Assessment of the results of the study by McCann et al. (2007) on the effect of some colours and sodium benzoate on children’s behaviour

Scientific Opinion of the Panel on Food Additives, Flavourings, Processing Aids and Food Contact Materials (AFC)

(Question No EFSA-Q-2007-171)

Adopted on 7 March 2008

PANEL MEMBERS


SUMMARY

Following a request from the European Commission, the Panel on Food Additives, Flavourings, Processing Aids and Food Contact Materials (AFC) was asked to assess the results of a recent study on the effect of mixtures of additives on children’s behaviour and provide an opinion on the findings, taking into account, if possible, other available scientific literature in the related area.

A recent study by McCann et al. (2007) has concluded that exposure to two mixtures of 4 synthetic colours plus a sodium benzoate preservative in the diet result in increased hyperactivity in 3-year old and 8- to 9-year old children in the general population. In an earlier study by the same research team there was some evidence for adverse behavioural effects of a

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* Two members of the Panel did not participate in the discussion on the subject referred to above because of possible conflict with declared interests.
mixture of 4 synthetic colours and sodium benzoate in 3-year old children on the Isle of Wight (Bateman et al., 2004).

In this recent study the effects of two combinations of Tartrazine (E102), Quinoline Yellow (E104), Sunset Yellow FCF (E110), Ponceau 4R (E124), Allura Red AC (E129), Carmoisine (E122) and sodium benzoate (E211) on children’s behaviour were studied. Five of the six food colours belong to the class of synthetic azo dyes and one, Quinoline Yellow (E104), is a quinophthalone. Sodium benzoate is used as a preservative.

The study involved one hundred and fifty three 3-year old and one hundred and forty four 8- to 9-year old children, selected to represent a broad range of behaviour in the general population including children with normal to high level behavioural activity. Children who were medicated for ADHD were not included. A global hyperactivity aggregate (GHA) score was the main outcome of the study, and this parameter was based on aggregated z-scores of observed behaviours and ratings by teachers, classroom observers and parents, plus, for 8- to 9-year old children, a computerised test of attention.

Mix A containing Tartrazine (E102), Ponceau 4R (E124), Sunset Yellow FCF (E110), Carmoisine (E122) and sodium benzoate significantly increased GHA scores for all 3-year old children compared to the placebo control GHA scores (effect size 0.20 [CI 0.01 to 0.39], p<0.05).

Mix B containing Sunset Yellow FCF (E110), Carmoisine (E122), Quinoline Yellow (E104), Allura Red AC (E129) and sodium benzoate had no effect on GHA scores in 3-year old children as compared to the placebo control GHA scores (effect size 0.17 [CI -0.03 to 0.36]).

This result persisted when analysis was restricted to 3-year old children who consumed more than 85% of juice and had no missing data (complete case group); in this analysis the effect of Mix A in the 3-year old children was still significantly increased compared to placebo control (effect size 0.32 [CI 0.05 to 0.60, p<0.05]) but for Mix B no significant effect on GHA scores was observed (effect size 0.21 [CI -0.06 to 0.48]).

For the 8- to 9-year old children a significant effect of Mix A (effect size 0.12 [CI 0.02 to 0.23], p<0.05) or Mix B (effect size 0.17 [CI 0.07 to 0.28], p<0.01) was seen when analysis was restricted to those children consuming at least 85% of drinks with no missing data (complete case group). When all 8- to 9-year old children that completed the study were taken into account, Mix A had no effect on the GHA scores compared to the placebo control (effect size 0.08 [CI -0.02 to 0.17]) and Mix B had a significant effect on GHA scores (effect size 0.12 [CI 0.03 to 0.22] p<0.05).

The authors concluded that exposure to synthetic colours or a sodium benzoate preservative (or both) in the diet result in increased hyperactivity in 3-year old and 8- to 9-year old children in the general population.

Based on surveys conducted from 2002 to 2005, the target colours are more frequently used in sweets but also occur commonly in soft drinks and benzoate is frequently present in soft drinks. Children consuming brightly coloured sweets may be exposed to levels comparable to those considered in the protocol of the McCann et al. study for one or more of the food colours studied. Comparable levels may also be reached in those children who consume brightly coloured soft drinks. The level of exposure to sodium benzoate is also likely to occur.

The Panel considers that the steps taken for score normalisation and aggregation are mathematical transformations that might affect the assumptions of normality and independence of the data which are essential for the whole statistical analysis. Therefore, the authors’ primary analysis was repeated using a more justifiable and conventional statistical model, and this was
supplemented by a set of additional analyses with the aim of aiding the interpretation of the results.

The Panel considers the re-analysis undertaken by EFSA, in which all single variables (minus the individual baseline value for that variable) were considered without normalisation, so that each subject served as its own reference, as the most adequate. This re-analysis was undertaken both at the level of the individual parameters as well as on the aggregated scores.

Based on the results obtained it was concluded that the analysis with the recalculated GHA score led to broadly similar conclusions to that in the original paper by McCann et al, except for the following:

1. The Mix A versus placebo comparison was not statistically significant for the 3-year olds when all subjects were included (entire sample), while the significance for the \( \geq 85\% \) consumption and complete case groups was increased slightly;

2. For the 8- to 9- year age group, the Mix A versus placebo comparison was no longer statistically significant in any of the three consumption groups.

In addition the data were analysed on the basis of a modified GHA score in which the parental scores were not included. The results from this analysis no longer revealed any statistically significant effects of Mix A or Mix B versus placebo, except for Mix B versus placebo in 8- to 9-year old completers.

A further analysis was carried out on the whole data set, comprising analysis of the single variables of parental scores, teacher scores and observer scores, and, in the case of 8- to 9-year old children, computer-based scores. There is a suggestion from these analyses that the statistically significant effects seen in the 3-year olds (Mix A versus placebo) and in the 8-to 9-year olds (Mix B versus placebo) are largely driven in the data by the parental scores and, in the older males in both comparisons, by the computer score.

The Panel notes that some, but not all, earlier studies have also reported effects of food colours on child behaviour, the majority of these studies being conducted on children described as hyperactive or with a clinical diagnosis of ADHD.

The Panel concludes that the McCann et al. study provides limited evidence that the two different mixtures of synthetic colours and sodium benzoate tested had a small and statistically significant effect on activity and attention in some children selected from the general population, although the effects were not observed for all children in all age groups and were not consistent for the two mixtures. The findings may thus be relevant for specific individuals within the population, showing sensitivity to food additives in general or to food colours in particular.

However, it is not possible to assess the overall prevalence of such sensitivity in the general population and reliable data on sensitivity to individual additives are not available.

The clinical significance of the observed effects also remains unclear, since it is not known whether these small alterations in attention and activity would interfere with schoolwork and other intellectual functioning. The clinical significance could possibly be clarified by assessments that used scales for functional impairment and diagnostic interviews, especially if a high proportion of children with high symptom scores were to be included in such a study.
There are thus a number of uncertainties that are apparent from this new research, some of which are echoed in earlier research. These include:

- the limited consistency of the results with respect to age and gender of the children, the effects of the two mixtures of additives tested and the type of observer (parent, teacher or independent observer);
- the unknown clinical relevance of the novel metric, i.e. the GHA score;
- the unknown relevance of the small effect size (as was also seen in the meta analysis of earlier studies by Schab and Trinh, (2004));
- the fact that the study has not been designed to identify the effects of individual additives;
- a lack of information on dose-response;
- the lack of a biologically plausible mechanism for induction of behavioural effects from consumption of food additives.

The Panel concludes that the McCann et al. study provides limited evidence that the two different mixtures of synthetic colours and sodium benzoate tested had a small and statistically significant effect on activity and attention in children selected from the general population excluding children medicated for ADHD, although the effects were not statistically significant for the two mixtures in both age groups.

Since mixtures and not individual additives were tested in the study by McCann et al., it is not possible to ascribe the observed effects to any of the individual compounds.

The clinical significance of the observed effects also remains unclear.

In the context of the overall weight of evidence and in view of the considerable uncertainties, such as the lack of consistency and relative weakness of the effect and the absence of information on the clinical significance of the behavioural changes observed, the Panel concludes that the findings of the study cannot be used as a basis for altering the ADI of the respective food colours or sodium benzoate.

**Key words:**
Hyperactivity, ADHD, children’s behaviour, Southampton study, McCann, food additives, food colours.
Tartrazine, FD&C Yellow No. 5, E102, CAS 1934-21-0, Trisodium-5-hydroxy-1-(sulfonatophenyl)-4-(4-sulphonatophenylazo)-H-pyrazole-3-carboxylate, food colouring substance, EINECS number 217-699-5.

Ponceau 4R, New Coccine, E124, CAS Registry Number 2611-82-7, Trisodium 2-hydroxy-1-(4-Sulphonato-1-naphthylazo)-naphtalene-6,8-disulphonate, food colouring substance, EINECS number: 220-036-2.

Carmoisine, Azorubine, CI Acid Red 14 and CI food red 3, E122, CAS 3567-69-9, Disodium 4-hydroxy-3-(4-sulphonato-1-naphthylazo)naphthalene-1-sulfonate, food colouring substance EINECS number 222-657-4.
Quinoline Yellow, D&C Yellow No. 10, E104, CAS 8004-92-0, 2-(2-quinolyl)indan-1,3-dione-disulphonate, food colouring substance, EINECS number 305-897-5.

Allura Red AC, E129, CAS 25956-17-6, Food Red No. 40, FD&C Red No. 40, disodium, 2-hydroxy-1-(2-methoxy-5-methyl-4-sulphonatophenylazo)naphthalene-6-sulphonate, food colouring substance, EINECS number 247-368-0.

Sunset Yellow FCF, E110, Food Yellow No. 5, FD&C Yellow No. 6, E 110, CAS 2783-94-0, Disodium 2-hydroxy-1-(4-sulphonatophenylazo)naphthalene-6-sulphonate.

Sodium benzoate, benzoic acid, E 211, E 210, CAS 532-32-1, CAS 65-85-0, food preservative, EINECS number 208-534-8.
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BACKGROUND AS PROVIDED BY THE COMMISSION

The European Commission has been informed of a recent study, funded by the UK Food Standards Agency, examining the effect which the consumption of certain food additives may have on children’s behaviour. The studies were undertaken with two age groups (3-year old children and 8- to 9-year old children) and involved the following food additives in 2 different mixtures/formulations: Tartrazine, Ponceau 4R, Carmoisine, Quinoline Yellow, Allura Red AC and sodium benzoate.

Before additives are authorised they must first be evaluated for their safety. Council Directive 89/107/EEC states that all food additives must be kept under continuous observation and must be re-evaluated whenever necessary in the light of changing conditions of use and new scientific information. Therefore when the European Commission is informed about new scientific evidence relating to a permitted food additive it requests the European Food Safety Authority to give an opinion on this new research.

TERMS OF REFERENCE AS PROVIDED BY THE COMMISSION

In accordance with Article 29 (1) (a) and 31 of Regulation (EC) No 178/2002, the European Commission asks the European Food Safety Authority to assess the results of the study and provide an opinion on the findings, taking into account, if possible, other available scientific literature in the related area.

ACKNOWLEDGEMENTS

The European Food Safety Authority wishes to thank the members of the ad hoc Working Group on the study from the Southampton University on the effect of some colours and benzoic acid on hyperactivity for the preparation of this opinion:

Jan Buitelaar, John Chr. Larsen, Manfred Doepfner, Manfred Gerlach, Michael Kenward, Alicja Mortensen, Iona Pratt (WG Chair), Diane Purper-Ouakil, Ivonne Rietjens, Terje Sagvolden, Stephen Senn, Stephan Strobel.

The Panel wishes to thank Prof. J. Stevenson, one of the authors of the McCann et al. study, for the fact that he has met with the ad hoc Working group to discuss methodological and other issues. In addition, the Panel wishes to thank the Food Standards Agency and Prof. J. Stevenson for providing EFSA with the study details.

The Panel also wishes to thank the Assessment Methodology Unit of EFSA for their assistance with the statistical re-analysis.
INTRODUCTION

It has been suggested for a number of years that exposure to synthetic food colours and other food additives may have behavioural effects, especially in young children, resulting in overactive, impulsive and inattentive behaviour (Feingold, 1975; Overmeyer and Taylor, 1999; Schab and Trinh, 2004). If severe, children who show this behaviour are likely to be diagnosed with Attention Deficit Hyperactivity Disorder (ADHD).

Attention Deficit Hyperactivity Disorder (ADHD) or Hyperkinetic Disorder (HKD) is a behavioural disorder, characterised by problems with sustained attention, impulsivity and hyperactivity, which adversely affects these children’s behaviour. ADHD typically has onset in early childhood (WHO, 2007; American Psychiatric Association, 2000). Hypotheses about the cause of ADHD have evolved from simple one-cause theories to the view that it is a complex, multi-factorial disorder caused by the confluence of many different types of risk factors (i.e., genetic, biological, environmental, psychosocial), with each factor contributing to the vulnerability to the disorder (Biederman and Faraone, 2005; Sagvolden et al., 2005). This multi-factorial view of ADHD is consistent with the observed heterogeneity in the genetics, pathophysiology and clinical manifestation of the disorder.

Most recently, a study by McCann et al. (2007) has concluded that synthetic colours plus a sodium benzoate preservative in the diet result in increased hyperactivity in 3-year old and 8- to 9-year old children in the general population. An earlier study by the same research team (the so-called Isle of Wight study) reported some evidence for adverse behavioural effects of a mixture of 4 synthetic colours and sodium benzoate, as measured by parental ratings for 3-year old children on the Isle of Wight (Bateman et al. 2004). In addition a meta analysis of double-blind Placebo-controlled trials has shown a small but statistically significant effect of synthetic food colours on the behaviour of children with hyperactivity (Schab and Trinh 2004). Other older studies however failed to identify similar behavioural effects (NIH, 1982; Harley et al., 1978a; Harley et al., 1978b; Mattes and Gittelman, 1981; Karvale and Forness, 1983).

The earlier study (Bateman et al., 2004) did not allow firm conclusions about the clinical significance of the observed effects of a series of food colours and the preservative sodium benzoate on children’s behaviour, mainly because of limitations in the study design (COT, 2007). These limitations included among others the presence of a large Placebo effect, and the fact that statistically significant effects on behaviour were only observed with parental observations and not with assessments made by independent researchers.

Subsequently the UK Food Standards Agency (FSA) set up an ad-hoc working group to consider these limitations in study design and to make recommendations on a new study design. Based on the findings of this working group, the FSA commissioned a new study via open competition in 2004, incorporating the study design changes that had been recommended by the working group. The results of this new study were recently published (McCann et al. 2007).

In this study, the effects of two combinations of Tartrazine (E102), Quinoline Yellow (E104), Sunset Yellow FCF (E110), Ponceau 4R (E124), Allura Red AC (E129), Carmoisine (E122) and sodium benzoate (E211) on children’s activity levels and attention were evaluated. Table 1 presents the chemical structures of the six synthetic colours included in the study. Five of the six food colours belong to the class of synthetic azodyes and one, Quinoline Yellow (E104), is a quinophthalone. Sodium benzoate is used as a preservative.
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Table 1. Chemical structure of the additives included in the study.

<table>
<thead>
<tr>
<th>Substance</th>
<th>Chemical formula</th>
<th>Chemical structure</th>
</tr>
</thead>
<tbody>
<tr>
<td>Allura Red AC (E129)</td>
<td>C_{16}H_{14}N_{2}Na_{2}O_{8}S_{2}</td>
<td><img src="image" alt="Allura Red AC" /></td>
</tr>
<tr>
<td>Azorubine = Carmoisine  (E122)</td>
<td>C_{20}H_{12}N_{2}Na_{2}O_{7}S_{2}</td>
<td><img src="image" alt="Azorubine" /></td>
</tr>
<tr>
<td>Ponceau 4R (E124)</td>
<td>C_{20}H_{11}N_{2}Na_{3}O_{10}S_{3}</td>
<td><img src="image" alt="Ponceau 4R" /></td>
</tr>
<tr>
<td>Quinoline Yellow (E104)</td>
<td>C_{18}H_{9}N_{2}NaO_{5}S</td>
<td><img src="image" alt="Quinoline Yellow" /></td>
</tr>
<tr>
<td>Sunset Yellow FCF (E110)</td>
<td>C_{16}H_{10}N_{2}Na_{2}O_{7}S_{2}</td>
<td><img src="image" alt="Sunset Yellow FCF" /></td>
</tr>
<tr>
<td>Tartrazine (E102)</td>
<td>C_{16}H_{9}N_{4}Na_{3}O_{9}S_{2}</td>
<td><img src="image" alt="Tartrazine" /></td>
</tr>
<tr>
<td>Sodium benzoate (E211)</td>
<td>NaC_{7}H_{5}O_{2}</td>
<td><img src="image" alt="Sodium benzoate" /></td>
</tr>
</tbody>
</table>

ASSESSMENT

EFSA’s AFC Panel, in addressing the Terms of Reference provided by the European Commission, has assessed this new study in the light of previous opinions on the compounds, and has also considered more recent studies which have become available since the publication of the available opinions on these colours and benzoate, in order to evaluate the relevance of these findings for human health.
To assist the Panel in this task a number of experts in behaviour, child psychiatry, allergy, and statistics were invited to join the ad hoc Working Group (see Acknowledgements). This ad hoc Working Group met on four occasions and prepared a detailed analysis of the McCann et al. study.

1. Study design and conduct

The study design and conduct are described in detail in McCann et al. (2007). The study consisted of a community-based double-blind, Placebo-controlled randomised cross-over food challenge in 3-year old children and in 8- to 9-year old children with two mixtures (Mix A and Mix B), each consisting of 4 different colours and sodium benzoate. A mixed fruit juice drink was used as vehicle for the food colour / sodium benzoate mixtures, and a Placebo drink was used in the wash-out periods. The Placebo and the two additive mixes were identical except for the additives, and there were no nutritional differences in the composition of the drinks. Table 2 presents an overview of the additive composition of the mixtures A and B and also presents data reflecting the actual intake levels of the additives achieved.

### Table 2. Overview of the composition of the mixtures A and B and of the actual intake levels achieved.

<table>
<thead>
<tr>
<th>Additive (E number)</th>
<th>ADI mg/kg bw</th>
<th>Mix A Daily dose in mg for 3-/8- to 9-year old</th>
<th>Mix B Daily dose in mg for 3-/8- to 9-year old</th>
<th>Mix A Daily dose in mg/kg bw* for 3-year olds (% ADI)</th>
<th>Mix B Daily dose in mg/kg bw* for 3-year olds (% ADI)</th>
<th>Mix A Daily dose in mg/kg bw* for 8- to 9-year olds (% ADI)</th>
<th>Mix B Daily dose in mg/kg bw* for 8- to 9-year olds (% ADI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tartrazine (E102)</td>
<td>7.5</td>
<td>7.5 / 9.4</td>
<td>-</td>
<td>0.5 (6.7%)</td>
<td>-</td>
<td>0.3 (4%)</td>
<td>-</td>
</tr>
<tr>
<td>Ponceau 4R (E124)</td>
<td>4.0</td>
<td>5.0 / 6.3</td>
<td>-</td>
<td>0.33 (8.25%)</td>
<td>-</td>
<td>0.2 (5%)</td>
<td>-</td>
</tr>
<tr>
<td>Sunset Yellow FCF (E110)</td>
<td>2.5</td>
<td>5.0 / 6.3</td>
<td>7.5 / 15.6</td>
<td>0.33 (13.2%)</td>
<td>0.5 (20%)</td>
<td>0.2 (8%)</td>
<td>0.5 (20%)</td>
</tr>
<tr>
<td>Carmoisine (E122)</td>
<td>4.0</td>
<td>2.5 / 3.1</td>
<td>7.5 / 15.6</td>
<td>0.17 (4.25%)</td>
<td>0.5 (12.5%)</td>
<td>0.1 (2.5%)</td>
<td>0.5 (12.5%)</td>
</tr>
<tr>
<td>Quinoline Yellow (E104)</td>
<td>10.0</td>
<td>-</td>
<td>7.5 / 15.6</td>
<td>0.5 (5%)</td>
<td>-</td>
<td>0.5 (5%)</td>
<td>-</td>
</tr>
<tr>
<td>Allura Red AC (E129)</td>
<td>7.0</td>
<td>-</td>
<td>7.5 / 15.6</td>
<td>0.5 (7.1%)</td>
<td>-</td>
<td>0.5 (7.1%)</td>
<td>-</td>
</tr>
<tr>
<td>Sodium benzoate (E211)</td>
<td>5.0</td>
<td>45 / 45</td>
<td>45 / 45</td>
<td>3.0 (60%)</td>
<td>3.0 (60%)</td>
<td>1.45 (36.3%)</td>
<td>1.45 (36.3%)</td>
</tr>
</tbody>
</table>

*The doses per kg bw were calculated by COT (2007) using average body weights for the two age groups obtained from UK National Diet and Survey data. For comparison the ADI values for these different additives are also included.

The children were selected from families volunteering from nurseries, preschool groups and playgroups for the 3-year old children and from schools in the Southampton area for 8- to 9-year old children. The children who were included in the study were selected to represent a
broad range of behaviour in the general population including children with normal to high level activity. Children who were medicated for ADHD were not included. In total one hundred and fifty three 3-year old and one hundred and forty four 8- to 9-year old children were included in the study.

The families were instructed to maintain the children during the course of the study on diets that were free of the food colours used in the study and also free of sodium benzoate. Compliance was monitored by means of a diary in which parents reported consumption levels of the test mixtures as well as compliance with the dietary requirements.

In order to investigate the hypothesis that the children’s behaviour in response to the challenge with the food colours in question could be influenced by allelic variation in a number of genes that have previously been implicated in ADHD (Thapar et al., 1999; Swanson et al., 2000; Kuntsi and Stevenson, 2001), buccal swabs were collected from the children for genotypic analyses of cellular DNA (COT, 2007). The genes studied included genes from the dopamine, adrenergic and histamine neurotransmitter systems (COT, 2007).

The amounts of the different colours in Mix A given to the 3-year old children were identical to those used in the Isle of Wight study (Bateman et al., 2004). For the 8- to 9-year old children the intakes of the different colours in Mix A were lower on a kg bw basis than for the 3-year old children whereas for Mix B the intakes on a mg per kg bw basis were the same for both age groups and higher than for Mix A. For sodium benzoate the intake in mg/kg bw was about 2 times higher for the 3-year old children as compared to the 8- to 9-year old children, but similar for Mix A and Mix B for each of age groups. The researchers indicated that the intakes of the different colours in both Mix A and Mix B for the 3-year olds and for Mix A in the 8- to 9-year olds were approximately equivalent to the amount of food colouring in two 56 gram bags of sweets. The intakes for Mix B for the 8- to 9-year old equated to about four bags of sweets a day.

During the 6 weeks of the study children received batches of the drinks on a weekly basis. Wash-out weeks (week 1, 3 and 5) in which the children received a Placebo drink, were alternated with challenge weeks (week 2, 4 and 6) during which the children received either Placebo drink, Mix A or Mix B in randomised order. The ingredients of the Placebo drink were free of the colours and preservative being tested in the challenge, and for 8- to 9-year olds the volumes of the different juices that made up the mixture and were consumed on a daily basis were as follows: 150 ml tropical juice, 80 ml red grape juice, 10 ml prune juice, 140 ml blackcurrant juice, 10 ml beetroot juice, 20 ml pear juice, 160 ml orange juice and 55 ml water, together making up a final volume of 625 ml per day. For the 3-year olds the volumes were reduced proportionately to provide 300ml a day. The Placebo drink was developed so that when each of the additive mixes in turn was introduced there were no detectable differences in taste, colour or smell. The Placebo and the two additive mixes were therefore identical except for the additives. It was thought essential to ensure that any behavioural effect attributable to ingredients of the Placebo mix, including idiosyncratic reactions from individual children were kept constant across the challenge types. In order to make the Mix palatable for children the sweetener aspartame was included, since studies on aspartame and hyperactivity have produced uniformly negative results (e.g. Wolraich et al., 1994). In contrast there is evidence that sugar can affect inattention using one of the same computerised tests adopted in this study (Wender & Solanto, 1991).

Table 3 presents a schematic overview of the crossover trial.
Assessment of the results of the study by McCann et al. (2007) on the effect of some colours and sodium benzoate on children’s behaviour

Table 3. Schematic overview of the crossover trial

<table>
<thead>
<tr>
<th>Sequence</th>
<th>Week 0</th>
<th>Week 1</th>
<th>Week 2</th>
<th>Week 3</th>
<th>Week 4</th>
<th>Week 5</th>
<th>Week 6</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Normal diet*</td>
<td>WO** Mix A</td>
<td>WO Mix B</td>
<td>WO Mix C</td>
<td>Mix C</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>(27,25***</td>
<td>(27,24)</td>
<td>(23,24)</td>
<td>(21,23)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>Normal diet</td>
<td>WO Mix A</td>
<td>WO Mix C</td>
<td>WO Mix B</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>(25,24)</td>
<td>(24,22)</td>
<td>(23,21)</td>
<td>(22,21)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>Normal diet</td>
<td>WO Mix B</td>
<td>WO Mix A</td>
<td>WO Mix C</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>(26,23)</td>
<td>(25,22)</td>
<td>(25,23)</td>
<td>(20,21)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>Normal diet</td>
<td>WO Mix B</td>
<td>WO Mix C</td>
<td>WO Mix A</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>(24,24)</td>
<td>(24,24)</td>
<td>(21,21)</td>
<td>(20,21)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>Normal diet</td>
<td>WO Mix C</td>
<td>WO Mix A</td>
<td>WO Mix B</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>(27,25)</td>
<td>(24,21)</td>
<td>(21,22)</td>
<td>(23,22)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>Normal diet</td>
<td>WO Mix C</td>
<td>WO Mix B</td>
<td>WO Mix A</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>(24,23)</td>
<td>(23,20)</td>
<td>(19,20)</td>
<td>(18,20)</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*Normal diet to set baseline levels
** Wash-out period
*** Number of children with a GHA score (Number of children in 3-Year Group, Number of children in 8- to 9-Year group) (derivation of the GHA score is described below)

Behaviour at home was assessed by parents, behaviour in the classroom was assessed by teachers and by independent observers. For the 8- to 9-year old children behaviour was also assessed by a computerised attention test. Behaviour was scored at the end of each treatment week (week 2, 4 and 6) using standardised and validated ADHD-behaviour assessment tools. The following measurement tools were used:

1. ADHD rating scale IV (teacher version). A questionnaire was completed to describe the frequency of the specific behaviours displayed over the past week, for every week of the study.

2. The abbreviated Weiss-Werry-Peters (WWP) hyperactivity scale. The WWP has been used in a number of studies to assess hyperactivity. Parents rated their child’s behaviour during the previous week for seven items previously used.

3. Classroom Observation Code (COC): The COC assesses the occurrence of 12 mutually exclusive behaviours during structured didactic teaching and during periods of independent work under teaching supervision.

4. Conners’ Continuous performance test II (CPTII). This is a test using visual stimuli of 14 minutes duration and is widely used to evaluate attention and the response inhibition component of executive control. This test was only used for the 8- to 9-year old children.

Ratings of behaviour from each of the individual measurement tools were combined, unweighted, into an overall Global Hyperactivity Aggregate (GHA) score, representing a novel metric developed by the researchers, combining observational and computerised behavioural scores in one parameter. ADHD rating, WWP and COC were used to calculate the GHA score for 3-year old children, with an additional measure (CPTII, being itself an aggregate of four computer scores) as a fourth instrument for 8- to 9-year old children.
The primary analysis of the data was based on the GHA score. A high GHA score indicates more activity. The authors indicated that although the designs for the two age groups were similar, the difference in composition of the GHA score, and in the dose of the additives used, meant that data from the two age groups could not be analysed jointly. Therefore the age groups were analysed in parallel but independently.

The data were analysed using linear mixed model methods in SPSS (Gueorguiva and Krystal, 2004; Mallinckrodt et al., 2004). Two linear mixed models were fitted for each age group. Although in the original publication (McCann et al., 2007) no details were given on the parameterisation of the model, the raw output of the analysis provided to EFSA gave detailed information. The first model is a basic mixed effect model where a random effect is put on the subject and a fixed effect on the treatment. Model 2 undertaken by the authors included some additional fixed effects (see the separate statistical report for further details).

A compound symmetry covariance matrix provided best fit for the models fitted to the data of the first age group while an unstructured covariance matrix gave the best fit for the second age group. The choice of covariance structure was done based on log-likelihood ratio comparisons for Model 1 only, accounting for the total number of parameters to be estimated.

The analyses were replicated for three sets of data: the full study population sample, a high consumption subset (data included if the child consumed \( \geq 85\% \) of drinks in each treatment week) and a complete case subset data (high consumption and no missing GHA scores).

To test whether there was evidence of carry-over effects, the scores of the previous active challenge period and baseline were added as factors in the mixed model. No effect due to the type of challenge in the previous period on the current scores could be demonstrated. From this, it was concluded by the study authors that the wash-out periods were sufficiently long to have prevented carry-over effects.

### 2. AFC Panel comments on study design and conduct

The Panel notes that:

- The study tested two mixtures and no individual compounds. Testing of mixtures cannot identify the hazards of individual compounds. The choice to test mixtures was based on the fact that the Isle of Wight study (Bateman et al. 2004) had also tested a mixture, and part of the objective of the new study was to investigate whether the findings of the Isle of Wight study could be replicated with a better study design. Mix A reflects the mixture tested in this earlier study. Mix B reflects a mixture representative for sweets as they are consumed in the UK;

- Parents, teachers and independent observers scored the behaviour of the children. The outcome of their scores was combined, and for 8- to 9-year old children the results of the CPTII computer tests were also included, to give the GHA scores. In this way observational scores were combined with computer scores;

- Including parental scores into the GHA score the study design does not completely overcome the criticisms of the earlier Isle of Wight study that effects on behaviour were only observed via parental observations and not via assessments made by independent researchers;

- The GHA score combines three measures of behaviour for the 3-year old children, adding the computer-based measure (CPTII) for the 8- to 9-year old children only. The Panel noted that using this aggregated score is adequate from a statistical point of view,
because it is considered not to affect the integrity of the statistical approach. Combining the measures does not increase the chance of introducing statistical differences which do not actually exist;

• The aggregated score is adequate to score an overall change in behaviour, but that it is not a clinically accepted and validated outcome in that it has not been assessed whether it shows meaningful relations with external variables such as prognosis and impairment of functioning in other areas of behaviour;

• The combined GHA scores reflect a global score that may be difficult to interpret behaviourally and statistically. Therefore the Panel notes that also subsequent analyses on each of the individual behaviour variables would be helpful in order to assess the relative contributions of each behavioural pattern;

• The drinks were given at home to guarantee better compliance. However, the time of day at which the children consumed the drink was not regulated. In addition, the durations of the possible behavioural effects are not known. These factors combined may have influenced the results since transient effects may not have been observed in some of the tests;

• The study was not designed to explore possible dose: response relationships or possible subgroups of “responders”;

• Self-volunteering of the subjects included in the study might have introduced a selection bias;

• Although the study designs for the two age groups were similar, the difference in composition of the GHA scores and the dose of additives used meant that data from the two age groups cannot be analysed jointly;

• A one week wash-out period was chosen which was also the period used in the Isle of Wight study (Bateman et al. 2004). There was no evidence of carry-over effects from a challenge week into the next challenge week based on statistical analyses. Analysis of the assessment of behaviour during the wash-out period would have provided a clue on the efficiency of the wash-out period and/or possible effects of the Placebo;

• Based on data on toxicokinetics in experimental animals, it may be concluded that the test substances, with the possible exception of Quinoline Yellow, will be eliminated from the body during the wash-out periods. Quinoline Yellow has a longer half-life and some parent compound or breakdown products thereof may still be present in the body after the one week wash-out period (see Appendix A).

2.1. Comments on statistical analysis

General remarks and discussion on the design

Basic principles of cross over trials are that every subject (child) receives each of the treatments being evaluated over a standard period of time while each subject serves as its own control and the outcome variables are assessed in the same way in each period of the treatment. The major advantage of such a design is the reduction of the sample size (number of subjects) needed to achieve a certain statistical power. However, several disadvantages are inherent to a cross over trial design: the treatment must have a reversible effect; the statistical analysis is rather complicated and if not carried out correctly, may lead to erroneous conclusions. Also carry-over effects are possible, i.e. the residual effect of one treatment on the outcome of a
Assessment of the results of the study by McCann et al. (2007) on the effect of some colours and sodium benzoate on children’s behaviour

subsequent treatment. This is particularly a potential concern if no prior pharmacokinetic (PK) and pharmacodynamic (PD) assessment of the tested food additives is available to support the chosen length of the wash-out period. In the McCann et al. (2007) study the claim of absence of carry-over effects was not supported by preliminary investigation on the pharmacokinetics of the compounds, although a formal statistical test for carry-over was performed and no statistically significant carry-over effect was found.

Discussion on the statistical methodology used in the McCann et al. paper

Analyses were based on two linear mixed models. The first and simplest model contained treatment effects only, in addition to those for random variability. With a cross-over design it is recommended that adjustment be made for possible period effects; indeed, in this study, there is clear evidence of differences between the different treatment weeks.

The second model was an extension of the first in which, in addition, period effects and a number of between-subject terms, including baseline GHA score, were introduced. The incorporation of between-subject effects is pointless in a cross-over design like this because such effects are anyway effectively eliminated from the analysis by the within-subject nature of the treatment comparisons. This is illustrated clearly in Tables 3 and 4 in the McCann et al. study (McCann et al., 2007), where the differences in treatment effects estimates from the two models are of little or no importance. The choice of covariance structure is very unlikely to be crucial in the analysis of data from a design like this, provided within-subject dependence is accommodated.

The steps taken for score normalisation and aggregation are mathematical transformations that might affect the assumptions of normality and independence of the data which are essential for the whole statistical analysis. Moreover, in such cross-over trials, since each subject serves as its own control, individual scores should be compared to individual baseline scores, not to the group mean baseline score.

2.2. Comments on dietary exposure levels used in the study

The researchers indicated that the daily dietary exposure to the different colours for both Mix A and Mix B for the 3-year olds and for Mix A in the 8-to 9-year olds were approximately equivalent to the amount of food colouring in two 56 g bags of sweets and that the daily dietary exposures for Mix B for the 8- to 9-year olds equated about four bags of sweets a day (McCann et al., 2007).

The Panel was provided with information allowing it to assess if the level of exposure considered in the study was likely to occur, based on current legislation, current actual uses and use levels in foods consumed by children and current levels of consumption of these foods by children.

The amounts in Mix A given to 3-year olds were identical to those used in the previous (Isle of Wight) study, while for 8- to 9-year olds the amounts of the colours in Mix A were increased by 25% to reflect the greater food intake by the older children.

According to COT (COT, 2007, based on information from the Food Standards Agency, UK), for 8- to 9-year olds, the amounts of the colours in Mix B reflected what a child could reasonably consume in one day, based on average consumption of foods containing colours at their maximum permitted levels (MPL).

The two main sources of added colours in children are soft drinks and confectionery. The Panel noted that according to current legislation, all six target colours may be used singly or in
Assessment of the results of the study by McCann et al. (2007) on the effect of some colours and sodium benzoate on children’s behaviour

combination to a cumulative MPL of 300 mg/kg in confectionery and to a cumulative MPL of 100 mg/l in soft drinks. Mix A provided overall 20 mg of artificial colours for 3-year old children and 25 mg for children aged 8 to 9 years. Mix B provided overall 30 mg of colours for 3-year old children and 62.4 mg for children aged 8 to 9 years. These levels of dietary exposure could be reached through consumption of 67 g to 208 g of confectionery containing artificial colours at their cumulative MPL. They could also be reached through consumption of 200 ml to 624 ml of soft drinks containing artificial colours at their cumulative MPL.

In addition, three of the target colours (Sunset Yellow FCF, Carmoisine and Ponceau 4R) have a lower individual MPL in the legislation: 50 mg/l in soft drinks and 50 mg/kg in confectionery. The individual dietary exposure to these colours in the study protocol was highest in Mix B: up to 7.5 mg in 3-years old children and up to 15.6 mg in 8- to 9-years old children (for Sunset Yellow FCF and Carmoisine). These levels of dietary exposure to either Sunset Yellow FCF or Carmoisine could be reached through consumption of respectively 150 g and 312 g of sweets or through the consumption of respectively 150 ml and 312 ml of soft drinks containing one of these colours at its individual MPL.

The main source of benzoates in children are soft drinks and the MPL for benzoates is 150 mg/l. Both Mix A and Mix B provided 45 mg of sodium benzoate in the two age groups. This level of exposure could be reached through consumption of 300 ml of soft drink containing benzoate at its MPL.

The Panel noted that the dietary exposures to the colours used in the study, were well below the ADIs of the individual substances. As shown in Table 2, the dietary exposures in the 3-year old children ranged from 4.3% to 13.2% of the ADIs for the individual colours in Mix A. For the 8- to 9-year old children these values for Mix A ranged from 2.5% to 8% of the ADIs for the individual colours. For the colours in Mix B the dietary exposure of the 3-year old children and the 8- to 9-year old children were similar, ranging from 5 to 20% of the ADIs of the individual colours. The dietary exposures for the two age groups were different for sodium benzoate, amounting to respectively 60% and 36.3% of the ADI for sodium benzoate for the 3-year and 8- to 9-year old children respectively.

Connolly and co-workers have investigated the frequency of occurrence of the food colours and sodium benzoate used in the McCann et al. study in a recent 7-day dietary survey of 594 Irish children aged 5-12 years (Unpublished data by Connolly et al., 2008). The food consumption data, coded at brand level, were combined with the Irish National Food Ingredient Database in which all ingredients listed on the label of food items, including additives, are recorded, (Gilsenan et al., 2002). In the case of sodium benzoate, the presence as a natural ingredient in the food was not considered. Among the 5,551 individual food items coded at brand level that were consumed during the survey, 279 (5%) contained at least one of the target additives. The percentage of child food-eating occasion containing the target artificial colour ranged from 138/72,024 (0.2%) for Tartrazine to 555/772,024 (0.8%) for Sunset Yellow FCF. Tartrazine, which is authorised for use in “processed mushy and garden peas (canned)” was found to occur most frequently in the food group “Peas, Beans and Lentils”; The other five colours occurred most frequently in “Chocolate and non chocolate confectionary”. Other food groups containing the target colours were “Cakes, Pastries & Buns” (in particular for Carmoisine) and carbonated beverages (in particular for Sunset Yellow FCF). The frequency of occurrence for sodium benzoate was 2183/72,024 (3%); its most frequent source was beverages. The total number of observed child-days was 4158. At least one target additive occurred in 30.5% of child-days, two additives occurred in 7.7% of child-days, three additives in 5.1%, four additives in 2.8%, five additives in 2.2% and six or seven additives in 0.7% of child days. The Panel noted that the data provided do not allow the estimation of the percentage of children who were exposed to the colours or their combination in at least one of the survey day. This percentage would have been useful but was not provided in the Connolly Report.
A usage survey conducted by the Union of European Beverage Associations (UNESDA) in 2005 was made available to the Panel (Tennant, 2006). The survey report indicates that all but one of the six colours considered in the study by McCann et al. are commonly used in soft drinks, with Carmoisine being an uncommon artificial colour in these products. However, other surveys described underneath suggest that Carmoisine is also commonly used.

Data from three *ad hoc* surveys in which analytical determinations of artificial colours were performed in retail products were also provided to the Panel: an unpublished survey conducted in 2005 by the Food Safety Authority of Ireland (FSAI) in 34 retail ready to drink soft drinks, a survey by the UK Food Standards Agency in 201 retail ready to drink soft drinks selected for being distinctly coloured (FSA, 2003) and a survey by the UK Food Standards Agency in 196 retail samples of brightly coloured packaged sweets (FSA, 2002).

The frequency of occurrence of each of the artificial colours under study and the range of analytical values observed in soft drinks and sweets (when the colour was present) are reported in Table 4.

Among colours, Tartrazine was present with the lowest frequency: in respectively 6%, 1.5% and 3% of Irish soft drinks, UK soft drinks, and UK sweets. The colour most frequently present was Quinoline Yellow: in respectively 21%, 37% and 56% of Irish soft drinks, UK soft drinks, and UK sweets. In sweets, the overall concentration of the target colours was up to 208 mg/kg in sweets, lower than the cumulative MPL of 300 mg/kg in sweets. In the same table, the quantity of either sweets or beverages that needs to be consumed to lead to the level of exposure of the experimental protocol was calculated for each group of children in order to verify if the levels of exposure considered are in line with potential level of exposure in children consuming products present on the market. Calculations were based on the upper concentration values observed in the surveys on retail products. For sweets the quantity varied from a minimum of 42 g needed to reach the dietary exposure to Allura Red AC in Mix B for 3-year old children to a maximum of 363 g needed to reach the dietary exposure to Carmoisine in Mix B for 8- to 9-year old children. In the case of beverages, the quantity varied from a minimum of 42 ml needed to reach the dietary exposure to Carmoisine in Mix A for 3-year old children to a maximum of 371 ml needed to reach the dietary exposure to Allura Red AC in Mix B for 8- to 9-year old children.

According to UK FSA diary survey on the consumption of soft drinks by young children (FSA 2003 b), high level consumers of 1.5 to 4.5 years drank around 500 ml (just over one and a half 330 ml cans) a day.

In the UK survey, co-occurrence of two of the colours under study was observed in 41 soft drinks, co-occurrence of three of the colours under study was observed only in two soft drinks, co-occurrence of three or four of the colours under study was not observed. In the Irish survey, co-occurrence of two of the colours under study was observed in 4 soft drinks, co-occurrence of three of the colours under study was observed in one soft drink, co-occurrence of three or four of the colours under study was not observed. The co-occurrence of two, three or four of the target colours was more frequent in sweets, as observed in the UK survey. When two or more colours under study occurred in the same product, the overall concentration reached the cumulative MPL of 100 mg/l in a number of soft drinks (up to 106 mg/l in the UK survey).

In conclusion, the target artificial colours and benzoate were found to be used in foods consumed by children in surveys conducted in years 2002 to 2005. The target colours were more frequently used in sweets but also occurred commonly in soft drinks, benzoate was very frequently present in beverages. Children consuming brightly coloured sweets may be exposed to levels comparable to those considered in the protocol of the McCann *et al.* study for one or more of the food colours studied. Comparable levels may also be reached in those children who...
consume brightly coloured soft drinks. The level of exposure to sodium benzoate is also likely to occur.
Assessment of the results of the study by McCann et al. (2007) on the effect of some colours and sodium benzoate on children’s behaviour

Table 4. Actual use levels of colours used in the Southampton study and quantities of sweets and beverages corresponding to the different Mixes

<table>
<thead>
<tr>
<th>Analytical survey of the UK Food Standards Agency conducted in 2002 in England in brightly coloured retail ready to drink soft drinks (UK FSA, 2003)</th>
<th>Survey of the Food Safety Authority of Ireland conducted in 2005 (unpublished)</th>
<th>Quantity of soft drink (ml) at the highest observed concentration(*) corresponding to the exposure to individual colours in Mix A and Mix B</th>
<th>Analytical survey of the UK Food Standards Agency conducted in England in 2000/2001 in brightly coloured sweets (UK FSA, 2002)</th>
<th>Quantity of sweets (g) at the highest observed concentration(*) corresponding to the exposure to individual colours in Mix A and Mix B</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Occurrence in excess of LOD</strong></td>
<td><strong>Range of analytical data above LOD (mg/l)</strong></td>
<td><strong>Occurrence in excess of LOQ</strong></td>
<td><strong>Range of analytical data above LOQ (mg/l)</strong></td>
<td><strong>3-year olds</strong></td>
</tr>
<tr>
<td>Allura Red AC</td>
<td>6/201</td>
<td>9-42</td>
<td>2 / 34</td>
<td>20–32</td>
</tr>
<tr>
<td>Ponceau 4R</td>
<td>32/201</td>
<td>1-47</td>
<td>6/34</td>
<td>3-22</td>
</tr>
<tr>
<td>Tartrazine</td>
<td>3/201</td>
<td>3-28</td>
<td>2/34</td>
<td>5-25</td>
</tr>
<tr>
<td>Quinoline Yellow</td>
<td>75/201</td>
<td>1-92</td>
<td>7/34</td>
<td>1-34</td>
</tr>
<tr>
<td>Sunset Yellow FCF</td>
<td>61/201</td>
<td>1-61</td>
<td>7/34</td>
<td>11-49</td>
</tr>
<tr>
<td>Carmoisine</td>
<td>64/201</td>
<td>1-45</td>
<td>13/34</td>
<td>1-59</td>
</tr>
</tbody>
</table>

*The concentration value used is the highest range of analytical data observed in the UK and Ireland survey (evidenced in bold character in the columns reporting the ranges of analytical data)
3. Results of the study

3.1. Overview of the findings of the study (as reported by the authors)

For reasons unrelated to effects, 16 of the one hundred and fifty three 3-year old children and 14 of the one hundred and forty nine 8- to 9-year old children did not complete the study.

Tables 5 and 6 provide an overview of the results from Model 1 of the two linear mixed models used by the authors, obtained for three groups of participants from each age group: (1) the full study population sample (2) those with at least 85% consumption of the trial drinks (≥ 85% consumption), and (3) those with at least 85% consumption of the trial drinks and observations from all three periods (complete case). The latter, (2) and (3), represent subgroups of the original trial samples.

Table 5. GHA score estimates during challenge period for 3-year old children (taken from McCann et al. 2007‡)

<table>
<thead>
<tr>
<th></th>
<th>Entire sample (n=140)</th>
<th>Group with ≥85% consumption (n=130)</th>
<th>Complete case group, ≥85% consumption and no missing data (n=73)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mix A vs. Placebo</td>
<td>0.20 (0.01 to 0.39)*</td>
<td>0.28 (0.05 to 0.51)*</td>
<td>0.32 (0.05 to 0.60)*</td>
</tr>
<tr>
<td>Mix B vs. Placebo</td>
<td>0.17 (−0.03 to 0.36)</td>
<td>0.19 (−0.04 to 0.41)</td>
<td>0.21 (−0.06 to 0.48)</td>
</tr>
</tbody>
</table>

Values given in the Table represent estimates (95% CI) of the differences in GHA mean scores between the Placebo and the challenge for each population group, based on the mean baseline scores at week 0 and the mean scores following treatment.

*p<0.05.
‡ The figures quoted above are those used by the authors in their discussions of the new findings and represent the outcomes from application of statistical model 1 or model 2.

Table 6. GHA score estimates during challenge period for 8- to 9-year old children (taken from McCann et al. 2007‡).

<table>
<thead>
<tr>
<th></th>
<th>Entire sample (n=136)</th>
<th>Group with ≥85% consumption (n=119)</th>
<th>Complete case group, ≥85% consumption and no missing data (n=91)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mix A vs. Placebo</td>
<td>0.08 (−0.02 to 0.17)</td>
<td>0.09 (−0.01 to 0.19)</td>
<td>0.12 (0.02 to 0.23)*</td>
</tr>
<tr>
<td>Mix B vs. Placebo</td>
<td>0.12 (0.03 to 0.22)*</td>
<td>0.15 (0.05 to 0.25)†</td>
<td>0.17 (0.06 to 0.28)†</td>
</tr>
</tbody>
</table>

Values given in the Table represent estimates (95% CI) of the differences in GHA mean scores between the Placebo and the challenge for each population group, based on the mean baseline scores at week 0 and the mean scores following treatment.

*p<0.05. †p<0.01
‡ The figures quoted above are those used by the authors in their discussions of the new findings and represent the outcomes from application of statistical model 1 or model 2.

Mix A significantly increased GHA scores for all 3-year old children compared to the Placebo control GHA scores (effect size 0.20 [CI 0.01 to 0.39], p<0.05).

Mix B had no effect on GHA scores in 3-year old children as compared to the Placebo control GHA scores (effect size 0.17 [CI -0.03 to 0.36]).
This result persisted when analysis was restricted to 3-year old children who consumed more than 85% of juice and had no missing data; in these analysis the effect of Mix A in the 3-year old children was still significantly increased compared to Placebo control (effect size 0.32 [CI 0.05 to 0.60, p<0.05) but for Mix B no significant effect on GHA scores was observed (effect size 0.21 [CI -0.06 to 0.48]).

For the 8- to 9-year old children a significant effect of Mix A (effect size 0.12 [CI 0.02 to 0.23, p<0.05) and of Mix B (effect size 0.17 [CI 0.07 to 0.28, p<0.01) was observed when analysis was restricted to those children consuming at least 85% of drinks with no missing data. When all children that completed the study were taken into account in 8- to 9-year old children Mix A had no effect on the GHA scores compared to the Placebo control (effect size 0.08 [CI -0.02 to 0.17], and Mix B had a significant effect on GHA scores (effect size 0.12 [CI 0.03 to 0.22] p<0.05).

Post-hoc analysis reported in the statement of the Committee on Toxicity (COT, 2007) revealed that the parental reports were the main contributors to the changes in GHA scores for the 3-year old children, whereas in the 8- to 9-year old children the largest contribution to the GHA score was reported to come from the computer-based task (COT, 2007). The researchers suggested that parents may have been more sensitive to or more exposed to behavioural changes in their children than the independent observers or teachers.

As also reported by the UK COT, the research team found that the observed increases in the GHA scores with Mix A in 3-year olds and 8- to 9-year olds and with Mix B in 8- to 9-year olds were statistically significantly associated with differences in genotype, specifically with two genetic polymorphisms thought to impair histamine clearance (COT, 2007). This analysis was carried out in the subgroup of children with at least 85% consumption of the trial drinks.

The authors concluded on the basis of these results that synthetic colours and/or a sodium benzoate preservative in the diet may exacerbate hyperactive behaviours (inattention, impulsivity, and overactivity) in 3-year old and 8- to 9- year old children in the general population.

### 3.2. AFC Panel assessment of observed effects in the study

The Panel noted that:

- Small but statistically significant effects of Mix A but not of Mix B on GHA scores in 3-year old children were described. In 8- to 9-year old children, when taking all children that completed the study into account, a small but significant effect of Mix B but not of Mix A on GHA scores was observed in 8- to 9-year old children. Thus, the statistically significant effects were not found for the same mixture in the two age groups;

- For Mix A the doses on a mg/kg bw/day basis were higher in the 3-year old children and this may have contributed to the difference in the magnitude of the effect of Mix A in the two age groups. For Mix B the doses in both age groups were similar but the effects were significant only in the 8- to 9-year old age group;

- The effects of Mix A on behavioural parameters in the 3-year old children were consistent with those of the Isle of Wight study, showing an increase in GHA score in the 3-year old children in the present McCann et al. study and an effect on the basis of only parental observations in the Isle of Wight study;
• The main contributors to the statistically significant effect on the GHA scores in the 3-year old children were the parental scores, as described in more detail in Section 3.3 below. The scores from teachers and independent observers were not a major component in the overall GHA scores. The use of the GHA scores does not therefore completely overcome the criticisms on the earlier Isle of Wight study;

• Since each subject serves as its own control, no further explanation of variation in the trial can be achieved by fitting subject level covariates. Only covariates that changed over time for a given subject could have any further explanatory power. Therefore the extra information obtained from model 2 used by the authors of the study is minimal, as shown in their publication in Tables 3 and 4. Apart from the fact that a period effect was fitted the additional factors will not explain the variation of the GHA scores;

• The clinical significance of the observed effects (a) for the individual children in the study and (b) for the population as a whole remains unclear, since the effects were small in magnitude and these small alterations in attention and activity may not interfere with schoolwork and other intellectual functioning.

3.3. Statistical re-analysis and AFC Panel assessment

The Panel considered that the steps taken for score normalisation and aggregation are mathematical transformations that might affect the assumptions of normality and independence of the data which are essential for the whole statistical analysis. Therefore, the authors’ primary analysis was repeated using a more justifiable and conventional statistical model, and this was supplemented by a set of additional analyses with the aim of aiding the interpretation of the results.

Details of the statistical re-analysis can be found in the separate statistical report.

The re-analysis consisted of two parts. First, the authors’ primary analysis was repeated, with minor changes to reflect a more appropriate statistical treatment and, second, a set of supplementary analyses were carried out.

For the primary analysis the Global Hyperactivity Aggregate (GHA) score was recalculated following the same steps as in the original analysis, except for the omission of the final re-normalisation step.

The remainder of the supplementary analyses consisted of the calculation of various descriptive statistics and formal analysis of each of the individual component measures.

For all formal analyses, both primary and supplemental, a linear mixed model was used that was similar to that of the first analysis reported in the Lancet paper (McCann et al. 2007). The model included only within-subject effects, namely those associated with the experimental intervention and with periods. Random subject effects were also included, and in this setting imply an analysis identical to that with a compound symmetry covariance structure. The “week” variable was also included as a fixed effect in the model. Consumption subgroup analyses matched those of the original paper.

The p-values were calculated for the contrast ‘Mix A vs. Placebo’ and ‘Mix B vs. Placebo’ for the whole dataset, the ≥ 85% consumers and the children consuming at least 85% of drinks with no missing data (complete case group). Models were run for all combinations of sexes and age groups.
Table 7 presents a summary of all statistically significant cases found in the re-analysis using the Global Hyperactivity Aggregate (GHA) score similar to that carried out in the McCann et al. study.

Table 7. Summary of all significant cases found in the statistical reanalysis in the treatment group comparisons for the GHA score, using a similar approach to that used in the McCann et al. study.

<table>
<thead>
<tr>
<th>Test</th>
<th>Year Group</th>
<th>Sex</th>
<th>Estimate</th>
<th>Std Err</th>
<th>P-Value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td><strong>entire sample</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mix B vs. Placebo</td>
<td>8/9Y</td>
<td>M</td>
<td>0.1115</td>
<td>0.04394</td>
<td>0.0124</td>
</tr>
<tr>
<td>Mix B vs. Placebo</td>
<td>8/9Y</td>
<td>both</td>
<td>0.05963</td>
<td>0.02981</td>
<td>0.0466</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td><strong>≥85% consumption</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mix B vs. Placebo</td>
<td>8/9Y</td>
<td>both</td>
<td>0.08348</td>
<td>0.03276</td>
<td>0.0116</td>
</tr>
<tr>
<td>Mix A vs. Placebo</td>
<td>3Y</td>
<td>both</td>
<td>0.1962</td>
<td>0.08074</td>
<td>0.0161</td>
</tr>
<tr>
<td>Mix B vs. Placebo</td>
<td>8/9Y</td>
<td>M</td>
<td>0.1116</td>
<td>0.04644</td>
<td>0.0179</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td><strong>complete case</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mix B vs. Placebo</td>
<td>8/9Y</td>
<td>both</td>
<td>0.08546</td>
<td>0.03536</td>
<td>0.0167</td>
</tr>
<tr>
<td>Mix A vs. Placebo</td>
<td>3Y</td>
<td>both</td>
<td>0.2359</td>
<td>0.09764</td>
<td>0.0169</td>
</tr>
<tr>
<td>Mix B vs. Placebo</td>
<td>8/9Y</td>
<td>M</td>
<td>0.1118</td>
<td>0.04993</td>
<td>0.0273</td>
</tr>
</tbody>
</table>

Based on these results it is concluded that the primary analysis with the recalculated GHA score led to broadly similar conclusions to that in the original paper by McCann et al., except for the following:

1. The Mix A versus Placebo comparison was not statistically significant for the 3-year olds when all subjects were included (entire sample), while the significance for the ≥ 85% consumption and complete case groups was increased slightly;

2. For the 8- to 9-years age group, the Mix A versus Placebo comparison was no longer statistically significant in any of the three consumption groups.

The Panel considers the re-analysis presented here in which all single variables (minus the individual baseline value for that variable) were reanalysed without normalisation, so that each subject served as its own references baseline, as the most adequate.

In addition the data were analysed on the basis of a modified GHA score in which the parental scores were not included. The results from this analysis no longer revealed any statistically significant effects of Mix A or Mix B versus Placebo, except for Mix B in the 8- to 9-year old completers when both sexes are pooled (p=0.042).

A further analysis was carried out on the whole data set, comprising analysis of the single variables of parental scores, teacher scores and observer scores, and, in the case of 8- to 9-year old children, computer-based scores. Table 8 presents a summary of all statistically significant cases found in the re-analysis of the single variables for all three in treatment groups.
Table 8. Summary of all statistically significant cases found in the re-analysis of single variables for all three treatment groups.

<table>
<thead>
<tr>
<th>Test</th>
<th>Score</th>
<th>Year Group</th>
<th>Sex</th>
<th>Estimate</th>
<th>Std Err</th>
<th>P-Value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Entire sample</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mix A vs. Placebo</td>
<td>CPTCom</td>
<td>8/9Y</td>
<td>M</td>
<td>5.0607</td>
<td>2.1497</td>
<td>0.0206</td>
</tr>
<tr>
<td>Mix B vs. Placebo</td>
<td>Parent</td>
<td>8/9Y</td>
<td>both</td>
<td>0.9017</td>
<td>0.4186</td>
<td>0.0322</td>
</tr>
<tr>
<td></td>
<td></td>
<td>&gt;85% consumption</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mix A vs. Placebo</td>
<td>CPTCom</td>
<td>8/9Y</td>
<td>M</td>
<td>5.0867</td>
<td>2.0922</td>
<td>0.0172</td>
</tr>
<tr>
<td>Mix B vs. Placebo</td>
<td>CPTCom</td>
<td>8/9Y</td>
<td>M</td>
<td>6.0375</td>
<td>2.0077</td>
<td>0.0035</td>
</tr>
<tr>
<td></td>
<td>Mix B vs. Placebo</td>
<td>Parent</td>
<td>both</td>
<td>0.9687</td>
<td>0.4584</td>
<td>0.0359</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Complete case</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mix A vs. Placebo</td>
<td>CPTCom</td>
<td>8/9Y</td>
<td>M</td>
<td>5.5414</td>
<td>2.2611</td>
<td>0.0166</td>
</tr>
<tr>
<td>Mix B vs. Placebo</td>
<td>CPTCom</td>
<td>8/9Y</td>
<td>M</td>
<td>6.3442</td>
<td>2.2427</td>
<td>0.006</td>
</tr>
<tr>
<td></td>
<td>Mix A vs. Placebo</td>
<td>Parent</td>
<td>both</td>
<td>1.2448</td>
<td>0.5585</td>
<td>0.0274</td>
</tr>
</tbody>
</table>

The main results are as follows.

- No statistically significant component effects were observed that did not coincide with effects seen already in the authors’ overall GHA analysis;
- For the 3-year olds, only the Mix A versus Placebo effect with the parental score was statistically significant in the complete case group. The teacher or observer scores showed no evidence of an effect in the 3-year olds in any consumption group;
- For the 8- to 9-year olds, statistically significant Mix B versus Placebo effects were seen for the parental scores in the entire sample group and the >85% consumption group, and for the computer scores of the males in the >85% consumption group and the complete case group;
- For the computer scores there were statistically significant Mix A versus Placebo effects in the 8- to 9-year old males in all three consumption groups;
- Consistency across consumption groups could not be observed except for the computer score for males.

In conclusion, there is a suggestion from these analyses that the statistically significant effects seen in the 3-year olds (Mix A versus Placebo) and in the 8- to 9-year olds (Mix B versus Placebo) are largely driven in the data by the parental scores and, in the older males in both comparisons, by the computer score.

From the data presented in the separate statistical report, it can be derived that a ‘Week’ effect was shown on both single and aggregated scores but only for the “Week 4 vs. Week 6” comparison in the latter. The general trend was that hyperactivity generally went up from Week 2 to Week 4 and then significantly decreased from Week 4 to Week 6. The size of this period effect was globally of the same order of magnitude as those observed for the treatment effects, in both single and aggregated scores. It illustrates a large intra-individual variability over time and interpretation of all statistical results should be done in the light of this result.
4. Overview of previous studies of the effect of food colours and sodium benzoate on behaviour.

Since Feingold’s (1975) initial report that a diet free of synthetic food colours and flavours and naturally occurring salicylates resulted in an improvement in behaviour of hyperactive children, several studies have investigated the possible relationship between exposure of young children to synthetic food colours and other food additives and behavioural effects. The AFC Panel has (briefly) reviewed these previous studies, in particular reports involving the food additives used in the McCann et al. study. These studies are summarised in Table 9.

Table 9. Summary of behavioural toxicity studies carried out on food colours and other additives

<table>
<thead>
<tr>
<th>Reference</th>
<th>Numbers of subjects, additive tested and dose levels</th>
<th>Outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td>Feingold 1975</td>
<td>194 HDL children from five separate dietary programs following a diet eliminating synthetic food colours (including Tartrazine = Yellow dye 5), flavours and naturally occurring salicylates.</td>
<td>Improvement in behaviour and learning abilities.</td>
</tr>
<tr>
<td>Conners et al., 1976</td>
<td>15 hyperkinetic children (6-13 yr) in a double-blind crossover trial involving a control diet and a diet eliminating synthetic flavours, colours and natural silicates as recommended by Feingold.</td>
<td>Teachers and parents observed the children for one month prior to treatment, using standardised rating scales. Both teachers and parents reported fewer hyperkinetic symptoms on the experiment test diet as compared to the pre-treatment baseline. The control diet ratings did not differ from the baseline period ratings. The teachers noted a significant reduction in symptoms on the experimental diet compared to the control diet but parents did not.</td>
</tr>
<tr>
<td>Williams et al., 1978</td>
<td>26 Hyperactive children (5-12 yr) given active or Placebo medications in combination with challenge cookies with synthetic food colours (red dyes 2, 3 and 4; blue dyes 1 and 2; Yellow dyes 5 and 6; green dye 3; orange dye B at levels estimated to be equal to one-half the daily dietary intake of children in the USA) or control cookies without the additives. The children were crossed over into each of 4 treatments and assessment was double blind by teachers and parents.</td>
<td>Stimulant medications were more effective than diet in reducing hyperactive behaviour. The behaviour of 3 to 8 children was diet responsive. It was concluded that especially the teachers (but not the parents) ratings provided support of Feingold’s hypothesis that food additives trigger the hyperactive response.</td>
</tr>
<tr>
<td>Harley et al., 1978a</td>
<td>9 Hyperactive male subjects, selected on the basis of showing a favourable response to the Feingold diet in an earlier study were maintained on a strict elimination (Feingold) diet for 11 weeks, and given multiple trials of Placebo and challenge food materials in a double-blind challenge experiment. Challenge food items (candy bars and cookies) contained a blend of half the average daily intake of 27 mg of certified food colours.</td>
<td>Parental and teacher ratings, classroom behaviour observations and neuropsychological test scores obtained during baseline, Placebo and challenge conditions in general were not found to be adversely affected by the synthetic colour challenge materials.</td>
</tr>
<tr>
<td>Reference</td>
<td>Numbers of subjects, additive tested and dose levels</td>
<td>Outcome</td>
</tr>
<tr>
<td>---------------------------</td>
<td>----------------------------------------------------------------------------------------------------------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Harley et al., 1978b</td>
<td>36 school-age hyperactive boys under experimental (restricted Feingold K-P) and control diet.</td>
<td>Parent’s behavioural ratings on 10 hyperactive children indicated a positive response to the experimental diet but laboratory observations showed no diet effect. It was concluded that teacher ratings, objective classroom and laboratory observational data, attention-concentration and other psychological measured yielded no support for the Feingold hypothesis.</td>
</tr>
<tr>
<td>Levy et al. 1978</td>
<td>22 children (4-8 years) selected as hyperactive tested before and after 4 weeks on the elimination diet, after a Tartrazine (Yellow dye no 5) and Placebo challenge, and after a 4 week wash-out period on the diet, by Conners parent-teacher ratings, objective tests of attention, standard perceptual motor tests and subtests from the Wechsler Intelligence Scale for Children (WISC).</td>
<td>Statistically significant improvement in the mother’s ratings of the children’s behaviour after the first 4 weeks of the diet, but the effects was not substantiated by the objective tests (teacher, clinician ratings). Tartrazine did not result in a statistically significant deterioration in the children’s behaviour when they were challenged under double blind testing conditions with Tartrazine in any of the ratings (mothers, teachers, clinicians or as measured by objective tests).</td>
</tr>
<tr>
<td>Goyette et al. 1978</td>
<td>Two double blind challenge trials:</td>
<td>1) Subjects demonstrated a 57 percent mean reduction in behaviour problems as rated by parents and a 34 percent reduction as rated by teachers when placed on the elimination diet. Double blind testing with challenge and Placebo materials revealed no significant challenge differences. The authors indicated that the ratings may have been insensitive due to the long time span of the observation period as compared to the short duration of the effect 2) subjects demonstrated a 45 percent mean reduction in behaviour problems when placed on the elimination diet. A significant challenge effect (p&lt;0.025) was observed with more behavioural problems reported during the active challenge period as compared to the Placebo period.</td>
</tr>
<tr>
<td>Rose 1978</td>
<td>Two 8-year-old females who had been on a Feingold diet for a minimum of 11 months were studied in a double-blind Placebo controlled study. Data were obtained by trained observers in the subjects regular class settings.</td>
<td>It was concluded that there was a functional relationship between the ingestion of synthetic food colours and an increase in both the duration and frequency of hyperactive behaviours and such effects were absent upon Placebo exposure.</td>
</tr>
<tr>
<td>Swanson and Kinsbourne 1980</td>
<td>40 children (20 hyperactive responsive to stimulant mediation and 20 control average 10 years of age) given a diet free of synthetic food dyes and other additives for 5 days and on day 4 and 5 oral challenge at 10.00 am with 100 or 150 mg of a blend of nine food dyes or Placebo</td>
<td>Performance of the hyperactive children on paired-associate learning tests on the day they received the dye blend was impaired relative to their performance after they received the Placebo. The performance in the nonhyperactive group was not affected by the challenge with the food dye blend.</td>
</tr>
</tbody>
</table>
Assessment of the results of the study by McCann et al. (2007) on the effect of some colours and sodium benzoate on children’s behaviour

<table>
<thead>
<tr>
<th>Reference</th>
<th>Numbers of subjects, additive tested and dose levels</th>
<th>Outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td>Weiss et al. 1980</td>
<td>22 Young children, maintained on a diet that excluded certain foods, were challenged intermittently with a blend of seven synthetic colours in a double-blind trial. Parents’ observations provided the criteria of response.</td>
<td>One child responded mildly to the challenge and one, a 34-month-old female, responded dramatically.</td>
</tr>
<tr>
<td>Mattes JM, Gittelman R. 1981</td>
<td>11 Children on Feingold diet and responsive to colours in double blind cross over with order randomized challenged with food colouring at 13 mg mixture of all FDA approved synthetic food colours in proportions to reflect normal consumption patterns) and Placebo. Evaluations by parents, teachers and psychiatrists and psychological testing</td>
<td>No evidence of a food colouring effect</td>
</tr>
<tr>
<td>Kavale and Forness 1983</td>
<td>Meta analysis integrating findings from 23 studies testing the Feingold hypothesis.</td>
<td>The primary findings indicate that diet modification is not an effective intervention for hyperactivity.</td>
</tr>
<tr>
<td>Egger et al., 1985</td>
<td>76 Selected overactive children were treated with an oligoantigenic diet. 28 Children who improved their behaviour on the diet completed a double-blind crossover Placebo-controlled trial in which foods thought to provoke symptoms were reintroduced.</td>
<td>62 Children improved on the oligoantigenic diet and a normal behaviour was achieved in 21 of these children. In the cross over study, symptoms returned or were exacerbated much more often when children were on active material than on Placebo based diet. Synthetic colours and preservatives were the most common provoking substances but no child was sensitive to these alone.</td>
</tr>
<tr>
<td>David, 1987</td>
<td>Double blind challenges with 50 mg or 250 mg Tartrazine or benzoic acid in 24 children (1.6 to 12.4 years) with history of behavioural adverse effects to these colours in clinical settings</td>
<td>In no child was any change in behaviour noted by the parents or the nursing staff after administration of Placebo or test substance.</td>
</tr>
<tr>
<td>Gross et al., 1987</td>
<td>39 Children were given the Feingold diet for 1 week followed by administration for 1 week of food containing the synthetic additives and salicylates. All children were classified by public school psychologists as having moderate to severe learning disorders; 18 were also hyperkinetic and 17 were taking medication for motor restlessness. Three raters blind to the respective diets rated the children’s behaviour (monitored by video taping for 4 minute intervals at mealtime) for motor restlessness, disorganized behaviour and misbehaviour.</td>
<td>No significant differences were found in behaviours during week 1 and 2. The authors conclude that the Feingold diet has no beneficial effect.</td>
</tr>
<tr>
<td>Rowe K.S., 1988</td>
<td>From 55 children who participated in a 6-week open trial of the Feingold diet, 8 of 14 suspected reactors were involved in a double blind Placebo controlled repeated measures study in which 50 mg doses of the azo dyes Tartrazine and Carmoisine were used.</td>
<td>For 2 children there was a clear association between the ingestion of both dyes and behavioural symptoms of irritation, restlessness and sleep disturbance.</td>
</tr>
</tbody>
</table>
### Reference Numbers of subjects, additive tested and dose levels

<table>
<thead>
<tr>
<th>Reference</th>
<th>Numbers of subjects, additive tested and dose levels</th>
<th>Outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rowe K.S., 1988</td>
<td>Of 222 children referred for suspected hyperactivity, 55 were subjected to a 6 week trial of the Feingold diet. A double-blind crossover study, employing a single-subject repeated measures design was conducted using 8 of 14 children for which parents claimed that that a particular cluster of behaviours was associated with ingestion of foods containing synthetic colours. Subjects were maintained on a diet free from synthetic additives and were challenged daily for 18 weeks with Placebo or 50 mg of either tartrazine or Carmoisine each for 2 separate weeks.</td>
<td>Forty (72.7%) demonstrated improved behaviour and 26 (47.3%) remained improved following liberalization of the diet over a period of 3-6 months. In the double-blind crossover study two significant reactors were identified whose behavioural pattern featured extreme irritability, restlessness and sleep disturbance.</td>
</tr>
<tr>
<td>Kaplan et al., 1989</td>
<td>24 Hyperactive preschool-aged boys Within subject cross-over design (3 weeks baseline, 3 weeks Placebo, 4 weeks experimental diet, low in simple sugars, and eliminating synthetic colours and flavours, chocolate, monosodium glutamate, preservatives, caffeine and any substances the family reported might affect the specific child (i.e. dairy). According to parental reports (ten-item version of the Conners rating Scale, known as the Abbreviated Symptom Questionnaire (ASQ) asking about restlessness, impulsivity, disturbing other children, short attention span, fidgeting, distractibility, frustration, crying, mood changes and temper outbursts) more than half of the subjects exhibited a reliable improvement in behaviour and negligible Placebo effects. Several non-behavioural variables also tended to improve (halitosis, night awakenings, and latency to sleep onset).</td>
<td></td>
</tr>
<tr>
<td>Pollock and Warner, 1990</td>
<td>39 Children whose behaviour was observed by their parents to improve on a synthetic food additive free diet were included in a double-blind Placebo-controlled challenge. 19 Children completed the study. Synthetic food colours included in the study were 50 mg tartrazine, 25 mg sunset yellow, 25 mg Carmoisine, and 15 mg amaranth, all given in one capsule at breakfast. In these 19 children who completed the study food colours were shown to have an adverse effect on a daily Conners' rating of behaviour, but most parents could not detect these changes.</td>
<td></td>
</tr>
<tr>
<td>Carter et al., 1993</td>
<td>78 Children in clinical trial because of hyperactive behaviour placed on a few food items elimination diet. 19 of them (the ones for which foods or additives were disguised that reliably provoked behaviour) in a subsequent Placebo controlled double blind challenge protocol. 59 Children improved in behaviour. Crossover trial on the 19 children showed a significant effect for the provoking food to worsen ratings (by parents and other people with a role in child’s care) of behaviour and to impair psychological test performance</td>
<td></td>
</tr>
<tr>
<td>Rowe and Rowe, 1994</td>
<td>200 Children assessed for suspected hyperactivity. For the main study 50 reactors plus 34 other children (23 suspected reactors, 11 uncertain reactors) and 20 control subjects aged 2 to 14 years were studied in a 21 day double blind Placebo-controlled repeated measures study in which each child was used as its own control. Placebo or one of six dose levels of Tartrazine (1,2,5,10,20,50 mg) was administered randomly each morning and behavioural ratings were recorded by parents at the end of each 24 h. Parents of 150 children reported behavioural improvement with the diet and deterioration on the introduction of foods containing synthetic colouring. 24 children were clear reactors. Significant reactions at all dose levels and a dose response was obtained. It was concluded that behavioural changes in irritability, restlessness and sleep disturbance are associated with the ingestion of Tartrazine in some children.</td>
<td></td>
</tr>
</tbody>
</table>

*The EFSA Journal (2008) 660, 28-54*
<table>
<thead>
<tr>
<th>Reference</th>
<th>Numbers of subjects, additive tested and dose levels</th>
<th>Outcome</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Boris and Mandel, 1994</td>
<td>26 Children who met the criteria for ADHD were treated with a multiple item elimination diet and challenged with several foods, dyes and/or preservatives. A double blind Placebo controlled food challenge was completed in 16 of the 19 children responding favorably.</td>
<td>19 Children responded favorably. On open challenge all 19 children reacted to many foods, dyes and/or preservatives. In the double blind Placebo controlled study there was a significant improvement on Placebo days compared with challenge days. Atopic children with ADHD had a significantly higher response rate than the non-atopic group. It is concluded that the study shows a beneficial effect of eliminating reactive foods and synthetic colours in children with ADHD.</td>
<td></td>
</tr>
<tr>
<td>Schab and Trinh, 2004</td>
<td>Meta analysis: ten electronic databases were searched for double-blind Placebo controlled trials evaluating the effect of synthetic food colours. 15 Trials met the primary inclusion criteria.</td>
<td>Meta-analytic modelling determined the overall effect size of synthetic food colours on hyperactivity to be 0.283 (95% CI 0.079 to 0.488), falling to 0.210 (95% CA 0.007 to 0.414) when the smallest and lowest quality trials were excluded. Trials selected for responsiveness before enrolment demonstrated the greatest effects. The authors concluded that despite indications of publication bias and other limitations, this study is consistent with accumulating evidence that neurobehavioural toxicity may characterise a variety of widely distributed chemicals.</td>
<td></td>
</tr>
<tr>
<td>Bateman et al., 2004</td>
<td>Children (3-years old) screened for hyperactivity (HA) and atopy (AT). Four groups selected (HA/AT, not HA/AT, HA/not-AT and not-HA/not-AT (n=277) and subjected to a diet eliminating synthetic colours and benzoate preservatives for one week. In subsequent 3 weeks within subject double blind crossover study with, in random order dietary challenge with a drink containing synthetic colourings (20 mg/day)(Sunset Yellow, Tartrazine, Carmoisine, Ponceau 4R; 5 mg of each) and sodium benzoate (45 mg/day)(active period), or a Placebo mixture. Behaviour was assessed by a tester blind to dietary status and by parent’s ratings</td>
<td>Significant reductions in hyperactive behaviour during the withdrawing phase and significantly greater increases in hyperactive behaviour during the active than the Placebo period based on parental reports. Effects were not influenced by presence or absence of HA or AT. No significant differences detected based on testing in the clinic.</td>
<td></td>
</tr>
</tbody>
</table>

Initial studies, as summarized in Table 9, investigated the effects on behaviour of Feingold’s diet (a diet without synthetic colours and flavours) under double-blind conditions in relatively small groups of hyperactive children (Conners et al., 1976, Williams et al., 1978; Kaplan et al. 1989; Harley et al., 1978a). Several studies reported improved behavioural characteristics in part but not all children of their study population (Conners et al., 1976, Williams et al., 1978; Kaplan et al. 1989) whereas others did not provide support for the Feingold hypothesis (Harley et al., 1978a). In other studies children identified as possible responders were challenged in double-blind studies (Harley et al., 1978b; Goyette et al., 1978; Mattes et al., 1978; Levy et al., 1978; Weiss et al., 1980). These studies used generally only hyperactive and responsive children, small study groups and parental or teaching ratings, and reported either adverse or no effects. Generally all these studies did not study dose-response dependency and also did not link the adverse behavioural effect to a specific food additive.
The Panel noted that the findings from individual studies have in general not been conclusive. This has been ascribed by others to several logistic and methodological problems (Rowe and Rowe 1994 and references therein), including for example:

- the identification of children from heterogeneous populations;
- problems with dietary compliance;
- Placebo effects;
- the possible lack of inertness of the control substance;
- varying and imprecise diagnostic criteria for hyperactivity;
- doubts about the validity and reliability of behavioural outcome measures particularly those appropriate to the assessment of dye challenge effects;
- uncertainty about the detection of treatment effects when only a small number of children respond;
- confusion about suitable dosage levels of colourings for use in challenge trials, and
- lack of incorporating different dosages into the design.

The Panel also notes that a few controlled studies on the effects of ingested synthetic food colours on behaviour (Rose, 1978; Swanson, 1980; Mattes and Gittelman, 1981; Egger et al., 1985; David, 1987; Rowe, 1988) have reported inconsistent effects on behaviour (changes or no change) after the dye challenge as compared to Placebo.

Only a few studies deal with the possible behavioural effects of specific, individual, food colours. A dose response curve was reported in a study with responsive children exposed to increasing doses of Tartrazine, and this study again revealed that some but not all of the children in a selected responsive study population were responders (Rowe and Rowe, 1994).

A more recent meta-analysis of double blind Placebo-controlled trials combined fifteen individual trials that met the inclusion criteria (Schab and Trinh, 2004). Meta-analytic modelling determined the overall effect size of the artificial food colours on the hyperactivity score to be 0.283 (CI 95%, 0.079 to 0.488), falling to 0.210 (CI 95%, 0.007 to 0.414) when the smallest and lowest quality trials were excluded. Trials that screened for responsiveness before enrolment demonstrated the greatest effects. The authors concluded that despite indications of publication bias and other limitations, this study is consistent with accumulating evidence that neurobehavioural toxicity may characterise a variety of widely distributed chemicals.

On the other hand the older meta-analysis performed by Kavale and Forness (Kavale and Forness 1983) integrating findings from 23 studies testing the Feingold hypothesis concluded that diet modification is not an effective intervention for hyperactivity.

In addition to human behavioural studies, animal studies reporting effects of food colours on neurological and behavioural parameters have been reported (Vorhees et al., 1983; Tanaka, 1994; Tanaka, 1996, Tanaka, 2006a; Tanaka, 2006b). However, since the relationship between these experimental parameters in animal studies and ADHD-like symptoms in humans remain unclear the present opinion does not take these animal data into account. Novel behavioural methods have been developed measuring the key ADHD behaviours in children and animal models of ADHD (e.g. Sagvolden, 2000, Sagvolden et al., 2005) and it could be possible to use such methods to evaluate behavioural effects of doses of various synthetic colours and flavours in normal animals as well as animal models of ADHD in order to elucidate possible mechanisms and relations presently lacking in available studies of children.
In summary, the Panel considers that while a number of studies have reported a possible relationship between exposure of young children to synthetic food colours and other food additives and behavioural effects others have not identified an association between exposure to these substances and behavioural effects. The available literature is thus not consistent and does not allow a firm conclusion. The Panel notes that the majority of these studies have been conducted on children described as hyperactive or with a clinical diagnosis of ADHD, thus not being representative of the general population.

5. Possible mechanisms of action

One of the explanations for the suggested induction of neurobehavioural (ADHD-like) responses in children following exposure to synthetic food colours and other food additives, is a CNS-mediated origin. It is assumed that ADHD is linked to altered Central Nervous System (CNS) dopamine function, most likely mediated by faulty dopaminergic modulation of neuronal activity transmitted by the neurotransmitters glutamate and GABA (Sagvolden et al., 2005).

These responses may have a genetic basis. ADHD is not a uniform disorder and it is evident that there are subgroups of affected children responding to different triggers with behaviour classified as ADHD. Genetic factors (Shoukri & Donner, 2007), such as serotonin receptor polymorphisms (Brookes et al., 2006), dopamine receptor (Mill et al., 2004) and other receptor polymorphisms may play a role (Polanczyk et al., 2007). Some of these observations have not been confirmed and may be population-specific findings (Curran et al. 2001).

The findings of the present study also suggest that certain genetic polymorphisms, specifically two genetic polymorphisms thought to impair histamine clearance, may result in possible differential sensitivity to the particular additives used in the study, although the increases in GHA scores were not limited to individuals with the specific polymorphisms measured in the study (COT, 2007). There were no associations between behaviour and other genetic polymorphisms investigated in the study, including genetic polymorphisms selected from the dopamine neurotransmitter systems, which have previously been implicated in ADHD (COT, 2007).

Another possible mechanism is a hypersensitivity reaction in a small, sensitive subgroup of the population. ADHD sufferers can react negatively to “allergenic” foods such as milk, egg, wheat etc. (Carter et al., 1993; Boris & Mandel, 1994) and a connection with food allergies has been suggested (Marshall, 1989).

DISCUSSION

The study by McCann et al. (2007) reports effects of two combinations of Tartrazine (E102), Quinoline Yellow (E104), Sunset Yellow FCF (E110), Ponceau 4R (E124), Allura Red AC (E129), Carmoisine (Azorubine, E122) and sodium benzoate (E211) on children’s behaviour, as measured by the Global Hyperactivity Aggregate (GHA) score, a novel metric developed by the researchers, combining behavioural and computer-based measures in one overall parameter. The Panel notes that the children who were included in the study were selected to represent a broad range of behaviour in the general population including children with normal activity through to those with high activity levels, but that children who were medicated for ADHD were not included.

A small but significant effect of Mix A on GHA scores were observed in 3-year old children (effect size 0.20 [95% CI 0.01 to 0.40], p<0.05), while Mix B did not produce a significant
change in the GHA scores for this group. In contrast, in 8- to 9-year old children, a significant effect of Mix A (effect size 0.14 [95% CI 0.03 to 0.24], p<0.05) and Mix B (effect size 0.17 [95% CI 0.06 to 0.28], p<0.01) was observed, but only when analysis was restricted to those children consuming at least 85% of drinks with no missing data. When all children that completed the study were taken into account in 8- to 9-year old children only Mix B had a significant effect on GHA scores (effect size 0.12 [95% CI 0.03 to 0.22] p<0.05), while Mix A had no significant effect. Thus, the statistically significant effects were not found for the same mixture in the two age groups. Overall the increases in the GHA scores observed in the study were small, ranging from 12 to 20% increase in GHA scores for the entire sample.

A statistical reanalysis of the data from the Southampton study was undertaken by the Panel. All individual behaviour variables (minus the individual baseline value for that variable) were reanalysed without normalisation, so that each subject served as its own control. This reanalysis was undertaken both at the level of the individual behavioural variables as well as on the aggregated scores.

For the primary analysis the Global Hyperactivity Aggregate (GHA) score was recalculated following the same steps as in the original analysis, except for the omission of the final re-normalisation step.

Based on these results it is concluded that the primary analysis with the recalculated GHA score led to broadly similar conclusions to that in the original paper by McCann et al, except for the following:

1. The Mix A versus Placebo comparison was not statistically significant for the three year olds when all subjects were included (entire sample), while the significance for the ≥ 85% consumption and complete case groups was increased slightly;

2. for the 8- to 9-years age group, the Mix A versus Placebo comparison was no longer statistically significant in any of the three consumption groups.

In addition the data were analysed on the basis of a modified GHA score in which the parental scores were not included. The results from this analysis did no longer reveal any statistically significant effects of Mix A or Mix B versus Placebo, except for Mix B in the 8- to 9-year old completers when both sexes are pooled (p=0.042).

A further analysis was carried out on the whole data set, comprising analysis of the single variables of parental scores, teacher scores and observer scores, and, in the case of 8- to 9-year old children, computer-based scores. There is a suggestion from these analyses that the statistically significant effects seen in the 3-year olds (Mix A versus Placebo) and in the 8-to 9-year olds (Mix B versus Placebo) are largely driven in the data by the parental scores and, in the older males in both comparisons, by the computer score.

The Panel noted that the main contributors to the GHA scores were the parental scores. The scores from teachers and independent observers showed little positive trend and were not a major component in the overall GHA scores. The use of the GHA metric does not therefore completely overcome the criticisms of the earlier Isle of Wight study (Bateman et al. 2004).

The Panel thus concludes that the McCann et al. study (2007) provides limited evidence that the two different mixtures of synthetic colours plus sodium benzoate tested had a small but statistically significant effect on behaviour in children from the general population excluding children medicated for ADHD, as measured by an aggregated score of behavioural effects.

The Panel considers that the clinical significance of the observed effects (a) for the individual children in the study and (b) for the population as a whole remains unclear, since the effects
were small in magnitude and it is not known whether these small changes in attention and activity would interfere with schoolwork and other intellectual functioning.

The conclusions of McCann *et al.* were restricted to the hypothesis that some synthetic colours or sodium benzoate (or both) in the diet resulted in increased activity scores in children and did not implicate these agents as causative agents in ADHD. The Panel agrees that an “elevated score of activity / inattention” is by no means equivalent or even indicative of ADHD. The clinical diagnosis of this condition requires impaired social and behavioural functioning and not merely an “elevated score of activity / inattention”. Furthermore it is important to stress that ADHD is a condition with a multifactorial aetiology and exclusive focus on food additives may detract from the provision of adequate treatment for children with ADHD.

The Panel noted that some, but not all, earlier studies have also reported effects of certain food colours on child behaviour, the majority of these studies being conducted on children described as hyperactive or with a clinical diagnosis of ADHD.

A recent meta-analysis of double blind Placebo-controlled trials combined fifteen individual trials that met the inclusion criteria (Schab and Trinh, 2004). Meta-analytic modelling determined the overall effect size of the artificial food colours on the hyperactivity score to be 0.283 (CI 95%, 0.079 to 0.488), falling to 0.210 (CI 95%, 0.007 to 0.414) when the smallest and lowest quality trials were excluded. Trials that screened for responsiveness before enrolment demonstrated the greatest effects. The authors concluded that despite indications of publication bias and other limitations, this study is consistent with accumulating evidence that neurobehavioural toxicity may characterise a variety of widely distributed chemicals.

On the other hand the older meta-analysis performed by Kavale and Forness (1983) integrating findings from 23 studies testing the Feingold hypothesis concluded that diet modification is not an effective intervention for hyperactivity.

In the available studies, changes in behaviour, from either addition or withdrawal of additives from the diet, were not observed in all children, suggesting there may be a subpopulation of individuals who are sensitive to food additives in general or to food colours in particular. The findings of the present study suggest that certain genetic polymorphisms, specifically two genetic polymorphisms thought to impair histamine clearance, may result in possible differential sensitivity to the particular additives used in the study. The increases in GHA scores were however not limited to individuals with the specific polymorphisms measured in the study, and there were no associations between behaviour and other genetic polymorphisms investigated in the study (COT, 2007). The observed associations between polymorphisms in the histamine N-methyltransferase gene and the difference in behaviour with Mix A in 3-year olds and Mix A and Mix B in 8- to 9-year olds compared to Placebo, even if real and not merely chance effects, were not sufficiently strong that they could usefully be applied to identify at-risk groups or individuals (COT, 2007).

If a sensitive subpopulation does exist, it is not possible, from the currently available data, to assess the overall prevalence of such sensitivity and whether particular food additives may be implicated.

Based on surveys conducted from 2002 to 2005, the target colours are more frequently used in sweets but also occur commonly in soft drinks and benzoate is frequently present in soft drinks. Children consuming brightly coloured sweets may be exposed to levels comparable to those considered in the protocol of the McCann *et al.* study for one or more of the food colours studied. Comparable levels may also be reached in those children who consume brightly coloured soft drinks. The level of exposure to sodium benzoate is also likely to occur.
There are a number of uncertainties and limitations that are apparent from this new research, some of which are echoed from earlier research. These include:

- the limited consistency of the results with respect to age and gender of the children, the effects of the two mixtures of additives tested and the type of observer (parent, teacher or independent observer);
- the unknown clinical relevance of the novel metric, i.e. the GHA score;
- the unknown relevance of the small effect size (as was also seen in the meta analysis of earlier studies by Schab and Trinh (2004);
- the fact that the study has not been designed to identify the effects of individual additives;
- a lack of information on dose-response;
- the lack of a biologically plausible mechanism for induction of behavioural effects from consumption of food additives.

In the context of the overall weight of evidence, the Panel considers that the findings from the McCann et al. study are not sufficiently conclusive to be used as a basis for altering the ADI of the respective food colours or sodium benzoate.

CONCLUSIONS
The Panel concludes that the McCann et al. study provides limited evidence that the two different mixtures of synthetic colours and sodium benzoate tested had a small and statistically significant effect on activity and attention in children selected from the general population excluding children medicated for ADHD, although the effects were not statistically significant for the two mixtures in both age groups.

Since mixtures and not individual additives were tested in the study by McCann et al., it is not possible to ascribe the observed effects to any of the individual compounds.

The clinical significance of the observed effects also remains unclear, since it is not known whether these small alterations in attention and activity would interfere with schoolwork and other intellectual functioning.

In the context of the overall weight of evidence and in view of the considerable uncertainties, such as the lack of consistency and relative weakness of the effect and the absence of information on the clinical significance of the behavioural changes observed, the Panel concludes that the findings of the study cannot be used as a basis for altering the ADI of the respective food colours or sodium benzoate.
Assessment of the results of the study by McCann et al. (2007) on the effect of some colours and sodium benzoate on children’s behaviour

**DOCUMENTATION PROVIDED TO EFSA**

1. Study details provided by Prof. J. Stevenson and the Food Standards Agency.

**REFERENCES**


COT Committee on toxicity, 2007. Statement on research project (T07040) investigating the effect of mixtures of certain food colours and a preservative on behaviour in children. [http://www.food.gov.uk/multimedia/pdfs/committee/colpreschil.pdf](http://www.food.gov.uk/multimedia/pdfs/committee/colpreschil.pdf)

Assessment of the results of the study by McCann et al. (2007) on the effect of some colours and sodium benzoate on children’s behaviour


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APPENDICES

APPENDIX A

Details on Kinetics and metabolism data for substances investigated for neurobehavioural effects in a study by McCann et al., Lancet. September 2007.

TARTRAZINE

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Chemical name: Trisodium-5-hydroxy-1-(sulfonatophenyl)-4-(4-sulphonatophenylazo)-H-pyrazole-3-carboxylate

Chemical formula: C₁₆H₉N₄Na₃O₉S₂

Structural formula:

\[
\text{N} \quad \text{N} \quad \text{N} \\
\text{O} \quad \text{O} \\
\text{O} \quad \text{O} \\
\text{S} \quad \text{O} \\
\text{O} \quad \text{O} \\
+ 3 \text{Na}^+ 
\]

The JECFA evaluation (1966) describes several studies which have focussed on the toxicokinetic aspects of tartrazine. No further new literature has been published since except for some studies describing azoreduction by intestinal bacteria.

After intravenous injection of tartrazine in rats an average 1% of the dye was recovered from the bile. This low biliary excretion is believed to be associated with a free carboxyl group in the 3-position of the pyrazolone ring. After intraperitoneal administration of the colour a conjugated form of tartrazine was rapidly excreted in the urine. In both bile and urine no reduction products were detected. Based on these results the authors stated that tartrazine is a substituted phenylhydrazone rather than a true azo-compound (Ryan and Wright, 1961, 1962).

Rats were given tartrazine intraperitoneally and afterwards the urine was examined. The urine contained only the unchanged colour; conjugates or reduction products such as amines were absent. Rabbits were also given tartrazine by intraperitoneal injection and urine was examined. No details on the outcome of the study were however given.

After oral administration of tartrazine to rats, rabbits, and humans, sulfanilic acid (presumably the N-acetate) was found in the urine. As tartrazine was only reduced after oral administration it appeared that reduction was carried out by gastro-intestinal flora. Nonetheless, the authors concluded in favour of an absence of a true azo-linkage and mention that physical methods demonstrated tartrazine to be a keto-hydrazone tautomer (Wright, 1963; Jones et al., 1964).
Rabbits were fed tartrazine and subsequently a 48 hours urine sample was analyzed. Besides the unchanged colour (1%) the metabolites sulfanilic acid (74%), and p-acetamido-benzenesulfonic acid (22%; the N-acetyl conjugate of sulfanilic acid) were identified in the urine sample. The percentages are the fraction of the maximal theoretical amount, which indicate that tartrazine is virtually completely reduced in the azo bond (Daniel, 1962).
Assessment of the results of the study by McCann et al. (2007) on the effect of some colours and sodium benzoate on children’s behaviour

QUINOLINE YELLOW

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Chemical name: disodium 2-(2-quinolyl)indane-1,3-dionedisulfonate (principle component)

Chemical formula: C₁₈H₉NNa₂O₈S₂ (principle component)

Structural formula:

\[
\begin{align*}
\text{N} & \quad \text{O} \\
\text{O} & \quad (\text{NaSO}_3)_{1-3}
\end{align*}
\]

In the JECFA evaluation of 1984 the subject of toxicokinetics is addressed for the first time.

Blood levels of radioactivity were measured in male and female rats after intra-gastric doses of 1 mg/kg bw \(^{14}\)C-Quinoline Yellow. It appeared that the peak of radioactivity occurred between 0.5 and 1 hr. after dosing. All radioactivity was found in the plasma at a maximum concentration of 0.009% of the dose (no detail on how to interpret this percentage) and most of the radioactivity was bound to plasma proteins. No metabolites were found in the plasma. The kinetics of the blood levels fitted a two-compartment model with the following parameters: \(T_{1/2} = 0.6\) hour; \(T_{11/2} = 12\) hours; and \(T_{21/2} = 70\) hours. In a complementary study carcasses of male rats were dissected and residual tissue levels determined corresponding to 1/2, 1, 4, 8, 24 and 48 hours after dosing. The results confirmed that activity was selectively concentrated in the thyroid (LEMM, 1978).

Male rats received a single intra-gastric dose of 4 mg \(^{14}\)C-Quinoline Yellow. About 94% of the radioactivity was recovered in the faeces (within 120 hours) and about 2% was eliminated in the urine. Retention was approximately 0.14%. The compound was found to be metabolized to only a small extent. In the urine 10-15% of the activity was associated with an unidentified metabolite. After 120 hours males were sacrificed and residual tissue levels were determined. The activity was selectively concentrated in the thyroid (no quantitative details) (LEMM, 1978).

In rats dosed with 2.85 mg/kg bw \(^{14}\)C-Quinoline Yellow, after 31.5 hours only 1% of the dose was found to be excreted through the biliary route. No metabolites were found in the bile (LEMM, 1978).

After administering Quinoline Yellow to male rats whole body autoradiography demonstrated that after 1 hr. the activity was primarily associated with the gastro-intestinal tract and excretory organs. After 24 hours only the large intestine and, to a minor degree, the cortical zone of the kidney displayed activity (LEMM, 1978).

Tissue distribution studies after intra-gastric exposure of female rats to \(^{14}\)C-Quinoline Yellow showed that the small proportion of the dose that was absorbed from the gastro-intestinal tract (estimated 3-4%) was primarily associated with the liver (max. 1 %), kidney (max 0.02%), and
Bladder. Results expressed as concentration factors (radioactivity/g tissue) showed that a selective concentration of the thyroid persisted up to 48 hours, and a relatively high concentration was found in the ovaries in the first 24 hours (LEMM, 1978).

In dogs blood levels and excretion after intra-venous and intra-gastric administration of $^{14}$C-Quinoline Yellow (0.2 and 0.44 mg/kg bw respectively) were examined. After intra-venous administration the disappearance of radioactivity corresponded to a two-compartment pharmaco-kinetic model with $T_1/2 = 4$ hours and $T_2/2 = 43$ hours. About 22% of the dose was excreted in the faeces. Intra-gastric administration showed that peak blood levels occurred at 1-4 hours after dosing. From 0-72 hours the urine contained 1-4% of the radio-label, 42-60% was excreted in the faeces within 72 hours. After both routes of administration there was no indication of specific tissue accumulation, particularly in the thyroid. Examination of urine, faeces, and plasma indicated that Quinoline Yellow is metabolized to only a small extent (LEMM, 1978).
Assessment of the results of the study by McCann et al. (2007) on the effect of some colours and sodium benzoate on children’s behaviour

ALLURA RED AC

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Chemical name: Disodium 2-hydroxy-1-(2-methoxy-5-methyl-4-sulfonatophenylazo)naphthalene-6-sulfonate

Chemical formula: C\textsubscript{18}H\textsubscript{14}N\textsubscript{2}Na\textsubscript{2}O\textsubscript{8}S\textsubscript{2}

Structural formula:

\[
\begin{align*}
\text{N} & \quad \text{O} \\
\text{CH}_3 & \quad \text{CH}_3 \\
\text{S} & \quad \text{O} \\
\text{O}^- & \quad +2 \text{Na}
\end{align*}
\]

Three studies were reviewed by the JECFA (1980) to provide insight in the metabolic aspects of Allura Red AC.

Rats were fed a diet containing 5.19% of Allura Red AC (White, 1970). It was observed that 0.1% and 29% of the intact dye was excreted in the urine and faeces respectively. It was postulated that azo reduction by gut flora of the dye will yield the two components of the parent compound: 2-methoxy-5-methyl-aniline-4-sulfonic acid (cresidine-4-sulfonic acid), and 1-amino-2-naphthol-6-sulfonic acid (White, 1970)

In later studies, rats and dogs were pretreated daily with non-radioactive Allura Red AC. Subsequently, the animals were dosed with the \textsuperscript{35}S-labelled compound and studied for up to 72 hours for excretion and distribution patterns of the colour. Both species showed limited absorption of the compound with the major route of excretion being via the faeces. In the dog 92 to 95% of the recovered radioactivity appeared in the faeces within 72 hours while in the rat 76 to 92% of the recovered radioactivity appeared in the faeces within this time period. Urinary recoveries of the colour varied between 5.7 and 19.8% and 2.7 and 3.6% in rats and dogs, respectively. After sacrifice, significant retention of radioactivity was located in the intestinal contents of both species and in the washed intestines of the rats. This was thought to be due to adhesion of the compound to the intestinal wall, since the total carcass and viscera of these animals contained less than 0.4% of the administered dose (Guyton & Reno, 1975). Cresidinesulfonic acid was found to be the major metabolite of Allura Red AC in the urine of these two species, whereas the parent compound was not measurable. In addition, two other unidentifiable metabolites were found in the urine of the rats. In the rat faecal extracts, cresidinesulfonic acid was a major metabolite along with two unknowns and the parent compound. The dog faecal sample revealed an identical metabolite pattern as seen in the rat, and in addition, a third unknown was discovered. One of the urinary unknowns demonstrated an \textit{R}_f value which was identical to that of the one of the faecal unknowns suggesting that they were one and the same. The other unknowns exhibited distinctive \textit{R}_f values which indicated that these metabolites were different (Guyton & Stanovick, 1975).
Assessment of the results of the study by McCann et al. (2007) on the effect of some colours and sodium benzoate on children’s behaviour

PONCEAU 4R

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**Chemical name:** Trisodium 2-hydroxy-1-(4-Sulphonato-1-naphthylazo)-naphtalene-6,8-disulphonate.

**Chemical formula:** C_{20}H_{11}N_{2}Na_{3}O_{10}S_{3}

**Structural formula:**

![Structural formula of Ponceau 4R]

The JECFA evaluation of 1983 gives a brief understanding of the biochemical fate of Ponceau 4R.

Single oral dose studies of uniformly $^{14}$C-labeled Ponceau 4R of 0.5 or 50 mg/kg bw in rats, mice, and guinea-pigs show that substantially all of an orally administered dose of Ponceau 4R related material (e.g. $^{14}$C-label) is excreted in the urine, bile and faeces, with the majority being accounted for in the faeces (90%; 25-35% unchanged); metabolites are found in the urine (mainly naphthionic acid) and faeces (naphthionic acid and 7-hydroxy-8-aminonaphtalene-1,3-disulfonic acid); and finally, apart from some retention in foetuses, there is no marked accumulation in any tissue. Only some Ponceau 4R was absorbed by isolated intestinal loops (Phillips et al., 1982).

In a study in which rats received an intravenous dose of Ponceau 4R, 30-45% of the dye was excreted unchanged in the bile within six hours (Ryan and Wright, 1961).

Furthermore, it was found that after intraperitoneal administration of the dye the bile was coloured in mice and rats (Gaunt et al., 1967).

Finally, a study by Walker (1968) indicates that Ponceau 4R is reduced *in vitro* by rat caecal contents.
Assessment of the results of the study by McCann et al. (2007) on the effect of some colours and sodium benzoate on children’s behaviour

**SUNSET YELLOW FCF**

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**Chemical name:** Disodium 2-hydroxy-1-(4-sulfonatophenylazo)naphthalene-6-sulfonate  

**Chemical formula:** C\textsubscript{16}H\textsubscript{10}N\textsubscript{2}Na\textsubscript{2}O\textsubscript{7}S\textsubscript{2}  

**Structural formula:**

\[
\text{N} \quad \text{O} \\
\text{O} \quad \text{HO} \\
\text{O} \quad \text{S} \\
\text{O} \quad \text{O} \\
\text{N} \quad + 2 \text{Na}^+ \\
\text{O} \quad \text{S} \\
\text{O} \quad \text{O}
\]

The JECFA (1982) reports of five studies on the toxicokinetic aspects of Sunset Yellow FCF. In a study in rats given a single oral dose of Sunset Yellow FCF 0.8% of the administered dose was recovered from the faeces as intact colour. In bile and urine these percentages were 3 and 0.8 respectively. In the urine of rats given large oral doses of Sunset Yellow FCF, the azo-reduction products sulfanilic acid and 1-amino-2-naphtol-6-sulfonic acid were found. No qualitative or quantitative measurement of reduction products in the faeces was carried out. From these results, in combination with observations after intravenous and intrasplenic administration, the authors concluded that breakdown of Sunset Yellow FCF to (sulfonated) aromatic amines is due to reduction by intestinal bacteria rather than by liver enzymes (Radomski and Mellinger, 1962).

Rats that received Sunset Yellow FCF by gavage excreted 0.3% as intact colour and 37% as sulfanilic acid in the urine. In the bile 1.5% was excreted as intact colour (sulfanilic acid not measured). In the same study animals were gavage dosed with \(^{14}\text{C}\)-Sunset Yellow FCF (labelled at the C-8 position of the naphthalene ring). As a result 94.5% of the total radioactivity was retrieved from the faeces, 8.5% from the urine. After the first 24 hours 1-2% of the total dose in urine consisted of intact dye and 40% of the dose consisted of the molar equivalents of sulfanilic acid of which 24% was \(N\)-acyetylated. The other almost 60% of the radioactivity in the urine is unaccounted for (Honohan et al., 1977).

After an intravenous injection of Sunset Yellow FCF in rats (no specification on dose) 20-30% of the dose was found in the bile after 6 hours (Ryan and Wright, 1961).

The urine of rabbits which were fed a single dose of Sunset Yellow FCF contained unchanged colour (2%), and the two azo-reduction products sulfanilic acid (54%), and 1-amino-2-naphtol-6-sulfonic acid (55% in 24 hours). In addition the \(N\)-acyetylated form of sulfanilic acid, \(p\)-acetamidobenzene-sulfonic acid, was present in the urine (23%) (percentages indicate the ratio of the amount of the metabolite found to the theoretical amount, assuming complete breakdown) (Daniel, 1962).
AZORUBINE (= CARMOISINE)

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**Chemical name:** Disodium 4-hydroxy-3- (4-sulfonato-1-naphthylazo) naphthalene-1-sulfonate

**Chemical formula:** C_{20}H_{12}N_{2}Na_{2}O_{7}S_{2}

**Structural formula:**

IJCFA (1983) describes several studies on the toxicokinetic aspects of Azorubine

Mice (CD-1) (3-6 males/group) received single doses of $^{14}$C-Azorubine (5 μCi/mmol) by gavage (200 mg/kg bw, 6 μCi) or intravenously (200 mg/kg bw, 0.7 μCi).

After oral administration, peak levels of radioactivity occurred in plasma (0.08%/ml) and in the liver, lung, testes and spleen 8 hours after dosing. Radioactivity was almost completely excreted in faeces (74%) and urine (19%) within 16-32 hours.

After intravenous administration, most of the radioactivity (76%) was excreted in faeces (64%) and urine (12 %) 24 hours after dosing. The plasma $^{14}$C-radioactivity decay curve indicated a very rapid distribution of the compound into the tissues ($t_{1/2} = 10$ minutes) and an efficient excretion mostly through the gastrointestinal tract (92%) which was complete 48 hours after dosing (Galli et al., 1981).

In a study in rats, animals ($\geq 3$ males/group) were given $^{14}$C-Azorubine by gavage (200 mg/kg; 25 μCi), or by intravenous injection (200 mg/kg; 3 μCi). Radioactivity was measured in blood, tissue, faeces and urine 5, 10 and 30 minutes and 1, 2, 4, 8, 16, 32, 64 and 96 hours after dosing.

After gavage, no radioactivity was detected in the brain, adipose tissue, muscle, testes, spleen or lung (no time specification). After 32 hours, recovery of the administered radioactivity was 82% in faeces and 8% in urine. The radioactivity profile of the blood indicated rapid but poor absorption (maximum radioactivity content (0.01%/ml) being reached within 10 minutes).

After intravenous injection, the blood $^{14}$C-radioactivity decay curve indicated rapid distribution to the tissues and could be described in terms of a two-compartment model. Radioactivity was highest in the gastrointestinal tract and liver (no time specification). However, within 24 hours after injection all radioactivity was recovered in faeces (79%) and urine (not specified). The large quantity present in the faeces was considered to indicate active excretion of Azorubine.
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and its metabolites in the bile and poor re-absorption from the intestine. Based on the blood-radioactivity curves after oral and intravenous administration, bioavailability of $^{14}$C-Azorubine was calculated to be less than 10% (Galli et al., 1982a).

Rats were given an intravenous injection with approximately 1 mg Azorubine. The 6 hours recovery of Azorubine in the bile was an average of 38% (30-40%) of the administered quantity (Ryan and Wright, 1961).

In another study, rats received 200 mg/kg bw $^{14}$C-Azorubine (25 µCi) by gavage after which radioactive compounds in faeces and urine were investigated.

In addition to unmodified Azorubine, five radioactive compounds were present of which the predominant one was identified as naphthionic acid. After anaerobic incubation of $^{14}$C-Azorubine with a bacterial suspension isolated from faeces from humans and rats, similar metabolic patterns were found (Morovich et al., 1983).

Wistar albino Rats (both sexes), MF-1 mice (male), and Dunkin-Hartley guinea-pigs (male) were administered a single oral dose of either 0.5 mg/kg or 50 mg/kg $^{14}$C-Azorubine (20 µCi/kg).

In the first 24 hours the majority of radioactivity was excreted in the urine and faeces (18% and 73% in rats, 17% and 66% in mice, and 37% and 45% in guinea-pigs respectively). After 72 hours, substantially all of the radioactivity was recovered in the excreta, the majority being accounted for in the faeces.

In the urine of all three species 60 - 80% of the radioactivity was associated with naphthionic acid. Further, 10 - 20% of the radioactivity was identified as 2-amino-1-naphthol-4- sulfonic acid (2-ANS). In rats and mice ≥5% and in guinea-pigs 16% of the radioactivity was identified as 1,2-naphthoquinone-4-sulfonate (1,2-NQS). A fourth metabolite, accounting for 2 - 5% of the radioactivity in the urine, was not identified.

In the faeces of all three species naphthionic acid was also found; however, no 2-ANS or 1,2-NQS was detected. In addition, five unidentified metabolites (two hydrolysable by combined β-glucuronidase and sulfatase treatment) were found of which the proportions varied between species. No significant absorption of radioactivity from isolated small intestinal loops was noted in all three species.

Less than 0.03% of the administered radioactivity (50 mg/kg single dose) was recovered in the bile during 1 hour and only 0.04 - 0.7% during 5 hours. Less than 0.03% of the dose was eliminated as CO$_2$ (Phillips et al., 1982).

Pregnant rats eliminated a single oral dose of $^{14}$C-Azorubine (50 mg/kg at day 8 of pregnancy) at a similar rate to non-pregnant females. The concentration of radioactivity in the foetuses was similar to that in the other tissues (no further detail).

Pre-treatment of male rats with unlabelled Azorubine in the diet for 28 days (approximately 50 mg/kg bw/d) prior to dosing with $^{14}$C-Azorubine (50 mg/kg bw), had no effect on the route of excretion or time of total elimination. The only difference compared to single dosed animals was that the proportion of metabolites extracted from the faeces differed (no further detail) (Phillips et al., 1982).

In order to study the formation of (sulfonated) aromatic amines, the anaerobic reduction of Azorubine was investigated by incubating Azorubine with caecal content and hepatic microsomal fraction of rats. Caecal suspension exhibited higher azo reductase activity than that of hepatic microsomal fraction. The researchers consider that the reductive ability through caecal flora signifies the formation of sulfonated aromatic amines which may be re-absorbed through the intestine to be
either eliminated through urine as conjugates or retained in the target tissues (Singh et al., 1997).

Pregnant rats received 200 mg $^{14}$C-Azorubine/kg bw by gavage on gestational days (GD) 16-19 and were sacrificed to analyse maternal tissues, amniotic fluid, placentas, foetal membranes and foetuses for radioactivity. Male rats were given a single oral dose of 200 mg $^{14}$C-Azorubine/kg bw and sacrificed at different times after dosing.

In animals of both sexes, over 90% of radioactivity was excreted in faeces and urine within 64 h. This suggested that absorption of Azorubine is limited and that no significant accumulation occurred in any particular tissue. Of 5 metabolites determined, the principle one was identified as naphthionic acid. There was no evidence of transplacental transfer of $^{14}$C-Azorubine or its metabolites. Results demonstrated that pregnancy does not affect the toxicokinetic profile of Azorubine (Tragni et al., 1985).
Assessment of the results of the study by McCann et al. (2007) on the effect of some colours and sodium benzoate on children’s behaviour

NA-BENZOATE / BENZOIC ACID

<table>
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<tr>
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<td>Chemical formula: C7H6O2 / NaC7H5O2</td>
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<td>Structural formula:</td>
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(TAKEN FROM WHO, 2000; only information on oral absorption is included)

After oral ingestion of benzoic acid and sodium benzoate, there is a rapid absorption (of undissociated benzoic acid) from the gastrointestinal tract in experimental animals or humans (US FDA, 1972a, 1973). From the figures on excretion given below, 100% absorption can be assumed. In humans, the peak plasma concentration is reached within 1-2 h (Kubota et al., 1988; Kubota & Ishizaki, 1991).

In vivo dermal studies with benzoic acid in experimental animals (e.g., guinea-pigs, mice, rats, pigs, dogs, rhesus monkeys) confirm the results with humans (Hunziker et al., 1978; Andersen et al., 1980; Wester & Noonan, 1980; Bronaugh et al., 1982a; Reifenrath et al., 1984; Carver & Riviere, 1989; Maibach & Wester, 1989; Bucks et al., 1990). Absorption ranged from 25% in pigs (Reifenrath et al., 1984; Carver & Riviere, 1989) to 89% in rhesus monkeys (Wester & Noonan, 1980; Maibach & Wester, 1989; Bucks et al., 1990).

After oral and dermal uptake, benzoate is metabolized in the liver by conjugation with glycine, resulting in the formation of hippuric acid (Feldmann & Maibach, 1970; US FDA, 1972a; WHO, 1996; Feillet & Leonard, 1998). The rate of biotransformation in humans is high: after oral doses of 40, 80 or 160 mg sodium benzoate/kg body weight, the transformation to hippuric acid was independent of the dose -- about 17-29 mg/kg body weight per hour, corresponding to about 500 mg/kg body weight per day (Kubota & Ishizaki, 1991). Other authors obtained higher values of 0.8-2 g/kg body weight per day (US FDA, 1972a, 1973; WHO, 1996). Hippuric acid is rapidly excreted in urine. In humans, after oral doses of up to 160 mg/kg body weight, 75-100% of the applied dose is excreted as hippuric acid within 6 h after administration, and the rest within 2-3 days (Kubota et al., 1988; Fujii et al., 1991; Kubota & Ishizaki, 1991).

The limiting factor in the biosynthesis of hippuric acid is the availability of glycine. The utilization of glycine in the detoxification of benzoate results in a reduction in the glycine level of the body. Therefore, the ingestion of benzoic acid or its salts affects any body function or metabolic process in which glycine is involved; for example, it leads to a reduction in
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Another metabolite of benzoate is the benzoyl glucuronide. For example, the dog excretes considerable amounts of this metabolite in the urine (20% after a single dose of 50 mg/kg body weight; Bridges et al., 1970). In other species, this metabolite appears only after higher doses of about 500 mg/kg body weight (see above) of benzoic acid or sodium benzoate, after depletion of the glycine pool (Bridges et al., 1970; US FDA, 1972a; Kubota et al., 1988). In cats, glucuronidation is generally very low (Williams, 1967).

In some species, including humans, minor amounts of benzoic acid itself are also excreted in the urine (Bridges et al., 1970; Kubota & Ishizaki, 1991).

Experiments on the distribution and elimination of 14C-benzoate in the rat have shown no accumulation of sodium benzoate or benzoic acid in the body (US FDA, 1972a, 1973).

In the acid conditions of the stomach, the equilibrium moves to the undissociated benzoic acid molecule, which should be absorbed rapidly. Benzoate from sodium benzoate would change from the ionized form to the undissociated benzoic acid molecule. As a result, the metabolism and systemic effects of benzoic acid and sodium benzoate can be evaluated together.

(TAKEN FROM EFSA 2006)

Ring labelled 14C-benzoic acid was given orally at doses in the range of 1 – 400 mg/kg bw to various species including primates, pigs, rabbits, rodents, cats, dogs, hedgehogs, bats, birds, and reptiles. Hippuric acid was the primary urinary metabolite in most species. The ornithine conjugate of benzoic acid, ornithic acid, was the major urinary metabolite excreted within 24 hours in chickens and reptiles. Benzoyl glucuronide was predominant in bats. In humans, >99% of 14C was excreted as hippuric acid within 24 hours (Bridges et al., 1970).

Following oral administration of 375 mg [14C]-benzoic acid/kg bw to rats, 91 – 94% of the radioactivity was recovered in the urine of rats after 72 hours, and only 1 – 6% was present in the faeces. The following metabolites were identified: hippuric acid (70.2 – 84.2%), benzoic acid (0.4 – 12.8%), and 3-hydroxy-3-phenyl propionic acid (0.1–0.2%) (Nutley, 1990).

In order to investigate the types and quantities of beverages that increase urinary hippuric acid excretion, 137 healthy students were recruited and divided into quintiles based on their consumption of non-alcoholic beverages containing benzoic acid. HPLC was used to determine benzoic acid intake from beverages and urinary hippuric acid before, and 1.5 and 3 hours after consumption of various beverages. The range of benzoic acid in 13 beverages was 0 – 1.02 mg/ml and benzoic acid intakes from the beverages for groups 1 – 5, respectively, were: 0.4 mg ± 0.5; 23.4 mg ± 9.8; 55.2 mg ± 2.3; 76.3 mg ± 4.0; and 116.5 mg ± 16.5. Urinary hippuric acid geometric mean concentrations before consuming beverages in the five groups, respectively, were 0.276, 0.270, 0.207, 0.262, and 0.316 g/l; 1.5 hours after beverage consumption they were 0.210, 0.603, 1.026, 1.066, and 1.688 g/l and significantly increased (p<0.001) after adjustment for urinary hippuric acid before ingestion. Three hours after beverage consumption, urinary hippuric acid geometric mean concentrations in the five groups, respectively, were 0.160, 0.232, 0.306, 0.287, and 0.337 g/l (p<0.001). The authors concluded that beverages containing more than 100 mg benzoic acid may increase urinary hippuric acid significantly (Chang et al., 2000).
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**Sodium benzoate**

Male volunteers were given oral doses of 2000 to 5000 mg sodium benzoate. The 5000 mg dose group was given a 5000 mg dose of glycine one hour later and 2000 mg doses every two hours thereafter. Benzoate was excreted mainly as hippuric acid. No free benzoic acid was detected. Minor amounts of benzoyl glucuronide were detected at both doses. Co-administration of glycine with benzoate increased the rate of hippuric acid excretion, indicating that at high dose levels, glycine is rate limiting for formation of hippuric acid (Amsel & Levy, 1969).

After administration of oral doses of 40, 80, and 160 mg/kg bw of sodium benzoate to humans, the mean plasma AUCs of benzoic acid increased disproportionately to the dose, 3.7 and 12.0 times greater respectively for the higher dosages than for the lowest dose, while the mean AUCs for hippuric acid was proportional to dose. Peak plasma concentrations of benzoic acid increased with increasing dose, while peak hippuric acid concentrations did not change. The data suggest that the conjugation with glycine to form hippuric acid is a saturable process in humans (Kubota et al., 1988; Kubota & Ishizaki, 1991).

(TAKEN FROM SCF 1996; opinion released February 25, 1994)

Benzoate is a normal product of intermediary metabolism of phenylalanine and tyrosine and this results in human urinary excretion of a few tens of milligrams of benzoate/kg bw/day. Benzoate administered orally to man is rapidly absorbed and excreted in the urine within 14 hours. The main metabolite is its glycine conjugate, hippuric acid, with the glucuronyl conjugate and free benzoic acid as minor pathways of excretion. The rate limiting step in excretion of hippuric acid is the availability of glycine and this accounts for the glycine depletion which can occur when high doses of benzoate are administered. For example, in man the bolus dose of sodium benzoate causing 80% saturation of the maximal rate of hippuric acid secretion was found to be 28 mg/kg bw (expressed as benzoic acid).
REFERENCES TO CITED EVALUATIONS

For further data on the primary sources, the reader is referred to these reviews.

TARTRAZINE


QUINOLINE YELLOW


ALLURA RED AC


PONCEAU 4R


SUNSET YELLOW FCF


CARMOSINE / AZORUBINE


NA-BENZOATE / BENZOIC ACID


Assessment of the results of the study by McCann et al. (2007) on the effect of some colours and sodium benzoate on children’s behaviour

Glossary / Abbreviations

<table>
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<tr>
<th>Abbreviation</th>
<th>Definition</th>
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<tr>
<td>ADI</td>
<td>Acceptable Daily Intake</td>
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<tr>
<td>AFC</td>
<td>Scientific Panel on Food Additive, Flavourings, Processing Aids and Materials in Contact with Food</td>
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<tr>
<td>bw</td>
<td>Body Weight</td>
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<td>Central Nervous System</td>
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<td>JECFA</td>
<td>Joint FAO/WHO Expert Committee on Food Additives</td>
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<td>SCF</td>
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