 54 patients, whose cross-reactivity to CX and CO was evaluated and flow-charts analysed. The analysis of sIgE results indicated that the mean value of mean levels to AXO-PLL and the percentage of positive cases was 1.12 ± 3.65, with 4 out of 48 (8.3%) cases positive to BPO-PLL and 24 out of 48 (50%) to AXO-PLL (Table 2). Comparisons between groups A and B showed higher differences, although not discriminating, in terms of mean levels to AXO-PLL and the percentage of positive cases (76.5% vs 35.5%) for AXO-PLL in Group B (p = 0.038 and p = 0.007, respectively).

| Case |boys | girls | total | mean age | sex ratio |
|------|------|-------|-------|----------|-----------|
| P1   | 56.0 | 44.0  | 100.0 | 31.2     | 0.961     |
| P5   | 56.0 | 44.0  | 100.0 | 31.2     | 0.961     |
| P19  | 56.0 | 44.0  | 100.0 | 31.2     | 0.961     |
| P26  | 56.0 | 44.0  | 100.0 | 31.2     | 0.961     |

**Significant differences of recognition are only found at the lower concentration of cefadroxil**

The analysis of sIgE results indicated that the mean value of RAST to BPO-PLL and AXO-PLL was 1.12 ± 3.65 and 6.8 ± 9.4 respectively, with 4 out of 48 (8.3%) cases positive to BPO-PLL and 24 out of 48 (50%) to AXO-PLL (Table 2). Comparisons between groups A and B showed higher differences, although not discriminating, in terms of mean levels to AXO-PLL and the percentage of positive cases (76.5% vs 35.5%) for AXO-PLL in Group B (p = 0.038 and p = 0.007, respectively).

To study CX specific recognition, we performed RAST inhibition assays on 6 cases from each group (Fig. 4a).
Table 1  Demographic and clinical data of patients included in the study

| Pat | Group | Sex | Age | Drug      | Epi | Reaction           | IDR (min) | IRS (d) |
|-----|-------|-----|-----|-----------|-----|--------------------|----------|---------|
| 1   | A     | M   | 55  | AX-CLV    | 1   | Anaphylaxis        | 60       | 180     |
| 2   | B     | M   | 44  | AX-CLV    | 1   | Anaphylaxis        | 20       | 60      |
| 3   | A     | M   | 46  | AX        | 1   | Anaphylaxis        | 50       | 30      |
| 4   | B     | M   | 55  | AX        | 1   | Urticaria          | 30       | 300     |
| 5   | A     | M   | 62  | AX-CLV    | 1   | Anaphylaxis        | 60       | 90      |
| 6   | B     | F   | 45  | AX-CLV    | 1   | Anaphylaxis        | 10       | 30      |
| 7   | B     | M   | 24  | AX        | 1   | Anaphylaxis        | 10       | 30      |
| 8   | B     | F   | 45  | AX        | 1   | Anaphylaxis        | 20       | 90      |
| 9   | A     | M   | 47  | AX        | 2   | Urticaria          | 40       | 477     |
|     |       |     |     |           |     | Anaphylaxis        | 10       | 365     |
| 10  | A     | F   | 46  | AX-CLV    | 1   | Anaphylaxis        | 30       | 120     |
| 11  | A     | M   | 43  | AX-CLV    | 1   | Anaphylaxis        | 45       | 60      |
| 12  | B     | F   | 40  | AX/AX-CLV | 2   | Urticaria          | 30       | 112     |
|     |       |     |     |           |     | Anaphylaxis        | 5        | 109     |
| 13  | A     | F   | 16  | AX-CLV    | 1   | Urticaria          | 60       | 30      |
| 14  | A     | M   | 27  | AX-CLV    | 1   | Anaphylactic shock | 5        | 21      |
| 15  | A     | M   | 66  | AX        | 1   | Anaphylaxis        | 60       | 365     |
| 16  | B     | M   | 44  | AX        | 1   | Anaphylaxis        | 10       | 365     |
| 17  | A     | F   | 50  | AX-CLV    | 1   | Anaphylaxis        | 15       | 30      |
| 18  | A     | M   | 44  | AX-CLV    | 1   | Anaphylaxis        | 10       | 280     |
| 19  | A     | M   | 25  | AX-CLV    | 1   | Urticaria          | 60       | 120     |
| 20  | B     | F   | 33  | AX-CLV    | 1   | Anaphylaxis        | 15       | 6       |
| 21  | B     | F   | 36  | AX-CLV    | 1   | Anaphylaxis        | 30       | 60      |
| 22  | A     | M   | 30  | AX        | 1   | Anaphylaxis        | 5        | 20      |
| 23  | B     | M   | 45  | AX-CLV    | 1   | Anaphylactic shock | 5        | 30      |
| 24  | A     | M   | 30  | AX-CLV    | 1   | Anaphylaxis        | 30       | 210     |
| 25  | B     | M   | 57  | AX-CLV    | 1   | Anaphylaxis        | 5        | 60      |
| 26  | A     | M   | 49  | AX        | 1   | Anaphylaxis        | 10       | 120     |
| 27  | A     | F   | 39  | AX        | 1   | Urticaria          | 40       | 40      |
| 28  | A     | F   | 39  | AX-CLV    | 1   | Urticaria          | 60       | 95      |
| 29  | A     | F   | 21  | AX-CLV    | 1   | Anaphylaxis        | 15       | 230     |
| 30  | A     | M   | 46  | AX-CLV    | 1   | Anaphylaxis        | 2        | 137     |
| 31  | B     | M   | 49  | AX        | 1   | Anaphylaxis        | 5        | 180     |
| 32  | B     | F   | 37  | AX-CLV    | 1   | Anaphylaxis        | 20       | 30      |
| 33  | A     | F   | 57  | AX        | 1   | Anaphylaxis        | 30       | 90      |
| 34  | B     | M   | 26  | AX        | 1   | Anaphylaxis        | 20       | 10      |
| 35  | B     | M   | 48  | AX-CLV    | 1   | Anaphylaxis        | 60       | 146     |
| 36  | A     | M   | 23  | AX-CLV    | 1   | Anaphylaxis        | 50       | 60      |
| 37  | B     | F   | 42  | AX        | 1   | Anaphylaxis        | 5        | 120     |
| 38  | B     | M   | 57  | AX-CLV    | 1   | Anaphylaxis        | 30       | 28      |
| 39  | B     | M   | 30  | AX-CLV    | 1   | Urticaria          | 10       | 730     |
| 40  | A     | F   | 37  | AX-CLV    | 1   | Anaphylaxis        | 5        | 30      |
| 41  | A     | F   | 14  | AX-CLV    | 1   | Anaphylaxis        | 10       | 90      |
| 42  | A     | M   | 44  | AX        | 1   | Urticaria          | 30       | 120     |
| 43  | A     | M   | 28  | AX        | 2   | Anaphylaxis        | 20       | 230     |
| 44  | A     | M   | 63  | AX-CLV    | 1   | Anaphylaxis        | 10       | 30      |
| 45  | A     | M   | 51  | AX        | 1   | Urticaria          | 60       | 30      |
| 46  | A     | M   | 35  | AX        | 1   | Urticaria          | 30       | 90      |
| 47  | A     | M   | 32  | AX        | 1   | Anaphylaxis        | 45       | 180     |
As inhibitors, we included AX, CX, and CO at two concentrations, 10 and 100 mM (Fig. 4b). Results with AX showed, as expected, a high percentage of inhibition at both concentrations in all cases. Regarding CX, the percentage of inhibition was above 50% in most of patients at 100 mM, 5 out of 6 patients in each group, similarly to levels obtained with AX. However, these percentages decrease at 10 mM, being lower than those observed with AX especially Group A (Fig. 4a). In fact, comparison analysis of the percentage of inhibition between groups only shows significant differences for CX at 10 mM ($p = 0.034$) (Fig. 4c). Only one case (P38, Group B) showed a percentage above 50% with CO.

Synthetic determinants of cefadroxil showed better sIgE recognition in Group B

The design of the two synthetic determinants of CX was based on our degradation hypothesis of the aminocephalexosporin-protein conjugate, using butylamine as a model nucleophile emulating protein lysine (Fig. 3a). After covalent protein conjugation through BL ring, the dihydrothiazine ring is unstable and could degrade producing structures in which carbon 6 presents different functionalities. Two relevant candidates, according to previous immunological recognition results [30], are structures bearing hydroxyl and aldehyde functionality in carbon 6. In the case of hydroxyl functionality, it would generate the molecule 1 as determinant; whereas the aldehyde functionality can react with the amino group of R1 side chain generating the pyrazinone 2 as a novel determinant. The synthesis of the molecule 2 was achieved following the Ugi/Deprotect/Cyclize strategy (Fig. 3b) [39]. First, starting reagents (an isocyanide, a protected amine, a protected aldehyde, and a N-protected aminoacid) were assembled by following the one-pot Ugi four-component reaction to produce the Ugi adduct. The latter acid-mediated-cleavage of the protected groups may result in the amino-functionalised aldehyde intermediate that cyclises, through intramolecular imine formation, and aromatises affording target pyrazinone (2). This method allowed the straightforward synthesis of 2, for which other procedures resulted unsuccessful. Compounds 1 and 2 were purified and well-characterised, allowing the immunological recognition study of precise chemical structures.

RAST inhibition assays were performed using CX and the two synthetic structures (1 and 2), as inhibitors (Fig. 5b), in two cases from Group A and 6 from Group B. There was no inhibition with these structures in Group A (Fig. 5a). Higher percentages of inhibition were observed in Group B, being greater than 50% in 4 out of 6 cases at 100 mM, in which similar levels of inhibition to those obtained with CX were observed (Fig. 5a). However, significant lower percentage of inhibition with these synthetic structures was observed performing the RAST inhibition at 10 mM ($p = 0.0022$ for both) (Fig. 5c).

**Discussion**

BLs are the most widely used antibiotics and the drugs most frequently involved in IHR [1] in adults and children [40–42]. All BL compounds can potentially induce a specific immunological response and, due to their wide prescription, BL allergy is nowadays a worldwide health issue with relevant implications [43–45]. One of the main issues is establishing the risk of developing an allergic reaction to cephalosporins prescribed in patients previously diagnosed of penicillin IHR, with different unsolved questions like if this risk can be predicted by ST and/or DPT, or the role of the chemical structure, specifically the side chain, in this recognition [10, 46–49]. The main difficulty is that, despite efforts [28–30], the antigenic determinants of cephalosporins are unknown [31]. In this study we have found that, for predicting cross-reactivity, ST is not enough and, R1 side chain, although critical for recognition, is not the unique factor. Moreover, the use of chemical tools for SAR study is a promising approach.

**Table 1 (continued)**

| Pat | Group | Sex | Age | Drug       | Epi | Reaction         | IDR (min) | IRS (d) |
|-----|-------|-----|-----|------------|-----|------------------|-----------|---------|
| 48  | A     | M   | 53  | AX         | 1   | Anaphylaxis      | 30        | 90      |
| 49  | A     | M   | 44  | AX         | 1   | Urticaria        | 45        | 120     |
| 50  | A     | F   | 38  | AX-CLV     | 1   | Anaphylaxis      | 10        | 30      |
| 51  | A     | M   | 54  | AX-CLV     | 1   | Anaphylactic shock | 5       | 90      |
| 52  | A     | F   | 62  | AX-CLV     | 1   | Urticaria        | 60        | 210     |
| 53  | B     | M   | 49  | AX-CLV     | 1   | Anaphylaxis      | 30        | 120     |
| 54  | A     | F   | 44  | AX         | 1   | Anaphylactic shock | 5       | 240     |

Patients were classified into Group A (Good tolerance to cefadroxil) or Group B (Cross-reactivity with cefadroxil)

Pat, Patients; M, Male; F, Female; AX-CLV, Amoxicillin-clavulanic; AX, Amoxicillin; Epi, Number of episodes; IDR (min), Interval drug administration and development of symptoms in minutes; IRS (d), Interval last reaction and allergological study in days

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Table 2  Skin tests and RAST results in patients from Group A (Good tolerance to cefadroxil) and Group B (Cross-reactivity with cefadroxil)

| Pat | Group | Skin test | AX | CX | CO | RAST | AX-PLL |
|-----|-------|-----------|----|----|----|-------|--------|
| 1   | A     | Neg       | Neg| Neg| Neg| ND    | ND     |
| 2   | B     | Neg       | Neg| SPT + (5 × 5)| ID + (2x2)| Neg| 0      | 3.42   |
| 3   | A     | Neg       | Neg| SPT + (5 × 5)| Neg| Neg| 0.15   | 3.24   |
| 4   | B     | Neg       | Neg| SPT + (4 × 5)| SPT + (4 × 4)| Neg| 0      |        |
| 5   | A     | Neg       | Neg| Neg| Neg| 0    | 3.42   |
| 6   | B     | Neg       | Neg| SPT + (5 × 5)| ID + (1x1)| Neg| ND     | ND     |
| 7   | B     | Neg       | Neg| SPT + (6 × 6)| SPT + (3 × 3)| Neg| 0.34   | 14.59  |
| 8   | B     | Neg       | Neg| SPT + (8 × 1)| SPT + (5 × 5)| Neg| 0      | 3.51   |
| 9   | A     | Neg       | Neg| SPT + (5 × 5)| Neg| Neg| 0      |        |
| 10  | A     | Neg       | Neg| IDT + (3 × 2)| Neg| Neg| 0      |        |
| 11  | A     | Neg       | Neg| IDT + (2 × 2)| Neg| Neg| 0      |        |
| 12  | B     | Neg       | Neg| IDT + (3 × 3)| IDT + (2x2)| Neg| 0.2    | 0.46   |
| 13  | A     | Neg       | Neg| IDT + (4 × 4)| Neg| Neg| ND     | ND     |
| 14  | A     | Neg       | Neg| SPT + (3 × 3)| Neg| Neg| 0      |        |
| 15  | A     | Neg       | Neg| IDT + (2 × 3)| Neg| Neg| ND     | ND     |
| 16  | B     | Neg       | Neg| SPT + (4 × 5)| IDT + (2x2)| Neg| 1.72   | 11.71  |
| 17  | A     | Neg       | Neg| IDT + (2 × 3)| Neg| Neg| 0      |        |
| 18  | B     | Neg       | Neg| IDT + (3 × 4)| Neg| Neg| 0      |        |
| 19  | A     | Neg       | Neg| Neg| Neg| ND   | ND     |
| 20  | B     | Neg       | Neg| IDT + (2 × 2)| IDT + (2x1)| Neg| 0      | 3.2    |
| 21  | B     | Neg       | Neg| SPT + (5 × 6)| IDT + (1x2)| Neg| ND     | ND     |
| 22  | A     | IDT + (2x1)| IDT + (1x1)| SPT + (3 × 3)| Neg| Neg| 3.2    | 6.79   |
| 23  | B     | Neg       | IDT + (2x2)| SPT + (6 × 6)| SPT + (3 × 3)| Neg| 23.54  | 29.83  |
| 24  | A     | Neg       | Neg| SPT + (2 × 3)| Neg| Neg| 0      | 3.03   |
| 25  | B     | Neg       | Neg| SPT + (5 × 6)| IDT + (2 × 2)| Neg| 1.22   | 7.55   |
| 26  | A     | Neg       | Neg| Neg| Neg| Neg| 0      |        |
| 27  | A     | Neg       | Neg| IDT + (3 × 3)| IDT + (2x2)| Neg| 0.2    | 0.46   |
| 28  | A     | Neg       | Neg| IDT + (3 × 4)| Neg| Neg| 2.15   | 8.32   |
| 29  | A     | Neg       | Neg| IDT + (4 × 4)| Neg| Neg| 0.09   | 0.54   |
| 30  | A     | Neg       | Neg| SPT + (5 × 6)| Neg| Neg| 0      | 1.14   |
| 31  | B     | Neg       | Neg| SPT + (5 × 5)| SPT + (2+3)| Neg| 0      | 22.54  |
| 32  | B     | Neg       | Neg| SPT + (3 × 3)| IDT + (2 × 3)| Neg| 0      | 15.56  |
| 33  | A     | Neg       | Neg| SPT + (4 × 5)| Neg| Neg| 1.28   | 2.31   |
| 34  | B     | Neg       | Neg| SPT + (6 × 6)| IDT + (2 × 3)| Neg| 1.87   | 3.36   |
| 35  | B     | Neg       | IDT + (2x2)| SPT + (7 × 8)| SPT + (3 × 4)| Neg| 6.15   | 26.09  |
| 36  | A     | Neg       | Neg| IDT + (2 × 2)| Neg| Neg| 0      | 25.56  |
| 37  | B     | Neg       | Neg| IDT + (1 × 2)| Neg| Neg| 0.37   | 0.13   |
| 38  | B     | Neg       | Neg| SPT + (5 × 6)| IDT + (3 × 4)| Neg| 1.65   | 21.8   |
| 39  | B     | Neg       | Neg| IDT + (2 × 2)| Neg| Neg| 0      | 0.92   |
| 40  | A     | Neg       | Neg| SPT + (2 × 3)| Neg| Neg| 0      | 0.7    |
| 41  | A     | Neg       | Neg| IDT + (5 × 6)| Neg| Neg| 0      | 1.93   |
| 42  | A     | Neg       | Neg| IDT + (2 × 2)| Neg| Neg| 0      | 0.37   |
| 43  | A     | Neg       | Neg| IDT + (3 × 3)| Neg| Neg| 0      | 1.24   |
| 44  | A     | Neg       | Neg| IDT + (4 × 5)| Neg| Neg| 0.23   | 0      |
| 45  | A     | Neg       | Neg| IDT + (2 × 2)| Neg| Neg| 0      | 23.2   |
| 46  | A     | Neg       | Neg| SPT + (4 × 4)| Neg| Neg| 0      |        |
| 47  | A     | Neg       | ID + (2x2)| SPT + (5 × 6)| Neg| Neg| 0.84   | 1.12   |
for elucidating the chemical structures involved in these IHRs.

In this study we have found that the cross-reactivity of IHRs to AX with CX, a cephalosporin with the same R1, was 35% and with CO, cephalosporin with different R1, figures decrease to 1.8%. Results with CX are in agreement with those by Romano [19] reporting that 39.3% of patients with IHR to penicillins had positive tests for cephalosporins, with 37.7% positive to aminoccephalosporins, including CX and/or cefamandole. These results are similar to our previous data, with 38% of cross-reactivity between AX and CX using DPT [21]. Relevantly, we did not detect differences in cross-reactivity to CX among those selective to AX compared to those reacting to the whole group, confirming that R1 is not the only factor influencing cross-reactivity. Regarding CO tolerance, all patients had negative ST and only one had a positive DPT, showing a high degree of CO tolerance, in line with previous data [16–19, 25].

Comparisons of SPT results to AX between Group A and B (cross-reactivity to CX) showed a higher percentage of positivity (78.9 vs 34.3%) in the group tolerant to CX (Group A). These results agree with those by Romano [19] estimating an odds ratio of ST positivity to ampicillin for cross-reacting to at least one cephalosporin of 2.5 (CI, 1.4–4.5). Moreover, the analysis of the sIgE results showed significantly higher levels and positivity to AXO-PLL in Group B. This seems to indicate that patients that cross-react with cephalosporins have a high degree of reactivity, taking into account that the two cases that developed systemic symptoms after ST with penicillins belong to Group B and that patients reacted to small amount of CX (50 and 150 mg) and CO (50 mg) in DPT.

Regarding the role of ST for predicting cross-reactivity, a positive cephalosporin ST in patients allergic to penicillins may indicate not only cross-reactivity but also concomitant sensitivity. Of note, P34 with cross-reaction to CO also reacted to CX. Whether this patient has cross-reacting or co-existing antibodies was something we cannot clarify in the present study as the RAST level was not enough for performing RAST inhibition with both drugs. However, cross-reactivity is more probable since this patient had not been previously treated with cefuroxime or any cephalosporin. This percentage is in agreement with previous data [18] that found 2.9% of cross-reactivity with CO in patients with prior histories involving only a penicillin. If a negative cephalosporin ST predicts good tolerance is controversial [50]. Different studies showed that patients with a well-established IgE-mediated allergy to penicillin and with ST negative to cephalosporins tolerate cephalosporins [15–17]. However, others demonstrated that less than 3% of cases can have a DPT positive with cephalosporin despite having negative ST [18, 19]. In this study 2 out of 37 patients (5.4%) with ST negative to CX and 1 out of 54 patients (1.8%) with ST negative to CO had a positive DPT to CX and CO respectively, indicating a negative predictive value (NPV) of 94.6% for CX and 98.1% for CO. That means that although NPV are high, a negative ST does not mean tolerance even if R1 are different.

Our immunological study by RAST inhibition assays agrees with previous results on cross-reactivity between penicillins and cephalosporins, showing that AX presented a better recognition, followed by CX [21]. Data showed a discriminating capacity of the test between Group A and B using lower drug concentrations, 10 mM, observing a significantly lower recognition of CX in patients with good tolerance to CX (Group A).

Regardless of this discriminative capacity, these data indicate that, although important for IgE recognition, the R1 is not the only structure involved in the immunological response, as structural modifications or some fragments of the nuclear structure may be involved in the antigenic determinant. In penicillins, the penicilloyl

| Pat | Group | Skin test | RAST AX-PLL |
|-----|-------|-----------|-------------|
|     |       | PPL/BP-OL | MDM/MD AX | CX | CO | BPO-PLL |
| 48  | A     | ID+(2x2)  | Neg | SPT+(4x5) | Neg | Neg | 8.13 | 13.87 |
| 49  | A     | Neg      | Neg | IDT+(2x2) | Neg | Neg | 0    | 0.14  |
| 50  | A     | Neg      | Neg | SPT+(5x4) | Neg | Neg | 0.54 | 0.41  |
| 51  | A     | Neg      | Neg | IDT+(5x7) | Neg | Neg | 0    | 14.68 |
| 52  | A     | Neg      | Neg | IDT+(3x4) | Neg | Neg | 0    | 7.93  |
| 53  | B     | Neg      | Neg | SPT+(4x5) | Neg | Neg | 0    | 16.84 |
| 54  | A     | Neg      | Neg | IDT+(3x2) | Neg | Neg | 0.06 | 33.02 |

Pat, Patients; PPL/BPO-OL, Benzylpenicilloyl-poly-L-lysine/benzylpenicilloyl-octa-L-lysine; MDM/MD, Minor determinant mixture/minor determinant; AX, Amoxicillin; CX, Cefadroxil; CO, Cefuroxime; BPO-PLL, Benzylpenicilloyl-poly-L-lysine; AXO-PLL, Amoxicilloyl-poly-L-lysine; SPT, Skin prick test; IDT, Intradermal test; Neg, Negative; ND, Not done
structure formed after protein conjugation is stable and, therefore, the thiazolidine ring could also play a role in the antigenic determinant [51–53]. On the contrary, the equivalent cephalosporyl structure is unstable, thus the R2 substituent is expelled [54, 55] and the dihydrothiazine ring suffers different fragmentations, producing a complex mixture in which structures are difficult to elucidate [29, 31]. We have addressed this issue, by using chemical tools, for performing SAR studies in which precisely defined structures, consisting on the R1 side chain coupled to the open BL ring with the carbon 6 of the original drug represented by a methyl group, were recognised by sIgE from patients with IHR to the cephalosporin containing either the same R1 or one structurally similar [29]. Further SAR studies involved similar synthetic determinants but with different functionalisation in such carbon 6, finding that hydroxymethyl and aldehyde functionality, compared with methyl group, increased recognition [30]. Based on these results, synthetic determinants of CX, involving the whole intact R1 or a modified R1 side chain, have been immunologically evaluated, showing higher-recognition by sIgE from patients cross-reactive to CX (Group B).

The structure 1 (HPhG-Ser-Bu), consisting on the R1 side chain of CX and open BL ring with hydroxymethyl functionality at carbon 6 [30], was not previously evaluated with sIgE to aminocephalosporins. These determinants containing the intact corresponding

| Pat | Group | Drug | Reaction | IDR (min) | TCD (mg) |
|-----|-------|------|----------|----------|---------|
| 1   | A     | AX   | Urticaria| 160      | 500     |
| 2   | B     | PV/CO| Good tolerance | – | – |
| 3   | A     | PV/CO| Good tolerance | – | – |
| 4   | B     | PV/CO| Good tolerance | – | – |
| 5   | A     | AX   | Anaphylaxis| 15       | 50      |
| 6   | B     | PV/CO| Good tolerance | – | – |
| 7   | B     | PV/CO| Good tolerance | – | – |
| 8   | B     | PV/CO| Good tolerance | – | – |
| 9   | A     | PV/CO| Good tolerance | – | – |
| 10  | A     | PV/CO| Good tolerance | – | – |
| 11  | A     | PV/CO| Good tolerance | – | – |
| 12  | B     | PV/CO| Good tolerance | – | – |
| 13  | A     | PV/CO| Good tolerance | – | – |
| 14  | A     | PV/CO| Good tolerance | – | – |
| 15  | A     | PV/CO| Good tolerance | – | – |
| 16  | B     | PV/CO| Good tolerance | – | – |
| 17  | A     | PV/CO| Good tolerance | – | – |
| 18  | A     | PV/CO| Good tolerance | – | – |
| 19  | A     | AX   | Urticaria| 140      | 500     |
| 20  | B     | PV/CO| Good tolerance | – | – |
| 21  | B     | PV/CO| Good tolerance | – | – |
| 22  | A     | CX/CO| Good tolerance | – | – |
| 23  | B     | CO   | Good tolerance | – | – |
| 24  | A     | PV/CO| Good tolerance | – | – |
| 25  | B     | PV/CO| Good tolerance | – | – |
| 26  | A     | AX   | Generalized pruritus and erythema| 60 | 150 |
| 27  | A     | PV/CO| Good tolerance | – | – |
| 28  | A     | PV/CO| Good tolerance | – | – |
| 29  | A     | PV/CO| Good tolerance | – | – |
| 30  | A     | PV/CO| Good tolerance | – | – |
| 31  | B     | PV/CO| Good tolerance | – | – |
| 32  | B     | PV/CO| Good tolerance | – | – |
| 33  | A     | PV/CO| Good tolerance | – | – |
| 34  | B     | PV/CO| Good tolerance | – | – |
| 35  | B     | CO   | Urticaria| 25       | 50      |
| 36  | A     | PV/CO| Good tolerance | – | – |
| 37  | B     | PV/CO| Good tolerance | – | – |
| 38  | B     | PV/CO| Good tolerance | – | – |
| 39  | B     | PV/CO| Good tolerance | – | – |

Time Interval and Total cumulative dose drug administration and the development of symptoms

Pat, Patients; AX, Amoxicillin; PV, Penicillin V; CX, Cefadroxil; CO, Cefuroxime; IDR (min), Interval between drug administration and development of symptoms in minutes; TCD, Total cumulative dose in mg

Table 3 (continued)
Aminocephalosporins R1 have been immunologically evaluated in a recent study with cefaclor-allergic patients (12% of positive cases) [28], and in the present study with AX- and/or CX-allergic patients (66% of positive cases at the maximum concentration), showing different extent of recognition depending on R1.

The pyrazinone 2 has been synthesised and immunologically evaluated in this study for the first time. Its structure derives from intramolecular reaction between the R1 amino group and the aldehyde at carbon 6. Inhibition results in six cases of Group B show that the pyrazinone 2, at 100 mM concentration, is recognised in 66% of cases, in agreement with IgE recognition observed for pyrazinones derived from cefaclor, with 63% of positive cases for the equivalent pyrazinone to that described here [28], and 60% of patients for an equivalent analog developed by Venemalm [32].

These synthetic determinants (1 and 2) were not recognised by the two selected patients with tolerance to CX (Group A). Importantly, greater differences in recognition...
between CX and the synthetic structures were observed in Group A than in Group B, using the higher concentration.

One could think that AX presents the amino group in R1 for the formation of additional determinants, as dike-topiperazin, considered as a minor determinant of AX [56]. However, it did not show sIgE recognition in previous studies [57], which is consistent with its lack of reactivity with proteins [56].

**Conclusions**

We have confirmed that cross-reactivity between penicillin and cephalosporins occurs when the R1 side chain is identical as previously reported, and that negative ST is not enough for predicting tolerance, being DPT necessary. The primary determinant of immunochemical recognition of aminoccephalosporins rested, with the structure of the R1, intact (molecule 1) or in its cyclised form as pyrazinone (molecule 2), although other parts of the molecule (excluding R2 substituents and most of the dihydrothiazine) are necessary for the formation of...
the antigenic determinant. These structures represent useful and safe alternatives for determining in vitro cross-reactivity to CX in AX-allergic patients. We think that other determinants, involving different patterns of recognition, could also participate in CX-allergic reactions; and more research is needed in this regard.

Supplementary Information

The online version contains supplementary material available at https://doi.org/10.1186/s13246-020-00368-1.

Additional file 1: Figure S1. Nuclear Magnetic Resonance (NMR) characterization of structure 2. (A) 1H-NMR (CH3OD)spectrum, (B) 13C-NMR (CH3OD) spectrum, and (C) heteronuclear single quantum coherence (HSQC) experiment with gradient pulse. Bidimensional NMR spectrum (left) and signal assignation (right).

Abbreviations
AX: Amoxicillin; AXO: Amoxicilloyl; BL: Betalactam; BP: Benzylpenicillin; BPO: Benzylpenicilloyl; BP‑OL: Benzylpenicilloyl‑octa‑L‑lysine; CO: Cefuroxime; CX: Cefadroxil; DPT: Drug provocation test; EAACI: European Academy of Allergy and Clinical Immunology; IDT: Intradermal tests; IHR: Immediate hypersensitivity reactions; IN: Interquartile range; MD: Minor determinant; MDM: Minor determinant mixture; NPV: Negative predictive value; PPL: Benzylpenicilloyl; SAR: Structure–activity relationship; sIgE: Specific IgE; SD: Standard deviation; SPT: Skin prick test; TCD: Total cumulative dose.

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Authors’ contributions
Concepts, Design, Definition of intellectual content—GB, CM, MIM, MJT. Literature search—GB, CM, MIM, MJT. Clinical studies: GB, EB, TP, MS, MIT. Clinical data analysis—GB, CM, TDF, MS, MIT. Immunoassays, experiments and data analysis—CM, AM‑S, RF‑S, AA, IMI‑S, MIM. Chemical synthesis and structural elucidation—AM‑S, MIM. Statistical analysis—CM, TDF, MIT. Manuscript preparation and editing—GB, CM, MIM, MIT. All authors have read and approved the final manuscript.

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Availability of data and materials
All data generated or analysed during this study are included in this published article and its supplementary Additional file 1.

Ethics approval and consent to participate
The studied group was obtained from the Regional University Hospital of Málaga Drug Allergy Database. The study was approved by the institutional review board, and informed consent for all procedures was obtained from all patients.

Competing interests
The group collaborates in research grants with Diater Laboratories (Madrid, Spain). The authors declare no other relevant conflicts of interest.

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References
1. Doña I, Barrionuevo E, Blanca‑López N, Torres M, Fernandez T, Mayorga C, et al. Trends in hypersensitivity drug reactions: more drugs, more response patterns, more heterogeneity. J Investig Allergol Clin Immunol. 2014;24(3):143–53.
2. Demoly P, Adkinson NF, Brockow K, Castells M, Chiriac AM, Greenberger PA, et al. International consensus on drug allergy. Allergy. 2014;69(4):420–37.
3. Torres MJ, Celik GE, Whitaker P, Atanaskovic‑Markovic M, Barbaud A, Bircher A, et al. A EAACI drug allergy interest group survey on how European allergy specialists deal with beta‑lactam allergy. Allergy. 2019;74(6):1052–62.
4. Sánchez‑Gómez FJ, González‑Moreno JM, Vida Y, Pérez‑Inestrosa E, Blanca M, Torres MI, et al. Amoxicillin haptenates intracellular proteins that can be transported in exosomes to target cells. Allergy. 2017;72:385–96.
5. Ariza A, Fernandez TD, Mayorga C, Barbero N, Martin‑Serrano A, Perez‑Sala D, et al. Hypersensitivity reactions to beta lactams: relevance of the hapten‑protein conjugates. J Investig Allergol Clin Immunol. 2015;25(1):12–25.
6. Montañez MI, Ariza A, Mayorga C, Fernandez T, Torres M. Cross‑reactivity in beta‑lactam allergy: alternative treatments. Current Treatment Options in Allergy. 2015;2:141–54.
7. Martín‑Serrano A, Barbero N, Agundez JA, Vida Y, Pérez‑Inestrosa E, Montañez MI. New advances in the study of IgE drug recognition. Curr Pharm Des. 2016;22:1–14.
8. WHO report on surveillance of antibiotic consumption. 2016–2018 early implementation. : Geneva: World Health Organization. Licence: CC BY‑NC‑SA 3.0 IGO ; 2018.
9. Romano A, Atanaskovic‑Markovic M, Barbaud A, Bircher AJ, Brockow K, Caubet JC, et al. Towards a more precise diagnosis of hypersensitivity to beta‑lactams—an EAACI position paper. Allergy. 2019. https ://doi.org/10.1111/all.14122.
10. Zagursky RJ, Pichichero ME. Cross‑reactivity in β‑Lactam allergy. J Allergy Clin Immunol Pract. 2018;6(1):72‑81.e1.
11. Renaudin JM, Beaudouin E, Porvert C, Demoly P, Moneret‑Vautrin DA. Severe drug‑induced anaphylaxis: analysis of 333 cases recorded by the Allergy Vigilance Network from 2002 to 2010. Allergy. 2013;68(7):929–37.
12. Pichler WJ. Immune pathomechanism and classification of drug hypersensitivity. Allergy. 2019;74(8):1457–71.
13. Kowalski ML, Agache I, Bavbek S, Bakirtas A, Blanca M, Bochenek G, et al. Diagnosis and management of NSAID‑exacerbated Respiratory Disease (N‑ERD)—a EAACI position paper. Allergy. 2019. https ://doi. org/10.1111/all.14122.
14. Kowalski ML, Agache I, Bavbek S, Bakirtas A, Blanca M, Bochenek G, et al. Diagnosis and management of NSAID‑exacerbated Respiratory Disease (N‑ERD)—a EAACI position paper. Allergy. 2019;74(1):28–39.
15. Brockow K, Ardern‑Jones MR, Mockenhaupt M, Aberer W, Barbaud A, Caubet JC, et al. EAACI position paper on how to classify cutaneous manifestations of drug hypersensitivity. Allergy. 2019;74(1):14–27.
16. Audicana M, Bernaola G, Urrutia I, Echechica S, Gastaminza G, Muñoz D, et al. Allergic reactions to beta lactams: studies in a group of patients allergic to penicillin and evaluation of cross‑reactivity with cephalosporin. Allergy. 1994;49(2):108–13.
16. Novalbos A, Sastre J, Cuesta J, De Las HW, Lluch-Bernal M, Bombín C, et al. Lack of allergic cross-reactivity to cephalosporins among patients allergic to penicillins. Clin Exp Allergy. 2001;31(3):438–43.

17. Romano A, Guéant-Rodriguez R-M, Viola M, Pettinato R, Guéant J-L. Cross-reactivity and tolerability of cephalosporins in patients with immediate hypersensitivity to penicillins. Ann Intern Med. 2004;141(1):16–22.

18. Caimmi S, Gelera C, Bousquet-Rouanet L, Arnoux B, Demoly P, Bousquet P. Safety of cefuroxime as an alternative in patients with a proven hypersensitivity to penicillins: a DAHD cohort survey. Int Arch Allergy Immunol. 2010;153(1):53–60.

19. Romano A, Valluzzi RL, Caruso C, Maggioletti M, Quarantino D, Gaeta F. Cross-reactivity and tolerability of cephalosporins in patients with IgE-mediated hypersensitivity to penicillins. J Allergy Clin Immunol Pract. 2018;6(5):1662–72.

20. Sastre J, Quijano LD, Novalbos A, Hernandez G, Cuesta J, delas Heras M, et al. Clinical cross-reactivity between amoxicillin and cephalaxin in patients allergic to amoxicillin and with good tolerance of penicillin. Allergy. 1996;51(6):383–6.

21. Miranda A, Blanca M, Vega JM, Moreno F, Carmona MJ, García JJ, et al. Cross-reactivity between a penicillin and a cephalosporin with the same side chain. J Allergy Clin Immunol. 1996;98(3):671–7.

22. Romano A, Valluzzi RL, Caruso C, Zaffiro A, Quarantino D, Gaeta F. Tolerability of cefazolin and cefitubein in patients with IgE–. J Allergy Clin Immunol. 2009. https://doi.org/10.1016/j.jaci.2009.02.025.

23. Romano A, Mayorga C, Torres MJ, Artesani MC, Suau R, Sanchez F, et al. Immediate allergic reactions to cephalosporins: cross-reactivity and selective responses. J Allergy Clin Immunol. 2000;106(6):1177–83.

24. Antunez C, Blanca-Lopez N, Torres MJ, Mayorga C, Perez-Inestrosa E, Montañez MI, Fernandez T, Blanca M. Immediate allergic reactions to cephalosporins: evaluation of cross-reactivity with a panel of penicillins and cephalosporins. J Allergy Clin Immunol. 2006;117(2):404–10.

25. de Vicente J, Gamboa P, García-Lirio E, Irazabal B, Jáuregui I, Martínez MD, et al. Tolerance to cephalosporins and carbapenems in penicillin-allergic patients. J Investig Allergol. 2003;13(5):174–80.

26. Barbero N, Fernández-Santamaría R, Mayorga C, Martin-Serrano A, Salas M, Bogas G, et al. Identification of an antigenic determinant of clavulanic acid responsible for IgE-mediated reactions. Allergy. 2019;74(8):1490–501.

27. Martín-Serrano A, Mayorga C, Barrionuevo E, Pérez N, Romano A, Moreno E, et al. Design of an antigenic determinant of cefaclor: chemical structure–IgE recognition relationship. J Allergy Clin Immunol. 2020;145(4):1301–4.e4.

28. Sanchez-Sancho F, Perez-Inestrosa E, Suau R, Montañez MI, Mayorga C, Torres MJ, et al. Synthesis, characterization and immunochromatographic evaluation of cephalosporin antigenic determinants. J Mol Recognit. 2003;16:148–56.

29. Montañez MI, Mayorga C, Torres MJ, Ariza A, Blanca M, Perez-Inestrosa E. Synthetic approach to gain insight into antigenic determinants of cephalosporins: in vitro studies of chemical structure–IgE molecular recognition relationships. Chem Res Toxicol. 2011;24(5):706–17.

30. Perez-Inestrosa E, Suau R, Montañez MI, Rodriguez R, Mayorga C, Torres MJ, et al. Cephalosporin chemical reactivity and its immunological implications. Curr Opin Allergy Clin Immunol. 2005;5(4):323–30.

31. Venema LM, Pyrazinone conjugates as potential cephalosporin allergens. Bioorg Med Chem Lett. 2001;11(14):1869–70.

32. Doña I, Romano A, Torres MJ. Algorithm for betalactam allergy diagnosis. Allergy. 2019;74(9):1817–9.

33. Antunez C, Fernandez T, Blanca-Lopez N, Torres MJ, Mayorga C, Canto G, et al. IgE antibodies to betalactams: relationship between the triggering hapten and the specificity of the immune response. Allergy. 2006;61(8):940–6.

34. Azajal J, El Matougui A, Pérez-Rubio JM, Coelho A, Fernández F, Sotelo E. Multicomponent assembly of diverse pyrazin-2(1H)-one chemotypes. J Org Chem. 2013;78(9):4402–9.

35. Rubio M, Bousquet P, Gomez E, Romano A, Demoly P. Results of drug hypersensitivity evaluations in a large group of children and adults. Clin Exp Allergy. 2012;42(1):123–30.

36. Gomez ER, Brockow K, Kyucyu S, Saretta F, Mori F, Blanca-Lopez N, et al. Drug hypersensitivity in children: report from the pediatric task force of the EAACI Drug Allergy Interest Group. Allergy. 2016;71(2):149–61.

37. Torres MJ, Blanca M. The complex clinical picture of β-lactam hypersensitivity: penicillins, cephalosporins, monobactams, carbapenems, and clavams. Med Clin North Am. 2010;94(4):805–20.

38. Mayorga C, Fernandez TD, Montañez MI, Moreno E, Torres MJ. Recent developments and highlights in drug hypersensitivity. Allergy. 2019;74(12):2368–81.

39. Barnicoat AD, Hamid QA, Luke NG, Mims EA, Heatley RV, Donaldson GC, et al. Diagnostic testing for penicillin allergy: a survey of practices and perspectives. Allergy. 2020. https://doi.org/10.1111/all.14159.

40. Romano A, Gaeta F, Arribas Poves MF, Valluzzi RL. Cross-reactivity among beta-lactams. Curr Allergy Asthma Rep. 2016;16(3):24.

41. Picard M, Robitaille G, Karam F, Daigle JM, Bédard F, Biron É, et al. Cross-reactivity among cephalosporins: evaluation of cross-reactivity with a panel of penicillins and cephalosporins. J Allergy Clin Immunol. 2006;117(2):404–10.

42. Romano A, Valluzzi RL, Caruso C, Zaffiro A, Quarantino D, Gaeta F. Tolerability of cefazolin and cefitubein in patients with IgE-. J Allergy Clin Immunol. 2009. https://doi.org/10.1016/j.jaci.2009.02.025.

43. Blanca M, Romano A, Torres MJ, Fernandez J, Mayorga C, Rodriguez J, et al. Update on the evaluation of hypersensitivity reactions to betalactams. Allergy. 2009;64(2):183–93.

44. Ariza A, Garzon D, Abánades DR, de Rios V, Vistoli G, Torres MJ, et al. Protein haptenation by amoxicillin: High resolution mass spectrometry analysis and identification of target proteins in serum. J Proteomics. 2012;77:504–20.

45. Torres MJ, Montañez MI, Ariza A, Salas M, Fernandez TD, Barbero N, et al. The role of IgE recognition in allergic reactions to amoxicillin and clavulanic acid. Clin Exp Allergy. 2016;46(2):264–74.

46. Ariza A, Mayorga C, Salas M, Doña I, Martín-Serrano Á, Pérez-Inestrosa E, et al. The influence of the carrier molecule on amoxicillin recognition by specific IgE in patients with immediate hypersensitivity reactions to betalactams. Sci Rep. 2016;6:35113.

47. Ariza A, Garzon D, Abánades DR, de Rios V, Vistoli G, Torres MJ, et al. Protein haptenation by amoxicillin: High resolution mass spectrometry analysis and identification of target proteins in serum. J Proteomics. 2012;77:504–20.

48. Torres MJ, Montañez MI, Ariza A, Salas M, Fernandez TD, Barbero N, et al. The role of IgE recognition in allergic reactions to amoxicillin and clavulanic acid. Clin Exp Allergy. 2016;46(2):264–74.

49. Ariza A, Mayorga C, Salas M, Doña I, Martín-Serrano Á, Pérez-Inestrosa E, et al. The influence of the carrier molecule on amoxicillin recognition by specific IgE in patients with immediate hypersensitivity reactions to betalactams. Sci Rep. 2016;6:35113.

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