

The Emperor Has No Symptoms: The Risks of a Blanket Approach to Using Epinephrine Autoinjectors for All Allergic Reactions



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Fatal anaphylaxis in humans is rare and unpredictable. We note a trend to provide allergic individuals with care plans that recommend immediate use of epinephrine autoinjectors if allergen ingestion is suspected, even in the absence of any allergic symptoms, without any supporting evidence base. Instructions to use an autoinjector device, irrespective of reaction severity and especially when symptoms are *actually* absent, are likely to add to parental and patient anxiety. Of greater concern is the possibility of epinephrine being administered “too early” to treat initial, mild symptoms that then progress to severe anaphylaxis. It is not hard to visualize a scenario where one or both epinephrine autoinjectors have been deployed for mild

symptoms, yet the reaction progresses to a severe reaction and no further epinephrine is available for administration. Epinephrine needs to be available as a rescue treatment for anaphylaxis, potentially buying valuable minutes while emergency medical services are activated to attend. Food-allergic individuals and their carers need to be provided with more constructive strategies and support than merely being told to “use your pen.” © 2016 The Authors. Published by Elsevier Inc. on behalf of the American Academy of Allergy, Asthma & Immunology. This is an open access article under the CC BY license (<http://creativecommons.org/licenses/by/4.0/>). (J Allergy Clin Immunol Pract 2016;4:1143-6)

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A new emergency care plan for food-allergic individuals was published by the Food Allergy Research & Education (FARE) earlier this year. We find some elements in the plan¹ controversial, and are unaware of the evidence base on which some of the directives (Figure 1) have been promulgated, particularly the recommendation to administer epinephrine even in the absence of symptoms.

It is very difficult to determine whether an allergic individual is “extremely reactive” or not. The term “reactivity” causes confusion: does it refer to dose sensitivity (or threshold) to allergen, or severity of the elicited reaction? Most individuals who react to food allergens at the bottom of the dose response curve (derived from food challenge data) experience only mild reactions.^{2,3} Furthermore, data suggest that individuals with a history of anaphylaxis are not more dose sensitive than those without a history of anaphylaxis.⁴⁻⁷ The vast majority of fatal reactions are due to substantial levels of exposure accidentally ingested, often with other significant cofactors present contributing to the severity of the reaction.⁸⁻¹⁰

Anaphylaxis to food is not uncommon¹¹; however, fatal food-induced anaphylaxis is very rare, with a case fatality rate at <0.0001%.^{12,13} Data suggest that, in many cases, anaphylaxis (admittedly at the less severe end of the anaphylaxis spectrum) resolves without any treatment¹⁴—a daily event in many emergency departments (ED), where patients present having experienced symptoms of anaphylaxis but failed to use their autoinjector or have not had their anaphylaxis correctly diagnosed in the ED. Arguably, our biggest challenge in the management of food-allergic individuals is in identifying those who are most at risk of severe, life-threatening reactions; although fatal anaphylaxis is rare, it is also very unpredictable.¹¹ As a result,

Abbreviations used

ED- Emergency departments

EMS- Emergency medical services

we have to apply the same management strategies to all food-allergic individuals, resulting in the widespread provision of rescue medication including epinephrine autoinjectors to anyone considered to be at risk of anaphylaxis. That in itself is a further challenge; the limited published data imply that up to 3 quarters of peanut-allergic children will have anaphylaxis if exposed to *sufficient* allergen.¹⁵

CAN USING EPINEPHRINE AUTOINJECTORS FOR MILD ALLERGIC SYMPTOMS CAUSE HARM?

We believe that the use of epinephrine for *any* exposure to an allergen is overtreatment, and may, paradoxically, place some individuals at greater risk of severe outcomes. The concept of “very early” epinephrine, given immediately at the onset of any symptom seems to be overinvasive medical advice, which has implications for negative patient perception of the burden in managing daily life with this condition. One can argue that given the difficulty for patients in identifying their own symptoms of anaphylaxis,^{16,17} it may simplify management by advising epinephrine to be administered for *all* reactions. However, given the tendency of patients, and their parents, *not* to use epinephrine autoinjectors even when appropriate to do so¹⁴—often out of anxiety for the consequences of an injection and/or underlying needle phobia—we would be concerned that a lowering of the threshold to give epinephrine would only increase the reluctance of individuals to administer an injection, even when it is warranted. Similar to the widespread use of precautionary allergen labelling on food packaging, blanket strategies often give rise to greater, rather than less risky behaviors.

We are concerned that a recommendation to use epinephrine autoinjectors for *any* symptoms of an allergic reaction might be driven, in part, by commercial interests. The patient literature provided by at least one manufacturer of epinephrine autoinjectors appears to imply that the device should be used for “critical symptoms,” which according to the instructions include mouth swelling and skin symptoms.¹⁸ This trend or “slippage”—where nonsevere symptoms are labeled “critical”—is concerning; guidelines and patient literature should be written by non-conflicted individuals with appropriate expertise, and not driven by a marketing agenda. From a patient perspective, this may further compound confusion over what constitutes “anaphylaxis,” giving rise to stress and anxiety, particularly for parents of young children.^{19,20}

Delayed use of epinephrine in evolving anaphylaxis has been associated with adverse outcomes including death.^{8,9}

Epinephrine is generally well tolerated by most individuals, even children, who receive it intramuscularly,^{21,22} and, of course, we do encourage our patients to use their epinephrine autoinjector if they have signs of airway/respiratory or cardiovascular involvement, or if they are uncertain what to do, because of difficulty in recognizing or identifying symptoms in themselves or in their child. We also advocate epinephrine use in the context of a rapidly progressing reaction. However, we do *not* advise patients to inject epinephrine for urticaria or angioedema, because, if they occur in isolation or rather in the absence of other symptoms or signs, they are nearly always minor self-limiting symptoms that, in our view, do not justify epinephrine injection. One also has to consider that tremor and/or shaking is not uncommon after epinephrine autoinjector use²²; this can be justified in the context of treating anaphylaxis, but if used for mild reactions, the allergic individual may develop a reluctance to use epinephrine that impacts on the decision to treat subsequent reactions—including anaphylaxis—with appropriate rescue medication.

Food-allergic individuals often receive differing advice on their level of reactivity from clinicians, who themselves have differing levels of expertise and experience of food allergy and anaphylaxis, which, incidentally, are not the same thing.²³ A food-allergic individual’s perception of his and/or her own risk—and likelihood of reaction—may differ from that of an experienced clinician. Parents are understandably motivated to protect their own child to the greatest possible extent in their role as parents, but the allergy literature is now full of reports of disproportionate parental anxiety impacting adversely on family life, interpersonal relationships, and general social functioning.^{19,20,24} Instructions to use an autoinjector device, irrespective of severity and especially when symptoms are *actually* absent, are likely to add to parental and patient anxiety; this is maladaptive and counterproductive, in that it results in lower self-efficacy and competence in emergency care,^{25,26} in turn leading to a greater adverse impact on food allergy-related quality of life.

The idea that consumption of any amount of allergen (eg, peanut) should motivate immediate administration of epinephrine, even when no symptoms have been elicited, is very controversial. Up to 50% of challenge-proven peanut-allergic individuals do not react to doses up to 100 mg of peanut protein (about half a peanut),³⁻⁵ although we acknowledge that the formal challenge scenario is an artificial setting and that community reactions may involve other covariables. In the panic caused by possible exposure, it can be difficult to discern whether true exposure has occurred. If a peanut-allergic individual has inadvertently eaten a candy bar and only then notices a “may contain” warning, should that individual be instructed to self-administer epinephrine? Almost two thirds of prepacked confectionery are labeled with some form of precautionary allergen label.²⁷ The majority of nut-allergic individuals will eat some of

Extremely reactive to the following foods: _____

THEREFORE:

☐ If checked, give epinephrine immediately for ANY symptoms if the allergen was likely eaten.

☐ If checked, give epinephrine immediately if the allergen was definitely eaten, even if no symptoms are noted.

FIGURE 1. Extract from the Food Allergy Research & Education (FARE) emergency self-care plan.¹

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