**Can dietary strategies in early life prevent childhood food allergy? A report from two iFAAM workshops**

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**Abstract**

Food allergy affects a small but significant number of children and adults. Food allergy is responsible for considerable morbidity and is the commonest cause of anaphylaxis in children. One of the aims of the European Union funded “Integrated Approaches to Food Allergen and Allergy Risk Management” (iFAAM) project was to improve our understanding of the best way to prevent the development of food allergy. Groups within the project worked on integrating the current prevention evidence base as well as generating new data to move our understanding forward. This paper from the iFAAM project is a unique addition to the literature on this topic as it not only outlines the recently published randomised controlled trials (as have previous reviews) but it also summarises two iFAAM-associated project workshops. These workshops focused on how we may be able to use dietary strategies in early life to prevent the development of food allergy and summarises the range of opinions amongst experts in this controversial area.

**Introduction**

Food allergy represents a considerable burden to affected individuals. Around 5% of preschool children and between 0.3-5.6% of older children and adults in Europe have food allergy (1-5). Typical causative allergens are hen’s egg, cow’s milk, peanuts, tree nuts, wheat, soya, shell fish and fish (1,2). Food allergy is the commonest cause of anaphylaxis (6), a systematic and potentially life-threatening systemic allergic reaction (7). Current management strategies for food allergy revolve around avoidance of the causative allergen and rescue therapy for reactions (7). Avoidance of foods that are often a ubiquitous part of our diet is challenging, resulting in inconvenience to the individual and their family. Inconvenience also results from the need to carry rescue therapy, such as adrenaline auto-injectors. Although with adequate avoidance strategies, allergic reactions are infrequent, food allergy can result in considerable anxiety mainly due to their unpredictability (8).Therefore food allergy represents a considerable burden to individuals, families and society.

In the past, the prevailing hypothesis to explain the aetiology of food allergy was that early ingestion of allergenic foods would result in the development of food allergy in at risk individuals. The widely accepted approach to reducing the likelihood of an infant developing food allergy was therefore to delay the introduction of dietary allergens into their diet (9-11). The failure of these avoidance strategies to reduce the incidence of food allergy has led to a reassessment and the formulation of the dual exposure hypothesis (12). This reassessment developed from a number of different lines of evidence and suggests that initial contact of an allergenic food via the gastrointestinal tract in early life leads to development of tolerance; this contrasts with when initial contact is via eczematous skin when allergic hypersensitivity is likely to develop. This new hypothesis has led to a number of randomised controlled trials to assess whether the dietary introduction of allergens at 4-6 months leads to a reduction in the development of clinical food allergy. A number of these trials have now reported.

Integrated Approaches to Food Allergen and Allergy Risk Management (iFAAM) was an European Union funded FP7 project. iFAAM aimed to reduce the burden of food allergy through a number of integrated activities. The project evaluated the evidence relating to the development of food allergy to improve our approach to the prevention of food allergy. This paper is a summary from two iFAAM workshops held on 12th December 2016 and 19th to 20th April 2018. The aims of these workshops were to review the recent key data on the prevention of food allergy; to make recommendations on the best approach to utilise infant diet to prevent food allergy; and consider what gaps remain in the evidence base. The discussion was focused on the introduction of complementary solid foods into the infant diet with “early” defined as less than 26 weeks of life. Participants were all involved in iFAAM and represented the breadth of stakeholders and range of opinions in this area for example paediatricians, dietitians, patient group representatives, and food scientists.

**Potential impact of altering the age of introduction of allergenic solids on infant nutrition**

In making recommendations about infant diet, a holistic view is required, as positive impacts in one clinical area may need to be balanced with negative ones in another. This can be seen by the effect of feeding recommendations on the overall composition of the infant diet. Historically infants were breast fed for their first year of life with solids introduced once they were able to physically eat, e.g. being able to hold head up independently. Initiation rates and duration of breast feeding started to decline in many European Countries in the 1950s (Figure 1). At that time, complementary foods could be started from as early as 6-8 weeks of age. From the 1970s there were recommendations to introduce complementary foods later (13,14). In 2003, the World Health Organisation recommended exclusive breastfeeding until 26 weeks of age with continued breastfeeding until two years of age (15). In practice, this is rarely achieved in many countries in the developed world (8). Where breastfeeding is not prolonged, delaying the introduction of complementary foods means that there is less or even no overlap between breast feeding and the introduction of solids and this may have a detrimental effect on allergy development (11). Additionally, guidelines may be directed at particular population groups (e.g. high risk infants) but are followed by all population groups (9,10).

The format in which a food is eaten is very important for infants and is affected by their age. Consequently this needs to be considered when deciding how to introduce allergenic foods into the diet. For example, egg is traditionally introduced into an infant’s diet in a well-cooked format to overcome texture issues and is consequently given at a relatively low dose. This observation is supported by available trial data which demonstrates that cooked egg is much better tolerated than raw egg and is consumed in lower doses (16). Likewise, the choking risk with peanuts also needs to be considered, children under 5 years of age should only consume these in a ground/smooth format such as peanut butter or satay sauce. Consideration is also needed to ensure that any recommended changes to the infant diet are in line with local dietary practice. For example, peanuts are not part of the local diet in many areas of the world. Finally, the impact of any recommendations requires reflection to ensure it has no negative nutritional or dietary consequences. For example, peanuts formulated as peanut puffs have a relatively high sodium chloride and fat content and nutritional data from the LEAP study showed that the early introduction did have a major impact on the relative intake of key nutrients, although no adverse effects on growth were found (17). Peanut would be better introduced as part of the protein content of meals rather than as a snack to reduce the potential for an infant having too much protein, protein-associated fat and sodium chloride in their diet which have long-term health consequences including obesity in later life (18). Finally, there is a need to look for potential long-term consequences as our dietary experience as infants affects our long-term diet and therefore health (19).

All these factors demonstrate why it is important that one clinical speciality should not drive public nutrition recommendations. The European Society of Paediatric Gastroenterology and Nutrition have published updated recommendations on complementary feeding (20). They recommended that “exclusive or full breast-feeding should be promoted for at least 4 months and exclusive or predominant breast-feeding for approximately 6 months is a desirable goal.” Additionally they recommend that “complementary foods should not be introduced before 4 months but should not be delayed beyond 6 months.” They explain that they base these recommendations on when an infant has the necessary maturity in gastrointestinal, renal and motor function to safely consume complementary foods.

**Figure 1. Illustration of the historic trends in breasting feeding and the introduction of allergenic solids into the infant diet in the UK**

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Figure illustrates the typical timelines for breast feeding and the introduction of solids into the infant diet in the last 4 decades in the UK where the LEAP and EAT were undertaken (13,14, 21-23). However, an overlap between breastfeeding and the introduction of solids can still be seen in other European countries such as Norway and Denmark (24).

**Randomised controlled prevention trials focused on early introduction of allergenic solids**

A literature search was undertaken in Medline and EMBASE ([food allergy or hypersensitivity or intolerance] and [prevention] and [SIGN randomised controlled trial search strategy (25)]). Additionally, Clinicaltrials.gov, [www.clinicaltrialsregister.eu](http://www.clinicaltrialsregister.eu), [www.controlled-trials.com](http://www.controlled-trials.com), and <http://www.anzctr.org.au> were searched for ongoing or planned randomised controlled trials (18th April 2018) and re-run for any new studies 7th December 2019. A total of 7 randomised studies focusing on the question of whether the early introduction of allergenic solids might prevent the development of food allergy have been published; a further one is ongoing (Table 1). Most of the studies focus on egg (16,26-29), one has focused on peanuts (30,31) and one on multiple foods (32). One trial of multiple food allergies (PreventADALL) is still ongoing (33). Lastly, there is one non randomised trial on peanut allergy in high risk infants (PEAP) that is also ongoing (34). To understand these studies fully, a number of different aspects need to be understood:

*Population* – studies have focused on either a high risk group (16,26,30,31), moderate risk (27,28) or a general population (29,32,33. High risk has been defined as the presence of moderate to severe eczema, which is known to be associated with food allergy (35). Moderate risk has been defined as a first degree relative with allergy (27,28). Some studies have aimed to recruit a general population although inevitably, those enrolled are likely to be more at risk than a truly representative general population (29,32,33). Studies based in an at risk population have a better chance of identifying any preventative impact of an intervention as more individuals develop food allergy than in the general population. There is though a possibility that such high risk infants have already developed food allergy. The successful impact of the LEAP intervention in sensitised infants (SPT < 4mm) would argue against this at least for peanut allergy. In the STAR study (26), which did not screen for sensitisation prior to inclusion in the study, many infants reacted to the egg intervention and these infants had significantly higher SpIgE to egg at recruitment than those who did not react. However in the HEAP (study (29) only infants who were not sensitised were included and there were still frequent reactions to the egg intervention.; this was not seen in the PETIT study (16) who did not screen by Egg specific IgE prior to inclusion. The likely explanation for this difference was the use of small doses of heated hen’s egg in the PETIT study.

*Intervention* – foods have been introduced in a number of formats and doses. For the egg interventions, a number of studies used a format that appears to be similar to raw egg in terms of allergenicity (16,27-29). But the EAT and PETIT studies used a cooked form of egg (16,32). Interestingly, it appears introduction of the “raw” egg into the infant diet does not prevent the development of egg allergy and results in lots of allergic side effects. Meanwhile, introduction of the cooked form into the infant diet possibly results in a reduction in the development of egg allergy and has been well tolerated. One challenge here though is that it is difficult to standardise the degree of cooking. Other factors may also be important, for example the matrix (Table 2). Dose wise, studies have used a wide range of amount of food. Most of the egg studies have used an amount equivalent to approximately ½-1 egg per week (26-29,32). The exception is the PETIT study, which started with a low daily egg protein intake (equivalent to 1/20th of an egg a week) from 6- 9 months, followed by daily intake equivalent to a ¼ of an egg a week from 10-12 months and was associated with a very large reduction in egg allergy (16). It is important to note that the hen’s egg in the PETIT and EAT study was well cooked. For the peanut studies, the LEAP study gave 6g peanut protein per week (30,31) whereas the EAT study used 4g of peanut protein (32).

*Timing of introduction of allergen solid(s)* – the egg studies have all introduced this allergen at 4-6 months of age (22-26) while the peanut study introduced the allergen from 4-11 months (mean 7.8 months) (27,28). In the EAT study, allergenic foods were introduced between 4-6 months except for cow’s milk which was introduced from 3 months (29).

*Comparator* – only some of the studies had a double-blind design (26-29); the other randomised controlled trials were open in design (26-28). Without blinding there is a potential for reporting bias; to overcome this, the primary outcome needs to be objective and robust. The success of the randomisation also needs to be examined, for example, there was an imbalance in baseline sensitisation to egg in the PETIT study although a *post hoc* analysis did suggest that this did not affect the result (16).

*Outcomes* – the patient relevant outcome is food allergy which is assessed in most studies (26-32); only some used a double-blinded challenge protocol (25,26-28). Challenge results can be difficult to define accurately in small infants. Several studies have used sensitisation as their primary outcome (28,29); given the data that sensitised infants can be protected from developing food allergy (30,31), it is difficult to know how to interpret these sensitisation results in isolation from accompanying challenge results.

**Table 1. Summary of randomised controlled studies investigating the impact of early introduction of allergenic foods on prevention of food allergy**

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| **Name of Trial** | **Country (institution)** | **Population** | **Study details** | **Results** |
| **Egg** |  |  |  |  |
| Hens’ Egg Allergy Prevention (HEAP)Bellach et al(29) | Germany (Charité Universitätsmedizin Berlin, Germany) | General population, non-sensitised (Screened at recruitment) | * RCT, placebo controlled
* n = 383
* Enrolled at 4-6 months then consumption of egg powder (“pasteurized egg white” equal in its allergenicity to raw hen’s egg) or placebo until 12 months of age; started with 800mg egg protein three times a week, increasing to 1.6g in week 2 and 2.5g in week 3.
* Outcome: primary: prevalence of egg sensitisation; secondary: placebo controlled challenge proven IgE-mediated egg allergy at 12 months of age
 | At 12 months there was a non-significant difference in egg sensitisation (2.6% control vs 5.6% egg, p=0.24) and egg allergy (0.6% control vs 2.1% egg, p=0.35) between the groups in ITT analyses; many infants reacted to the intervention. |
| Prevention of egg allergy ininfants with atopicdermatitis (PETIT)Natsume et al(16) | Japan (National Centre for Child Health and Development, Japan) | High risk (infants with atopic dermatitis) | * RCT, placebo controlled
* N=147
* Enrolled at 4–6 months then consumption of “heated egg powder” or placebo; started on a very small amount (25 mg of egg protein daily increasing to 125mg from 9 months)
* Outcome: prevalence of open challenge proven IgE mediated egg allergy at 12 months of age
 | Recruited finished early after an interim analysis; intervention lead to a significant reduction in egg allergy (38% control vs 8% egg, p=0·0001); no major safety issues.  |
| Solids Timing for Allergy Research (STAR)Palmer et al 2013(26) | Australia (University of Western Australia) | High-risk (infants with moderate / severe eczema) | * RCT, placebo controlled
* n = 86
* Enrolled at 4-6 months of age then consumption of egg powder (“pasteurized raw whole egg powder”) or placebo until 8 months of age; 900mg egg proteins given daily
* Outcome: prevalence of open challenge proven IgE-mediated egg allergy at 12 months of age
 | A third of infants randomized to egg reacted to the intervention; at 12 months, there was a non-significant difference in egg allergy between the groups in the intervention group (51% control vs 33% egg, ITT analysis, p=0.11). |
| Starting Time for Egg Protein (STEP)Palmer et al 2016(27) | Australia (University of Western Australia) | Moderate-risk (infants without eczema but atopic mothers) | * RCT, placebo controlled
* N=804
* Enrolled at 4-6 months of age then consumption of pasteurized raw whole egg powder or placebo until 12 months of age
* Outcome: prevalence of challenge proven IgE-mediated egg allergy at 12 months of age
 | At 12 months there was a non-significant increase in IgE-mediated egg allergy in the control group compared to the intervention group (10.3% control vs 7.0% egg p=0.20); there were a number of allergic reactions to the intervention. |
| Beating Egg Allergy (BEAT) Tan et al(28) | Australia (Sydney University Children’s Hospital) | Moderate-risk (sibling / parent with allergy). SPT to egg <2mm | * RCT, placebo controlled
* N=319
* Enrolled at 4 months of age then consumption of pasteurized whole egg powder, or placebo until 8 months of age
* Outcome: primary egg white sensitisation; secondary: prevalence of IgE mediated egg allergy at 12 months of age
 | Early introduction of egg was associated with a reduction in sensitization at 12 months (20% control vs 11% egg, p=0.03). There was no effect on the proportion of children with probable egg allergy. A number of infants reacted to the intervention. |
| **Peanut** |  |  |  |  |
| Learning Early About Peanut allergy (LEAP) Du Toit et al(30,31) | UK (Kings College, London) | High-risk (infants with moderate / severe eczema and / or egg allergy). SPT to peanut < 4mm | * Open-label RCT
* n = 640
* Enrolled at 4-11 months (mean age 7.8 months) then peanut consumption or avoidance until age 5; 6g peanut protein per week
* Outcome: prevalence of DBPCFC confirmed peanut allergy at 5 years of age
 | Significant reduction at 60 months in ITT analysis (13.7%control vs 1.9% overall, p<0.001) regardless of presence of initial (cutaneous) sensitisation; no significant between-group differences in serious adverse events. Significant reduction still seen after both groups then avoided peanut for a year (18.6% control vs 4.8% peanut, ITT analysis, p<0.001). |
| **Multiple foods** |  |  |  |  |
| Enquiring About Tolerance (EAT)Perkin et al(32) | UK (Kings College, London) | General population | * Cows’ milk, hens’ egg, peanut, cod, sesame, wheat
* Open-label RCT
* n = 1106
* Enrolled at 3 months of age then consumption of 6 allergenic foods until 6 months or exclusive breastfeeding until 6 months of age; 2g protein of each food given twice a week
* Outcome: prevalence of IgE-mediated food allergy to any of the 6 allergenic foods between 1 and 3 years of age
 | Non-significant reduction in ITT analysis (7.1%control vs 5.6% intervention, p=0.32). In PP analysis, a significant reduction was seen for any food allergy (7.3% control vs 2.4% intervention, p=0.01), peanut (2.5% control vs 0% peanut, p=0.003) and egg allergy (5.5% control vs 1.4% egg, p=0.009)  |
| Preventing atopic dermatitis and allergies in children (PreventADALL)(33) | Norway (Oslo University Hospital) | General population | * Hen’s egg, milk, wheat, peanut
* Open label RCT with four arms: observation, early introduction by 4 months, skin care, both early introduction and skin care
* N=2500
* Outcome: food allergy, atopic dermatitis
 | Ongoing |

DBPCFC, double blind, placebo controlled food challenge; ITT, intention to treat; PP, per protocol; RCT, randomised controlled trial.

**Synthesis of the randomised control prevention trials**

A systematic review and meta-analysis of these randomised controlled prevention trials has been published (36). Level of risk for developing food allergy differed between these populations and can be seen in Table 1. An updated figure from the review is reproduced as Figure 2. The analyses suggest that the early introduction of peanut and egg are effective in preventing the development of peanut and egg allergy respectively. The PETIT study is a cause of much of the heterogeneity (16), and it has been criticised given that it was stopped early, after an interim analysis, when the two groups were unbalanced for the presence of specific IgE to egg (36). However, meta-analysis of the egg studies remains positive for a preventative effect even if the PETIT study is removed from the analysis. These updated data give a number needed to treat to prevent egg and peanut allergy of 28 and 18 respectively. The analysis for milk allergy failed to demonstrate that early introduction is an effective preventative strategy.

**Figure 2. Meta-analysis of randomised controlled prevention studies**

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Updated from Ierodiakonou et al (36) where results for egg, milk and peanut allergy are presented separately. Error bars represent 95% confidence intervals.

A subsequent systematic review and meta-analysis of studies looking at early introduction of egg and allergy development indicated that neither the total amount nor pre-treatment of egg showed any effect on egg allergy development at the age of 12 months. The authors propose further studies should be performed to generate stronger data (35).

**Interpretation of the randomised controlled prevention trial data**

For egg, milk and peanut, there are important limitations in the currently available data. There is heterogeneity in both the egg and peanut analyses. For egg, this is largely explained by the imbalance in risk factors for egg allergy at baseline in the PETIT study, the early termination of recruitment in this trial and the usage of very different amounts of the allergen (16). For peanut the heterogeneity may be explained by very different adherence to the intervention seen in the LEAP and EAT studies and also the different population groups (Table 1) (30-32). The data for peanut allergy is based on two studies from one institution and it remains to be seen whether the findings are generalizable to other countries with different breastfeeding initiation rates and durations, different prevalences of eczema and different household peanut consumption. Peanut allergy is also not seen in many countries. We do not have data for other allergenic foods. Lastly, we need to consider the potential for adverse consequences of the early introduction of allergenic solids; this is discussed in the next section.

So while there appears to be a proof of concept that early introduction of allergenic solids in the right amounts, in the right formulation and at the right time can reduce the development of food allergy in infants, important gaps in the evidence base remain (Table 2). The heterogeneity of the published studies is a particular concern with differences in the included populations, format and dose of the intervention and how “early” was defined. This heterogeneity could be used to optimise the approach but a definitive study would then be required to assess this strategy.

**Can we extrapolate from the available evidence to other food groups?**

Allergic reactions to different foods present very differently. Although their presentation is usually temporally associated with when that food group first appears in the diet, it is possible that clinically relevant allergy develops sometime before the first known ingestion. This may be different for different foods. For example, trials have demonstrated that peanut interventions are well tolerated even by sensitised infants up to 11 months of age (30-32) whereas many younger infants have allergic reactions with the raw pasteurised egg intervention (26,29). However, this difference may just as easily be due to the form the foods were ingested and how the infants were screened prior to inclusion in the study.

Additionally, there may be different “windows of opportunity” for prevention to different allergenic foods during childhood. Also, whether or not the allergy is outgrown and the speed that this happens is very variable making extrapolation to other foods problematic. However, the mechanisms that underlie tolerance are likely to be similar for all foods and will involve either clonal deletion, clonal anergy, or suppression. At present we can only compare early introduction of different foods in the EAT study (32) and this was underpowered to look individually at the less prevalent food allergies. We need more evidence for all the food groups (Table 2).

**Is there adequate evidence to make recommendations about the best approach for solid introduction and allergy prevention?**

Participants at the first workshop (2016) were asked to consider, in four small groups, whether there was sufficient current evidence to make recommendations about the age of introduction of complementary solids into the infant diet to reduce the likelihood of developing food allergy. In the second workshop, groups were asked to consider i) the definition of a high risk infant; ii) when foods should be introduced to low risk infants; iii) when foods should be introduced to high risk infants; iv) what are the potential consequences of early introduction of allergenic foods; v) should different foods be introduced at different ages and vii) in what format should foods be introduced.

There was agreement by participants at the 2018 workshop that there is a paucity of evidence to be able to define absolutely high(er) risk populations for the development of food allergy. However, it was thought that whilst a family history of allergies is widely acknowledged to increase the risk of food allergy development, there are other factors such as early onset eczema, ethnicity and geography that also have an effect and need to be considered.

All the participants at the first workshop (2016) agreed that breastfeeding should be recommended until at least 6 months of age, not based on its impact on food allergy but due to its other proven health benefits. The most recent systematic review on this topic compared maternal and infant outcomes with exclusive breastfeeding for 6 months versus 3 to 4 months (38). A total of 23 eligible studies (11 from low- and 12 from high-income countries) were included. Some benefits for exclusive breastfeeding for 6 months were seen, for example reduced gastrointestinal infection even in high-income countries. A weakness of the evidence base is that most of the studies are observational with no randomised controlled trials in high-income countries. Moreover exclusive breastfeeding duration in these studies was often determined by the introduction of breastmilk substitutes rather than introduction of solids, and it is unclear whether early introduction of solid foods on its own increases risk for gastrointestinal infection. At the second workshop (2018) the question of breastfeeding duration for allergy protection was not discussed directly but continuing to breastfeed whilst introducing solids into the diet was recommended by the many participants when answering the other questions (detailed above).

In 2016, all the discussion groups discussed when the allergenic foods should be introduced into the infant diet. For egg, most thought that it could be introduced into the diet of all infants from 4-6 months of age, when they are developmentally ready. Given the trial evidence, it was agreed that this should probably be in small amounts of the well-cooked or baked form that is appropriate within the local diet. However one group believed that the heterogeneity in the design of the completed trials and the relatively high number of adverse reactions meant that firm conclusions could not be made. Pragmatically this results in a timeframe around 4-6 months of age. The lack of consensus between the workshop participants indicated the need for additional data focused on the format and dosage of egg used in the intervention (Table 2). Since the first workshop there have been two international consensus documents have made recommendations for egg. The Asia Pacific Association of Pediatric Allergy, Respirology & Immunology (APAPARI) consensus statement says that high risk infants with severe eczema should first be assessed by an allergy specialist if parents want to introduce egg from 5-6 months of age (39). Meanwhile the Australian Society of Clinical Immunology and Allergy (ASCIA) states “all infants should be given allergenic solid foods including peanut butter, cooked egg, dairy and wheat products in the first year of life” (40). The participants at the second workshop (2018) reached a consensus that egg should be introduced first in the heated or cooked form, after the establishment of weaning by the introduction of low allergenic foods, e.g. fruit and vegetables between 4-6 months of age.

For peanut there was no consensus in either of the discussion groups. One group thought that the peanuts should be introduced into the diet of infants in the general population from around 4 months based on the evidence of benefit from the LEAP study (30.31) and lack of evidence of harm in the EAT study (32). Another group believed that data from the LEAP and EAT studies (30-32) was sufficient to recommend introducing peanuts into the diet of infants with severe eczema and/or hens egg allergy from the age of 4-11 months after testing for sensitisation to peanut in the UK. They restricted their recommendation because of the selected population recruited into LEAP (30,31), the lack of replication in the intention to treat analysis in the EAT study (32) and lack of evidence from outside of the UK. This is similar to the stance taken by an international consensus group (41), ESPGHAN (20), the Australian society (40), BSACI (42,43) and Asian association (where peanut prevalence is high) (39) although these extend their recommendations to all countries with a high prevalence of peanut allergy. The more recent US National Institute of Allergy and Infectious Diseases extend the recommendations of peanut introduction of around 6 months to infants at moderate and low risk of developing peanut allergy, however, only in accordance with family preferences and cultural practice (44). Other discussion groups agreed that peanut should be introduced into the infant diet once they are developmentally ready and other solids have been introduced; they thought that this would ideally be 4-6 months of age given the potential risk of increased risk of peanut allergy if introduction is delayed. The range of views clearly identify this is still a topic where there is a substantial gap in available evidence with the perceived need for confirmatory data from inside and outside the UK and in low risk infants (Table 2). Finally the US National Institute of Allergy and Infectious Diseases has suggested that in countries where peanut products are not widely consumed by adults, early dietary introduction of peanut could lead to an increase in sensitization and allergic manifestations (44). This fact was mentioned in the discussion at the 2018 workshop when answering the question of what potential adverse consequences of early introduction of allergenic foods. Other potential adverse consequences were given as allergic reactions (including anaphylaxis) and more rarely FPIES and EoE, increased risk of sensitisation, and the adverse impact on other eating patterns and nutritional intake.

For allergenic foods, the 2018 workshop reached consensus that allergenic foods should be introduced at around the same time as other complementary foods are introduced as per country-specific recommendations and culinary and family practices. However, it was thought that there were insufficient data to make recommendations about when other allergenic foods should be introduced into the infant diet, a further gap in the evidence base (Table 2).

**Table 2. Research gaps**

|  |  |  |
| --- | --- | --- |
| Gap | Approach to closing gap | Priority |
| **General considerations** |  |  |
| Evidence that early introduction of solids is beneficial for low risk infants. | Needs very large randomised controlled trial in a low risk population and in different countries as current observational data are confounded by socioeconomic factors. | + |
| How early should allergenic solids being introduced into the infant diet? Eg is <4 better than 4 or 5 months?  | Randomised controlled trial but impact likely to be different for different foods so trials need to be powered to look at specific foods.  | + |
| Appropriate format (eg raw, cooked) and dosage of the food. | Secondary analysis of published trials to assess optimal format and dose. Further assessment of hypothesised optimal approach in a randomised controlled trial.  | ++ |
| Lack of evidence on how food processing and the food matrix may affect the prevention of food allergy.  | Further studies need to control for a potential matrix effect.  | + |
| Potential role of maternal consumption and transfer of dietary protein to infant via breast milk on the prevention of food allergy  | Future trials need to control for or investigate the potential impact of maternal consumption of allergenic foods on the prevention of food allergy. | + |
| The longevity of any preventative impact.  | Ongoing follow up of randomised controlled trials. Before-after controlled trials.  | + |
| Evidence that allergenic solid introduction with concurrent breast feeding is protective  | Need to examine in datasets of existing RCT studies, or to be examined in future studies. | ++ |
| Short term nutritional safety of early introduction of allergenic solids.  | Nutritional consequences of infant dietary interventions need to be assessed. Need to examine EAT (and other RCT) data sets | +++ |
| Longterm safety of early introduction of allergenic solids, eg hypertension, obesity and diabetes  | This requires longterm follow up of randomised controlled trial participants, potentially using routine health data. Alternatively, before-after controlled trial approach could be used. | ++ |
| **Egg** |  |  |
| Best dose and format of egg. | Positive studies use a wide variety of doses and formats of egg, need a comparative randomised controlled trial to assess the best approach.  | +++ |
| **Peanut** |  |  |
| Replication of LEAP data in other countries with a high, moderate and low prevalence of peanut allergy and different cultural practices around infant feeding.  | Multi-centre randomised controlled trial building on LEAP study protocols.  | ++ |
| **Other foods** |  |  |
| Will the LEAP approach work for tree nuts?  | Multi-centre randomised controlled trial building on LEAP study protocols. | ++ |
| Effective preventative approach for cow’s milk allergy. | Perhaps a clinical trial of early introduction of cow’s milk alongside breast feeding. | + |
| Lack of evidence about the best time to introduce other food allergens. | Further randomised controlled trials. | + |

**Summary and future perspectives**

Over the last years, we have seen some initial attempts to assess the role of early allergen introduction on food allergy development with intervention trials. Data are now available for egg with some also for peanut and milk allergy. Delaying the introduction of egg or peanut to the infant diet also appears to increase their risk for egg or peanut allergy. This may be particularly relevant for infants at high risk for developing food allergy, such as those with early onset troublesome eczema.

The randomised controlled trial data suggest that the early introduction of egg into the infant diet might be associated with the development of less egg allergy. However, questions remain given the heterogeneity in the format and dosage of egg used in different studies plus many infants reacted to the intervention with sometimes life-threatening symptoms. The majority of participants in the workshop thought that there was sufficient evidence to recommend that cooked egg should be introduced into the diet of all infants from 4-6 months of age, when they are developmentally ready, to reduce the likelihood of egg allergy developing. However, others thought that the heterogeneity in trial designs and results precluded any preventative recommendations. A further large trial using a cooked form of egg is clearly required to resolve this.

There is clear evidence of benefit from the early introduction of peanut in one study but limited replication data and no data from outside of the UK. Although National and European government and academy organisations have used these data to make wide ranging recommendations, consensus was not achieved in either of the workshops. Many participants were concerned by the lack of replication data and the generalisability of the results. Given the potential for the early introduction of peanuts into the diet as being a very successful public health preventative strategy, there is an obviously an urgent need for a replication study in a more general risk population outside of the UK.

There are obvious gaps in the evidence base for other foods although foods such as tree nuts represent important causes of food allergy. We have still much to understand about how the infant diet can be modified to reduce the likelihood of developing food allergy. However, we do have the basis to plan and conduct the much needed further trials using the best methodological approach possible. We also need to understand any potential adverse nutritional consequences of altering the infant diet and any long-term impact, both positive and potentially negative, of such strategies.

There remains the challenge as to how to advise the families of infants who are at low, medium and high risk of developing food allergies. There was a consensus at the 2016 workshop that we should be advising that infants are breast fed for at least 6 months but not for any impact on food allergy. The 2018 workshop added that cooked egg can be introduced between 4-6 months of age and peanut may be introduced between 4-11 months if culturally appropriate. There are now international guidelines (20,39-44) that recommend the early introduction of peanuts into the infant diet to prevent allergy in countries with a high prevalence of peanut allergy; but how this is achieved differs according the whether they are considered to be at high risk of developing food allergy. In alternative approach is to return to the situation where there in no deliberate delay of introducing any food to an infant but to introduce it when they are developmentally ready (rather than delaying introduction), this will usually be when they are 4-6 months of age.

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