Drug-induced anaphylactic reactions in children: A retrospective analysis of 159 validated spontaneous reports

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Abstract

Purpose: The main objective of this study was to analyze validated cases of drug-induced anaphylactic reactions in children with regard to incriminated drugs, clinical characteristics, and associated factors. A further objective was to compare differences in incriminated drugs and characteristics between validated cases and a reference excluding anaphylactic reaction cases (basic dataset).

Methods: Spontaneous reports of anaphylactic reactions in children (0-17 years) registered between January 2000 to December 2016 were extracted from the adverse drug reaction database of the German Federal Institute for Drugs and Medical Devices. These reports were restricted to drugs for which at least four cases were found. After case validation, 159 reports remained (validated dataset) and were compared with the basic dataset (n = 12,168 reports) using inferential statistics.

Results: Estimated yearly increase of reports (36.8 vs 0.1), most frequently incriminated drugs (antibiotics 30.2% vs 11%, analgesics/antipyretics 22.0% vs 5.6%; P values less than 0.001) and route of administration (38.4% vs 6.7%) differed between the validated dataset and the basic dataset. Validated cases differed in severity (higher with atracurium), reported symptoms (urticaria leading with analgesics), and associated factors (atopy/allergy rarely reported with antibiotics) depending on the incriminated drug class. In 13.8% (11.3% if excluding repeated readministration in one person) of the cases, the drug had not been tolerated before.

Conclusions: A heterogeneous clinical phenotype with differences in associated factors was observed, suggesting different underlying mechanisms triggered by the different drug groups. Occurrence of serious drug-induced anaphylactic reactions in children could be reduced by carefully considering patient history.

KEYWORDS
adverse drug reaction, anaphylactic reaction, anaphylaxis, atopy, pharmacoepidemiology, spontaneous reports
1 | INTRODUCTION

According to the allergy for global use nomenclature, anaphylaxis is defined as a severe, life-threatening generalized or systemic hypersensitivity reaction resembling an immediate-type reaction.2,3

The distal pathophysiological pathway in immune-mediated and non-immune-mediated anaphylaxis involves the release of mediators such as histamine, tryptase, and other bioactive mediators from basophils and mast cells.4

Drugs rank either second5-6 or third7-9 behind food and insect venoms as elicitors of anaphylaxis in children. One study reported an incidence of 0.5/100 000 person-years based on the clinical evaluation of these cases.10

Antibiotics, particularly beta-lactams, and nonsteroidal antiinflammatory drugs (NSAIDs) are reported as frequent elicitors of drug-induced anaphylaxis in children.11-15 However, these observations are based on a limited number of anaphylaxis cases in children (less than 100).

Some publications have reported atopy and allergy as risk factors for severe courses of anaphylaxis, whereas others have not.12,14,15,18 However, risk factors and cofactors may differ between age groups or according to the underlying pathophysiology and are not sufficiently studied in children.19

This paucity of data prompted us to further investigate drug-induced anaphylaxis in children on a larger scale and over a longer period of time (ie, 159 validated cases in 16 years) by exploring the adverse drug reaction (ADR) database of the German Federal Institute for Drugs and Medical Devices (BfArM).

The 242 reports were assessed by one of two (either B.S. or W.F) certified specialists in dermatology and allergology. Only cases in which (a) the correctness of the diagnosis “anaphylactic reaction” according to a national guideline3 and (b) the causal relationship with the incriminated drug according to WHO criteria27 was at least possible were considered for further analysis. Reports with only few symptoms or reports where symptoms were already transformed into the diagnosis “anaphylaxis” were also considered if

- respective treatment or treatment in an intensive/emergency care unit was reported,
- the patient had to be hospitalized,
- the event occurred under medical surveillance (eg, during anesthesia),
- the case was reported as life-threatening, or

December 2016 and originating from Germany (n = 14 508). Subsequently, we selected all anaphylactic reaction cases (n = 505) by application of the Standardized MedDRA Query (SMQ) “anaphylactic reaction” (version 19.1 as of September 2016).26 The 505 cases were restricted to reports where the “suspected/interacting” drug was reported more than three times in order to exclude influence by single reports. This resulted in 242 reports. All ADR reports coded as medication errors or with evidence of ADRs due to intentional suicide/self-injury were excluded by application of respective SMQs (pertain to each of the three datasets).

2 | MATERIAL AND METHODS

2.1 | BfArM’s ADR database

As described earlier,20,21 physicians in Germany are obliged by their professional conduct code to report ADRs to their professional councils, which forward these reports to either BfArM (responsible for chemically defined drugs)22 or Paul-Ehrlich-Institut (PEI) (responsible for monoclonal antibodies, vaccines, etc).23,24 These reports can also be reported directly to BfArM, PEI, or marketing authorization holders who then forward the reports to the authorities.

In BfArM’s ADR database, drugs are coded according to the World Health Organization (WHO) Drug Dictionary and the Anatomical Therapeutic Chemical (ATC) classification system.25 ADRs are coded using Medical Dictionary for Regulatory Activities (MedDRA) terminology.26

The data lock point of the present analysis was December 2016.

2.2 | Case identification

We identified all spontaneous ADR reports (no study reports) referring to children (0-17 years), registered between January 2000 and

KEY POINTS

- Only a few studies have investigated drug-induced anaphylactic reactions in children.
- The adverse drug reaction (ADR) database of the German Federal Institute for Drugs and Medical Devices provided the opportunity to examine this rare ADR on a larger scale.
- Intravenous administration was noted for 38% of incriminated drugs. In 13.8% of cases (11.3% if excluding repeated readministration in one person), previous hypersensitivity to the drug had been reported, and these cases appeared to be more severe than cases designated as “drug never used before.”
- Antibiotics, analgesics, and MRI contrast media were most frequently suspected of having induced the anaphylactic reaction in validated cases.
- Cefaclor accounted for 27% and amoxicillin for 8.3% of cases induced by antibiotics, although exposure to amoxicillin seems to outweigh cefaclor exposure.
the physician already had classified the anaphylactic reaction suggesting medical expertise concerning anaphylactic reactions.

For quality assurance, the final dataset was reviewed by a pharmacist. Eventually, the validated dataset consisted of 159 cases including 164 incriminated drugs (equal causal probability for two drugs in five cases). The analysis of the incriminated drugs and routes of administration referred to the 164 drugs, whereas all other analyses referred to the 159 cases (see Figure 1).

### 2.4 Quality of validated cases

The completeness of data in the validated cases was assessed according to a published score. Calculation of the score was modified as it was not computed for every reported drug-ADR pair (in case more than one ADR had been reported) and then aggregated to an average, to yield an overall score for the corresponding report. Instead, since our analysis focused on anaphylactic reactions, the calculation of the score referred only to the diagnosis anaphylactic reaction. A completeness score of 0.89 [0.81–0.95] was calculated (greater than 0.8, well-documented according to Bergvall et al). Most data in the variable dose (30.8% of reports) was missing.

### 2.5 Generation and comparison of additional datasets

In order to address the lack of a control group, we generated a reference group ("basic dataset") containing all other ADR reports on children 0 to 17 years excluding the 505 cases identified by the SMQ "anaphylactic reaction" (n = 12,168 reports). In addition, we created the "all-anaphylactic reactions" dataset in order to examine whether differences between the basic dataset and the validated dataset might have resulted from the validation process or from restriction to reports with drugs reported in more than three cases. This dataset was based on the 505 identified anaphylactic reaction cases and finally resulted in 472 reports. The same predefined inclusion and exclusion criteria of cases were applied for both datasets.

The three datasets were compared with regard to basic characteristics, incriminated drugs, and the seriousness criteria based on the legal (not clinical) definition, ie, outcome of the ADR is fatal, life-
threatening or leads to (prolonged) hospitalization, persistent or significant disabilities, or congenital anomalies/birth defects.\textsuperscript{29}

2.6 Analysis of the validated cases

Any analysis was based on the information provided in the complete report including narrative and follow-ups.

Cases were classified with regard to increasing severity (grade I-IV) according to a national guideline.\textsuperscript{3} Grade I reactions, for example, are characterized by cutaneous and subjectively perceived general symptoms only, whereas grade IV refers to cardiovascular and/or respiratory arrest (unclassifiable cases are denoted as NOS).

Cases were also analyzed concerning reported symptoms by analyzing assigned preferred terms\textsuperscript{26} and associated factors like atopy/allergy. Atopy is an individual susceptibility usually occurring in childhood to become sensitized and produce immunoglobulin E (IgE) antibodies in response to ordinary exposures to allergens. These individuals can develop allergic asthma, allergic rhinoconjunctivitis, or atopic dermatitis.\textsuperscript{1} No published algorithm to diagnose atopy was found. Hence, an individual was designated as atopic if either atopy or one of the following conditions was reported: atopic dermatitis/asthma/pollinosis, a total IgE greater than 100 kU/L, or IgE slightly elevated. A patient was designated as allergic if allergy (NOS or specified) was reported.

The classification “drug administered before” referred to the previous administration of drugs with the same active ingredient except in cases where excipients were also cosuspected (e.g., coloring agents or flavors). The classification “drug not tolerated before” referred to the occurrence of hypersensitivity-like symptoms after previous administration.

2.7 Statistical analysis

The descriptive analysis was carried out with means (±SD) (for age, estimated yearly increase, drugs per report) and frequency distributions with percentages (all other results). Because of unequal variances, Welch t test was performed to compare mean ages between drug subgroups and the remaining validated cases. For differences in frequency distributions between the two anaphylactic reactions datasets and the basic dataset and in the validated dataset between drug subgroups and the remaining cases (without the respective drug subgroup), the chi-square test was applied (in case of less than six cases: Fisher exact test).

3 RESULTS

3.1 Comparison of datasets

Table 1 shows the characteristics of the three datasets. The number of reports in the basic dataset increased by an average of 36 reports per year, whereas the annual number of validated cases remained stable with an average proportion of 1.4% (range: 0.7-2.2%) per year. The validated cases in comparison with the basic dataset more often reported the seriousness criteria life-threatening (23.3% vs 5.8%) or hospitalization (45.3% vs 30.0%) but less often death (0.6% vs 3.5%).

Female gender was more frequently reported in the validated than in the basic dataset (51.6% vs 43.4%). Gender differences were also noted depending on the drug administered (e.g., MRI contrast media [female gender] 73.7% vs 49.1%).

The drug classes most frequently suspected in the validated cases were less often reported in the basic dataset (antibiotics 30.2% vs 11%, analgesics/antipyretics 22.0% vs 5.6%; P values less than 0.001).

Intravenous administration was clearly more often reported in the validated compared with the basic dataset (38.4% versus 6.7%; P value less than 0.001, based on the number of suspected drugs) and differed depending on drug class.

For most parameters, larger (but similar) differences were observed between the validated and the basic dataset than between the all-anaphylactic and the basic dataset. However, the number of cases that reported the seriousness criterion death was larger in the all-anaphylactic (6.1%) than in the validated dataset (0.6%).

3.2 Analysis of validated cases

3.2.1 Demographic parameters

The mean age of validated cases was 8.9 years (SD = 5.4) (Table 2). Slightly more reports were found for preschoolers (≥3 to ≤6 years; 28.9%) and adolescents (≥16 to ≤17 years; 17.6%). Drug-related age and gender differences were observed, e.g., mean age: iron (14.7 years); gender: MRI contrast media (14 females vs five males).

These gender differences were also observed in the stratified age groups (female 0-5 years: 38.2%; female 13-17 years: 62.7%).

3.2.2 Classification and description of anaphylactic reactions

A total of 10.1% of the validated cases were classified as grade I, 67.3% as grade II, 17.0% as grade III, and 0.6% as grade IV. Grade I/II (moderate; 77.4%) and grade III/IV (severe; 17.6%) cases were pooled for subanalysis. More severe than moderate reactions were only reported in atracurium cases (Table 3).

The most frequently reported symptom was dyspnea (35.8%; 57/159 cases) followed by urticaria (33.3%; 53/159). Differences were noted for analgesics/antipyretics (urticaria: 40.0%) and for atracurium cases (anaphylactic shock: 60.0%) (Table 4). Urticaria (43.6%) was the leading symptom reported for the age class 0 to 5 years, whereas this was dyspnea for age classes 6 to 12 (32.7%) and 13-17 years (33.3%) (data not shown).

3.2.3 Atopy/allergy

Only 15.1% and 27.7% of the cases respectively yielded information on atopy (24/159) and allergy (44/159). A total of 13.8% (22/159) of the cases were designated as atopic, and allergy was determined in 18.2% (29/159) of the cases. In 23/29 of the allergy cases, specific information about the allergen was provided (pollen/house dust mites/animals [n = 13], food [nuts, milk, eggs, etc;
n = 9], antibiotics [n = 2], and hymenoptera [n = 1]) (some patients reported more than one allergen). Histamine intolerance was reported in one case. For subgroup analysis, the atopy cases (n = 22) and allergy cases (n = 29) were pooled (altogether 40 cases, since 11 cases reported atopy and allergy). This was considered reasonable since the reported allergens are common in immediate-type allergic reactions (eg, allergic rhinoconjunctivitis), which is also a characteristic of atopy.

### TABLE 1 Characterization of the three datasets

| Criteria                      | Spontaneous Reports from 2000 to 2016 Without Medication Errors and Intentional Overdose; Age: 0 to 17 years |
|-------------------------------|----------------------------------------------------------------------------------------------------------|
|                               | Basic dataset[^b^] (without anaphylactic reaction cases) (n = 12 168 cases)                          |
|                               | All-anaphylactic reactions dataset (determined by SMQ[^c^]) (n = 472 cases)                           |
|                               | Validated dataset (n = 159 cases)                                                                     |
| Estimated yearly increase (in cases ± SD) | y = 36.875 (±110.9)                                        | y = 0.0625 (±7.7)                                         | y = 0.0625 (±5.4)                                         |
| Number of suspected/interacting drugs[^d^] | 16 777                                                   | 576                                                       | 164                                                       |
| Drugs per report (±SD)         | 1.4 (0.4-2.4)                                           | 1.2 (0.5-1.9)                                             | 1.0 (0.8-1.2)                                             |
| Primary source                |                                                            |                                                           |                                                           |
| Physician                     | 61.1% (n = 7437)                                         | 67.4% (n = 318)                                           | 71.1% (n = 113)                                           |
| Consumer/non-HCP[^e^]          | 8.9% (n = 1084)                                          | 8.7% (n = 41)                                            | 5.7% (n = 9)                                             |
| Serious[^f^]                   | 82.5% (n = 10 041)                                       | 87.5% (n = 413)                                          | 88.0% (n = 140)                                          |
| Hospitalization               | 30.0% (n = 3647)                                         | 41.9% (n = 198)                                          | 45.3% (n = 72)                                           |
| Life-threatening               | 5.8% (n = 710)                                           | 22.0% (n = 104)                                         | 23.3% (n = 37)                                           |
| Death                         | 3.5% (n = 426)                                           | 6.1% (n = 29[^g^])                                  | 0.6% (n = 1)                                             |
| Mean age (years ± SD)         | 8.2 (2.0-14.4)                                           | 10.0 (4.4-15.6)                                      | 8.9 (3.5-14.3)                                           |
| Male                          | 50.2% (n = 6106)                                         | 48.7% (n = 230)                                         | 48.4% (n = 77)                                           |
| Female                        | 43.4% (n = 5278)                                         | 50.0% (n = 236)                                         | 51.6% (n = 82)                                           |
| Unknown                       | 6.4% (n = 784)                                           | 1.3% (n = 6)                                             |                                                           |
| Administration route[^h^]      |                                                            |                                                           |                                                           |
| Intravenous                   | 6.7% (n = 1121)                                          | 25.0% (n = 144[^i^])                                  | 38.4% (n = 63[^i^])                                  |
| Oral                          | 38.9% (n = 6519)                                         | 39.9% (n = 230)                                         | 39.6% (n = 65)                                           |
| Rectal                        | 0.8% (n = 139)                                           | 3.3% (n = 19)                                           | 4.3% (n = 7)                                             |
| Unknown                       | 21.2% (n = 3555)                                         | 19.4% (n = 112)                                         | 12.8% (n = 21)                                           |
| Analgesics (N02[^i^] and ibuprofen[^j^]) | 687 cases (5.6%)                                  | 56 cases (11.9%^[^k^])                                | 35 cases (22.0%^[^k^])                                |
| Mean age (years ± SD)         | 6.9 (0.7-13.1)                                           | 9.1 (4.2-14.0)                                           | 7.9 (3.2-12.6)                                           |
| Female                        | 40.8% (n = 280)                                         | 33.9% (n = 19)                                          | 34.3% (n = 12)                                           |
| Male                          | 52.0% (n = 357)                                         | 66.1% (n = 37)                                          | 65.7% (n=23)                                             |
| Unknown                       | 7.3% (n = 50)                                            |                                                           |                                                           |
| Antibiotics (J01[^i^])         | 1336 cases (11.0%)                                       | 89 cases (18.9%^[^k^])                                | 48 cases (30.2%^[^k^])                                |
| Mean age (years ± SD)         | 8.2 (2.2-14.2)                                           | 9.7 (4.0-15.4)                                          | 8.8 (3.4-14.2)                                           |
| Female                        | 48.1% (n = 643)                                         | 52.8% (n = 47)                                          | 54.2% (n = 26)                                           |
| Male                          | 48.1% (n = 643)                                         | 47.2% (n = 42)                                          | 45.8% (n = 22)                                           |
| Unknown                       | 3.7% (n = 50)                                            |                                                           |                                                           |
| Iron                           | 40 cases (0.3%)                                          | 9 cases (1.9%^[^k^])                                  | 7 cases (4.4%^[^k^])                                  |
| Mean age (years ± SD)         | 8.2 (1.6-14.8)                                           | 15.1 (11.3-18.9)                                        | 14.7 (10.4-19.0)                                         |
| Female                        | 60.0% (n = 24)                                          | 77.8% (n = 7)                                           | 71.4% (n = 5)                                           |
| Male                          | 25.0% (n = 10)                                          | 22.2% (n = 2)                                           | 28.6% (n = 2)                                           |
| Unknown                       | 15.0% (n = 6)                                            |                                                           |                                                           |
| Alglucosidase                  | 35 cases (0.3%)                                          | 12 cases (2.5%^[^k^])                                | 12 cases (7.5%^[^k^])                                |
| Mean age (years ± SD)         | 2.7 (–1.9-7.3)                                          | 3.3 (0.4-6.2)                                           | 3.3 (0.4-6.2)                                           |
| Female                        | 51.4% (n = 18)                                          | 33.3% (n = 4)                                           | 33.3% (n = 4)                                           |
| Male                          | 37.1% (n = 13)                                          | 66.7% (n = 8)                                           | 66.7% (n = 8)                                           |
| Unknown                       | 11.4% (n = 4)                                           |                                                           |                                                           |
| MRI (V08C[^i^])                | 57 cases (0.5%)                                          | 25 cases (5.3%^[^k^])                                | 19 cases (11.9%^[^k^])                                |
| Mean age (years ± SD)         | 12.0 (7.7-16.3)                                         | 12.1 (7.3-16.9)                                         | 11.5 (6.4-16.6)                                         |
| Female                        | 49.1% (n = 28)                                          | 72.0% (n = 18)                                          | 73.7% (n = 14)                                           |
| Male                          | 47.4% (n = 27)                                          | 28.0% (n = 7)                                           | 26.3% (n = 5)                                           |
| Unknown                       | 3.5% (n = 2)                                             |                                                           |                                                           |

(Continues)
Thirty-two (26.0%) of the pooled atopy/allergy cases were classified as grade I/II (n = 123) and n = 6 (21.4%) as grade III/IV (n = 28) reactions (two cases NOS).

The largest number of reports designated as atopic/allergic was observed in the analgesics/antipyretics drug class (42.9%; 15/35; \( P < 0.05 \)), followed by MRI contrast media (31.6%; 6/19) (Table 4), whereas only 14.6% (7/48; \( P < 0.05 \)) of the antibiotic cases were designated as atopic/allergic.

### 3.2.4 Drug-related findings

Table 5 shows the 10 drugs most frequently assessed as causal inducers.

Ibuprofen ranked first with 18.9% (30/159; 85.7% [30/35] of analgesic/antipyretic cases) and was observed more frequently in males (21 vs 9; \( P < 0.05 \)) and ages 0 to 12 years (86.7%). In 56.7% (17/30) of the reports, the drug had been administered orally. Of the oral formulations, 41.2% (7/17) contained flavors (eg, strawberry). Allergy/atopy was stated in 43.3% (13/30) of the reports.

Cefaclor ranked second and accounted for 52.0% (13/25) of the reports attributed to cephalosporins and for 27.1% (13/48) of the antibiotic cases. Of these cases, 46.2% (6/13) reported the seriousness criterion life-threatening (compared with 23.3% of all cases). Age-stratified analysis showed a larger number of reports for the ages 0 to 12 years (92.3%), and no gender differences were observed. None of the cefaclor cases reported allergy or atopy.

Three of five atracurium cases (rank 5) were classified as anaphylactic reactions grade III (1 grade IV fatal outcome), 1 NOS; four out of five of these cases were in males.

Four of seven iron-related cases referred to ferric carboxymaltose (intravenous; rank 6) and one case each to ferric gluconate (intravenous), ferric dextran (intravenous), and ferric sulfate (oral). In all cases, the reaction occurred within 30 minutes.

Four cases of anaphylactic reaction after intravenous corticosteroid therapy with asthma as comorbidity (rank 6) were identified.

Another four cases reported anaphylactic reactions (3/4 grade II, 1/4 NOS) after topical application of an ointment with the ingredients methyl nicotinate and Symphytum officinale (rank 6).

In 15.1% (24/159) of the reports, the drug had never been taken previously (Table 2). In 34.0% (54/159) of the cases, the drug had been given previously (not tolerated before: 40.7% [22/54]; 33.3% if excluding repeated readministration in one person); tolerated before: 44.4% [24/54]; unknown: 8/54. Cases reporting "not tolerated before" (13.8% of all cases [22/159] or 11.3% [18/159] if excluding repeated readministration in one person) were more often designated as severe (grade III/IV 22.7% vs 8.3%), life-threatening (36.4% vs 20.8%), and serious (100% vs 83.3%) than cases reporting "drug never used before."

### 4 DISCUSSION

The present study analyzed 159 validated cases of drug-induced anaphylactic reactions in children and compared this dataset with a...
### TABLE 2  Characterization of validated cases of anaphylactic reactions

| Category                        | All Validated Cases (n = 159) | Cases Attributed to Antibiotics (n = 48) | Cases Attributed to Analgesics/Antipyrretics (n = 35) | Cases Attributed to MRI Contrast media (n = 19) | Cases Attributed to Alglucosidase (enzymes) (n = 12) | Cases Attributed to Iron (n = 7) | Cases Attributed to Atracurium (n = 5) | All Other Cases (n = 36) |
|---------------------------------|------------------------------|----------------------------------------|-----------------------------------------------|-----------------------------------------------|-----------------------------------------------|---------------------------------|---------------------------------|--------------------------|
| Serious                         | 88.1% (140/159)              | 75.0% (36/48)                          | 100.0% (35/35)                                | 84.2% (16/19)                                 | 85.7% (6/7)                                   | 100.0% (5/5)                    | 91.7% (33/36)                  |
| Hospitalization                 | 45.3% (72/159)               | 43.8% (21/48)                          | 62.9% (22/35)                                 | 42.1% (8/19)                                  | 25.0% (3/12)                                  | 14.3% (1/7)                     | 40.0% (2/5)                    | 44.4% (16/36)              |
| Life-threatening                | 23.3% (37/159)               | 31.3% (15/48)                          | 22.9% (8/35)                                  | 5.3% (1/19)                                   | 8.3% (1/12)                                   | 14.3% (1/7)                     | 60.0% (3/5)                    | 27.8% (10/36)              |
| Mean age (years ± SD)          | 8.9 (3.5-14.3)               | 8.8 (3.4-14.2)                         | 7.9 (3.2-12.6)                                | 11.5 (6.4-16.6)                               | 3.3 (0.4-6.2)                                 | 14.7 (10.4-19.0)                | 9.4 (3.0-15.8)                 | 9.6 (4.3-14.9)            |
| Female                          | 51.6% (82/159)               | 54.2% (26/48)                          | 34.3% (12/35)                                 | 73.0% (14/19)                                 | 33.3% (4/12)                                  | 71.4% (5/7)                     | 20.0% (1/5)                    | 61.1% (22/36)              |
| Mean age (years ± SD)          | 8.9 (3.5-14.3)               | 8.8 (3.4-14.2)                         | 7.9 (3.2-12.6)                                | 11.5 (6.4-16.6)                               | 3.3 (0.4-6.2)                                 | 14.7 (10.4-19.0)                | 9.4 (3.0-15.8)                 | 9.6 (4.3-14.9)            |
| Male                            | 48.4% (77/159)               | 45.8% (22/48)                          | 65.7% (23/35)                                 | 26.3% (5/19)                                  | 66.7% (8/12)                                  | 28.6% (2/7)                     | 80.0% (4/5)                    | 38.9% (14/36)              |
| Mean age (years ± SD)          | 8.9 (3.5-14.3)               | 8.8 (3.4-14.2)                         | 7.9 (3.2-12.6)                                | 11.5 (6.4-16.6)                               | 3.3 (0.4-6.2)                                 | 14.7 (10.4-19.0)                | 9.4 (3.0-15.8)                 | 9.6 (4.3-14.9)            |
| Intravenous administration      | 39.6% (63/159)               | 20.8% (10/48)                          | 0%                                             | 78.9% (15/19)                                 | 100.0% (12/12)                                | 85.7% (6/7)                     | 80.0% (4/5)                    | 44.4% (16/36)              |

- **This table shows the validated cases (n = 159; validated dataset) stratified by drug class and seriousness criteria, age and gender, proportion of intravenous administration, and drug-specific history. In 48 antibiotic cases, 49 antibiotics (one case with cefotaxime and cefixim) were reported. One case reporting cefaclor and ibuprofen as suspected drugs was also counted for the drugs class analgesics. In 35 analgesic cases, 36 analgesics metoclopramide as suspected drugs and was therefore also counted in the group "all other cases.

- **One of the 5 atracurium reports included atracurium and propofol as suspected drugs and, thus, was also counted in the group "all other cases."

- **In 36 "all other cases," atracurium and propofol were reported as suspected drugs and, hence, were also counted for atracurium. One report included metamizole and metoclopramide as suspected drugs and was therefore also counted in the group analgesics.

- **A total of 159 case reports contained 164 suspected drugs. Cases with more than one drug were counted in each drug class. However, they were not counted twice if they belonged to the same drug class. Therefore, the sum of cases of all drug subgroups exceeds 159 cases.

- **Twelve case reports for alglucosidase. Among these 12 cases, there was one patient accounting for five cases (each at a different date). In these cases, there was no evidence that the reactions occurred in context with a desensitization procedure.

- **The "seriousness" assessment may not reflect the clinical severity of the reaction since they refer to the legal definition of the Medicinal Products Act: An adverse drug reaction (ADR) is considered serious when the ADR results in death, is life-threatening, requires in-patient hospitalization or prolongation of existing hospitalization, results in persistent or significant disability or incapacity, or is a congenital anomaly/birth defect. One case may contain more than one of these criteria.

- **One case with age unknown.

- **Since this table refers to the number of cases (n = 159), the calculation of percentages is also based on the number of cases per drug subgroup. The respective figures relating to the number of incriminated drugs (n = 164) are: all validated cases: 38.4% (63/164), antibiotics: 20.4% (10/49), analgesics/antipyrretics: 0%, MRI: 78.9% (15/19), alglucosidase 100% (12/12), iron: 85.7% (6/7), atracurium: 80.0% (4/5), and all other cases 44.4% (16/36).

- **The relative distributions if a "drug was tolerated" or "not tolerated" or "tolerated is unknown after previous administration" refer to the number reporting "drug administered before." The 13.8% (22/159) cases, which reported previous hypersensitivity to the administered drug, included repeated readministration (four times) in one patient (assigned to the drug subgroup alglucosidase). A total of 11.3% (18/159) of cases remained if these four reports were excluded.

- **Chi-squared test/Fischer exact test; P < 0.05. Further information on calculation of P values is included in Section 2.7."
TABLE 3 Classification of anaphylactic reactionsa

| Anaphylactic Reaction Grades | All Cases of Anaphylactic Reactions (n = 159) | Cases Attributed to Antibiotics n = 48 (30.2%) | Cases Attributed to Analgesics/Antipyretics n = 35 (22.0%) | Cases Attributed to MRI Contrast Media n = 19 (11.9%) | Cases Attributed to Alglucosidase n = 12 (7.5%) | Cases Attributed to Iron n = 7 (4.4%) | Cases Attributed to Atracurium (Muscle Relaxants) n = 5 (3.1%) | All Other Cases n = 36 (22.6%) |
|----------------------------|-----------------------------------------------|-----------------------------------------------|----------------------------------------------------------|------------------------------------------------|-----------------------------------------------|---------------------------------|------------------------------------------------|-----------------------------------------------|
| Anaphylactic reaction grades I-II (n = 123) | 77.4% of cases (123/159) | 66.7% (32/48)* | 80.0% (28/35) | 89.5% (17/19) | 91.7% (11/12) | 85.7% (6/7) | 0% (0/5) | 83.3% (30/36) |
| Anaphylactic reaction grades III-IV (n = 28) | 17.6% of cases (28/159) | 22.9% (11/48) | 17.1% (6/35) | 10.5% (2/19) | 8.3% (1/12) | 14.3% (1/7) | 80.0% (4/5)* | 13.9% (5/36) |
| Anaphylactic reaction NOS (n = 8) | 5.0% of cases (8/159) | 10.4% (5/48) | 2.9% (1/35) | 0% (0/19) | 0% (0/12) | 0% (0/7) | 20.0% (1/5) | 2.8% (1/36) |

aIn n = 8 cases, the anaphylactic reaction was classified as NOS (not otherwise specified). Only one out of 159 (0.6%) of cases (atracurium) had a fatal outcome. This table shows the stratification of the validated cases (n = 159, validated dataset) by drug class and assigned grade of anaphylactic reaction (moderate (grade I/II), severe (grade III/IV), classification not possible [NOS]).

bA total of 159 case reports contained 164 suspected drugs. Cases with more than one drug were counted in each drug class. However, they were not counted twice if they belonged to the same drug class.

cTwelve case reports for alglucosidase. Among these 12 cases there was one patient accounting for five cases (each at a different date). In these cases there was no evidence that the reactions occurred in context with a desensitization procedure.

*Chi-squared test/Fischer exact test; P < 0.05. Further information on the calculation of P values is included in Section 2.7.

4.1 | Comparison of datasets

The drugs most frequently suspected in the validated dataset compared with the basic dataset were antibiotics (30.2% vs 11.0%), analgesics/antipyretics (22.0% vs 5.6%), and MRI contrast media (4.4% vs 0%). These findings are also reported in other investigations.14

Hence, intravenous administration may entail a higher risk for anaphylactic reactions compared with the basic dataset (38.4% vs 6.7%). This finding is reassuring, but others have reported similar findings.12 A total of 164 suspected drugs were contained in 159 case reports, and it cannot be concluded whether it also applies in real life because of the limitations of the spontaneous reporting system. In contrast to the basic dataset, the average number of cases reporting anaphylactic reactions did not increase in the past 16 years (validated compared with the basic dataset, 38.4% vs 6.7%). Hence, intravenous administration may entail a higher risk for anaphylactic reactions compared with the basic dataset (38.4% vs 6.7%). This finding is reassuring, but others have reported similar findings.12
### TABLE 4  Distribution of designated allergy/atopy and reported symptoms according to suspected underlying pathophysiology

| Suspected pathophysiology according to literature<sup>a</sup>,<sup>35,38-43</sup> | Validated Dataset<sup>b</sup> (n = 159) | Cases Attributed to Antibiotics n = 48 (30.2%) | Cases Attributed to Iron n = 7 (4.4%) | Cases Attributed to Analgesics/Antipyretics n = 35 (22.0%) | Cases Attributed to Atracurium n = 5 (3.1%) | Cases Attributed to MRI Contrast Median = 19 (11.9%) | Cases Attributed to Alglucosidase (enzymes)<sup>c</sup> n = 12 (7.5%) |
|---|---|---|---|---|---|---|---|
| Immune mediated<sup>d</sup> | Non-immune mediated | Non-immune mediated<sup>e</sup> | Immune or non-immune mediated | Immune or non-immune mediated | Immune (IgE) or non-immune mediated |
| **Allergy/atopy**<sup>f</sup> | 25.2% (40/159) | 14.6%* (7/48) | 14.3% (1/7) | 42.9%* (15/35) | 20.0% (1/5) | 31.6% (6/19) | 0% |
| 35.8% dyspnea (57/159) | 50.0%* dyspnea (24/48) | 42.9% dyspnea (3/7) | 40.0% urticaria (14/35) | 60.0%* anaphylactic shock (3/5) | 42.1% dyspnea (8/19) | 58.3%* rash (7/12) |
| 33.3% urticaria (53/159) | 31.3% urticaria (15/48) | 42.9% urticaria (3/7) | 31.4% anaphylactic reaction (11/35) | 40.0%* bronchospasm (2/5) | 31.6%* erythema (6/19) | 50.0% urticaria (6/12) |
| 22.0% rash (35/159) | 27.1% rash (13/48) | 31.4% urticaria (3/7) | 31.4%* angioedema (11/35) | 31.6%* cough (6/19) | 31.4%* vomiting (6/12) |

<sup>a</sup>Non-immune-mediated reactions cover different pathomechanisms, like NSAID-induced inhibition of COX enzymes,<sup>5,35,38</sup> complement activation by intravenously administered iron,<sup>39</sup> direct degranulation of mast cells in non-IgE-mediated hypersensitivity reactions induced by MRI contrast media,<sup>40,41</sup> or by neuromuscular blocking agents like atracurium.<sup>38,43</sup> In this table, the validated cases (n = 159; validated dataset) are stratified according to drug class, the reported underlying allergic/atopic conditions, the assumed underlying pathophysiological mechanisms, and the three most frequently reported symptoms.

<sup>b</sup>A total of 159 case reports contained 164 suspected drugs. Cases with more than one drug were counted in each drug class. However, they were not counted twice if they belonged to the same drug class.

<sup>c</sup>Twelve case reports for alglucosidase. Among these 12 cases, there was one patient accounting for five cases (each at a different date). In these cases, there was no evidence that the reactions occurred in context with a desensitization procedure.

<sup>d</sup>This group also contained four fluoroquinolone cases. Both immune-mediated and non-immune-mediated reactions have been described for fluoroquinolones. The first is reported as being more common.<sup>38</sup>

<sup>e</sup>Five subtypes of NSAID-induced hypersensitivity reactions have been proposed,<sup>35</sup> including non-immune-mediated and immune-mediated reactions. In one publication, it is assumed that non-immune-mediated cases account for more than 75% of cases.<sup>38</sup>

<sup>f</sup>Cases with patients designated as atopic (n = 22) or allergic (n = 29) were pooled for subgroup analysis (see section Results). Not mentioned does not exclude allergic/atopic condition.

<sup>g</sup>Reported symptoms by analyzing the assigned preferred terms. The diagnosis "anaphylactic reaction" is based on specific symptoms reported. Some symptoms may be reported more often than others. In some cases only the diagnosis "anaphylactic reaction" is reported.

<sup>*</sup>Chi-squared test/Fisher exact test; P < 0.05. Further information on the calculation of P values is included in Section 2.7.
antipyretics-induced cases. Regarding the differentiation of NSAID-induced hypersensitivity, this finding could reflect a higher proportion of the “NSAID-induced urticaria/angioedema” type or the “NSAID-exacerbated cutaneous disease” type in our cases. Children aged 0 to 5 years more often reported urticaria and vomiting than older age classes. In contrast, decreased blood pressure was more frequent in adolescents (13-17; data not shown) as also reported by others.11

About one quarter of the cases was designated as atopic/allergic; similar results were reported in other studies. Although preferential underreporting cannot be excluded, atopy was not confirmed as a risk factor for severe reactions in our study, which is also in accordance with literature.12,15,18,37

Atopic patients are IgE antibody high responders. We found a lower percentage (14.6%) of patients reporting atopy/allergy in “antibiotics cases” with assumed preferential immune-mediated pathophysiology (according to literature). On the other hand, in the “analgesics/antipyretics cases” with assumed preferential non-immune-mediated pathophysiology (according to literature), a higher percentage (42.9%) was observed. No significant association with atopy for beta-lactam allergy in children was found in other studies either. Instead, varying associations of atopy with different phenotypes of NSAID-induced hypersensitivity have been described, suggesting that atopy may predispose to selected forms of NSAID hypersensitivity. However, in one study in patients of all ages, no differences were found. Therefore, our findings could also be due to chance or varying documentation.

Ibuprofen accounted for nearly every fifth of all incriminated drugs (18.9%; 30/164) and nearly every fourth in the age groups 0 to 5 and 6 to 12 years (data not shown). No matching exposure data are available. However, ibuprofen passed paracetamol in terms of exposure in 2007 and accounted for 76% of all analgesics prescribed to children within the statutory insurance system in Germany in 2013. Over-the-counter sales may further increase this exposure. Nevertheless, if the large number of reports is seen in context with the large exposure, we arrive at a more reassuring scenario.

Cefaclor accounted for 27.1% (13/48) of cases attributed to antibiotics, and nearly every second (46.2%; 6/13) was designated as life-threatening. Cefaclor accounted for 10.4% of all antibiotics prescribed to children (0-15 years) in Germany in 2004 and for 18.6% in 2013. In contrast, amoxicillin accounted for only four reports (none designated as life-threatening), although it was the most frequently prescribed antibiotic for children in Germany in 2013 (28.7% of all antibiotics); this ratio has remained relatively stable since 2004. However, because of the limitations of the spontaneous reporting system, we cannot determine whether this finding reflects drug-preferential reporting, different potentials of these drugs to induce anaphylactic reactions, or other reasons.

All five atracurium cases were designated as serious (one fatal). It remains unclear whether atracurium is associated with more severe anaphylactic reactions or whether severe anaphylactic reactions occurring under anesthesia are more likely to be noticed/reported. The latter would also apply to other drugs used in anesthesia, which were not seen in our analysis. Nevertheless, our finding could also reflect different exposure rates. An analysis in France also reported a higher ratio of grade III/IV hypersensitivity reactions for neuromuscular blocking agents.

In 13.8% of the cases (11.3% if excluding reported readministration in one person), previous hypersensitivity to the drug had been reported, and these reactions appeared to be more severe than cases designated as “drug never used before.” Hence, serious anaphylactic reactions might have been avoided in about every seventh case if taking the patient’s history had included previous hypersensitivity reactions and if this factor had been considered prior to treatment. Concerning the 22/54 (40.7%) cases where previous administration had been tolerated, sensitization could have occurred in the immune-mediated cases. Finally, we cannot rule out that there may have been cases for which no alternative medication was available.

The strengths of the spontaneous reporting system encompass the large number of potential cases, the inclusion of vulnerable patient populations (eg, children), and the possibility to detect very rare/long latency ADRs. Its limitations include underreporting, preferential and stimulated reporting, a varying degree of documentation in the reports, and the impossibility to calculate ADR frequencies due to lack of exposure data. Hence, epidemiological studies not based on spontaneous data are usually required to further investigate the signals observed.

In conclusion, a heterogeneous clinical phenotype with differences in associated factors was observed, suggesting different underlying mechanisms triggered by the different drug groups. Future studies may thus focus on defined drug groups. Exploration of larger databases like EudraVigilance could be helpful in order to gain access to further of such cases.

### Ethics Statement

The study has been approved by the local ethics committee (009/17).

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### Table 5

| Ranking | Drug Substance | Drug Class |
|---------|----------------|------------|
| 1.      | Ibuprofen (n = 30) | Analgesics |
| 2.      | Cefaclor (n = 13) | Antibiotics |
| 3.      | Alglucosidase (n = 12) | Alglucosidase |
| 4.      | Gadobutrol (n = 9) | MRI |
| 5.      | Azithromycin (n = 5) | Antibiotics |
| 5.      | Cefuroxime (n = 5) | Antibiotics |
| 5.      | Etoposide (n = 5) | Other |
| 5.      | Atracurium (n = 5) | Atracurium |
| 5.      | Gadopentetate (n = 5) | MRI |
| 5.      | Gadoteric acid (n = 5) | MRI |

A total of 159 cases with 164 incriminated drugs.
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