

Initial Pre-Meeting Comments from CASAC Oxides of Nitrogen Primary NAAQS Review Panel on EPA’s *Integrated Science Assessment for Oxides of Nitrogen – Health Criteria (Second External Review Draft) (EPA/600/R-07/093aB EPA/600/R-07/903bB, March 2008)*

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## Comments from Professor Ed Avol

### **General Document Comments**

This is an impressive compilation of a great deal of work that reflects the large amount of effort put into it. It is generally well-written and with sufficient detail and reference to provide readers with useful guidance and documentation, as needed. There is a great deal of important and useful information in each of the several chapters, although the respective chapters have a slightly different architecture to them. Several important points are made in some chapters (such as Chapter 2), but not always emphasized at a section or chapter conclusion (as they are, for example, in Chapter 3). Although an excellent summary of the chapters appears as Chapter 5, is it worth considering having a conclusion, summary, or results section at the close of each chapter to re-emphasize or focus on the critical points raised?

### Agency ISA Charge Questions

1. What are the views of the Panel on the characterization of the search strategy for identifying literature, criteria for study selection, the framework for scientific evaluation of studies and causality determination?

Everything seems appropriate and well –described, except for the operational assignments for causality (P1-16, lines 22-23). I agree with the utility of a gradation of confidence in conclusive statements, but it's a bit confusing as to the clear distinction between “sufficient to infer a causal relationship” and “sufficient to infer a likely causal relationship”. Isn't the clarification currently present (“more likely than not”) the definition for the next lower level of confidence (“suggestive but not sufficient to infer a causal relationship”)?

2. To what extent are the atmospheric chemistry and air quality characterizations clearly communicated, appropriately characterized, and relevant to the review of the primary NO<sub>2</sub> NAAQS? Are the properties of ambient oxides of nitrogen appropriately characterized, including spatial and temporal patterns and relationships between ambient oxides of nitrogen and human exposure? Does the information in Chapter 2 provide a sufficient atmospheric science and exposure basis for the evaluation of human health effects presented in later chapters?

In general, yes...However, P2-7 through P2-10, Section 2.3 provides useful (and much appreciated) information regarding the various approaches to measurement of ambient levels of nitrogen-containing species, but does not provide any succinct conclusions or directed message for the reader. Perhaps a closing paragraph or bullet to provide a “take-home” message would be valuable? What discrete message(s) are you are trying to convey in this section?

3. To what extent is the discussion and integration of evidence from the animal toxicology and controlled human exposure studies and epidemiologic studies technically sound, appropriately balanced, and clearly communicated? What are the views of the

Panel on the conclusions drawn in the draft ISA regarding the strength, consistency, coherence and plausibility of NO<sub>2</sub>-related health effects?

This portion is generally excellent - well-written, thoughtfully presented, and meticulously designed. The summary figures, visually presenting the results of numerous studies to gauge consistency of response for a given outcome, is a noteworthy accomplishment that adds to the accessibility of the document. The summary of evidence sections at the close of each section are invaluable and much-appreciated.

(P3-4, line 24) – A comment made here that “Relatively few new clinical and animal (NO<sub>2</sub>) toxicologic studies have been published since 1993...” seems inconsistent with a review of the literature. A PubMed search reveals over 200 publications reporting on NO<sub>2</sub> health outcomes, animal studies, etc...? Please clarify or correct this sentence.

4. What are the views of the Panel on the characterization of groups likely to be susceptible or vulnerable to NO<sub>2</sub> and the potential public health impact of NO<sub>2</sub> exposure?

(P4-9, Section 4.3 and elsewhere) A brief but clear delineation of the operational differences between susceptibility and vulnerability in this document would be useful; this is likely to otherwise create some confusion. Is vulnerable being used to describe “a lack of defenses against” or something more? Is susceptible being used to describe being “prone to a certain response” or something more? This section discusses children and older adults as being “...particularly susceptible to air pollution...”, but aren’t they more vulnerable? If they were missing a specific gene (in the oxidant stress pathway) or had pre-existing disease (such as emphysema, asthma, or possibly diabetes), then they would be susceptible.

5. What are the Panel’s views on the adequacy of this second external review draft ISA to provide support for future exposure and policy assessments?

Overall, this document provides sufficient support for future exposure and policy assessments.

### **Specific Comments**

Chapter I - Introduction P1-13, Fig 1.6-2 – shouldn’t “Mediated Effect” arrows from PM to NO<sub>x</sub> to O<sub>3</sub> go in both directions? NO<sub>x</sub> can affect PM formation which can then have an effect on observed outcomes, and O<sub>3</sub> can affect NO<sub>x</sub> levels which will affect the entire chain of events. Similarly, shouldn’t there be a “Surrogate” arrow from NO<sub>x</sub> to Other Pollutants?

Chapter 2 – Source to Tissue Dose

P2-18, line 1 – (spelling error in San Bernardino, and I don’t think I have ever heard this region referred to as the San Bernardino “Valley”...??)

P2-21, Figure 2.5-1 (spelling error, “In a residence”(68.7%)

Chapter 3 - Integrated Health Effects of NO<sub>2</sub> Exposure  
(No additional specific comments other than those above)

Chapter 4 – Public Health Significance  
(No additional specific comments other than those above)

Chapter 5 – Integrative Summary and Conclusions  
This chapter is an excellent compilation of the assembled data.

#### Comments from Dr. John Balmes

Agency Charge Question 3. To what extent is the discussion and integration of evidence from the animal toxicology and controlled human exposure studies and epidemiologic studies technically sound, appropriately balanced, and clearly communicated? What are the views of the Panel on the conclusions drawn in the draft ISA regarding the strength, consistency, coherence and plausibility of NO<sub>2</sub>-related health effects?

#### GENERAL COMMENTS

The second external review draft of the NO<sub>x</sub> ISA is much improved. In particular, the discussion in Chapter 1 on the “EPA Framework for Causal Determinations” provides an integrative approach to how the agency will assess the evidence presented in the subsequent chapters. Given that this is the first ISA to be produced under the agency’s new process for reviewing a “criteria” pollutant NAAQS, it is important to provide such an integrative approach as a precedent. My major concern about this draft of the NO<sub>x</sub> ISA is that in certain summaries of sections of Chapter 3, the framework described in Chapter 1 is insufficiently applied. I will provide specific examples below.

In my opinion, the discussion of healthy effects of short-term exposure to NO<sub>2</sub> in Chapter 3 is appropriate and fairly well integrated. My major concern with this discussion involves a statement is made on p. 3-14 and repeated on p. 3-15, 3-61, and p. 5-12 that “the onset of inflammatory responses in healthy subjects appears to be between 100 and 200 ppm-min, i.e., 1 ppm for 2 to 3 h.” Figure 3.1-1 is presented to demonstrate this threshold graphically, but appears to be the wrong figure, perhaps an alternative version of Figure 3.1-2. In any event, Figure 3.1-1 presents studies of the effect of NO<sub>2</sub> on allergen-induced lung function and inflammatory responses not the studies of the airway inflammatory effect of NO<sub>2</sub> alone that are described on pp. 3-12 to 3-14. This figure needs to be replaced with one that actually supports this important statement.

The discussion of respiratory effects associated with long-term exposure to NO<sub>2</sub> is important since several prospective cohort epidemiological studies have been published since the 1993 NO<sub>x</sub> NAAQS review that reported adverse effects on growth of lung function. There is a subsection on “Toxicological Studies” on p. 3-89 and 3-90 that is apparently intended to provide mechanistic support for the results of the epidemiological

studies. However, the discussion on pp. 3-95 and 3-96 of “Animal Studies of Long-Term Morphological Effects to the Respiratory System” is both more comprehensive and concise. I would delete the subsection on p.3-89.

On p. 3-90, the concluding paragraph for section 3.4.1 on “Lung Function Growth” provides no assessment of the strength of the association between long-term exposure to NO<sub>2</sub> and decreased rate of growth of lung function among children. This paragraph merely states that the epidemiological studies of long-term exposure to NO<sub>2</sub> are likely confounded by other ambient pollutants. Unfortunately, section 3.4.5, which is supposed to be the summary and integration of evidence on “Long-Term NO<sub>2</sub> Exposure and Respiratory Illness and Lung Function Decrements” also does not provide an assessment of the strength of the association. This is a critical deficiency given the relative importance of the issue of potential respiratory effects of long-term NO<sub>2</sub> exposure. The section also does not directly assess the strength of the association for either “Asthma Prevalence and Incidence” or “Respiratory Symptoms”. It is not until Chapter 5 that the strength of the association for long-term exposure to NO<sub>2</sub> and respiratory morbidity is assessed.

Another concern that I have regarding section 3.4.5 is the long discussion on animal toxicological studies that have found effects of NO<sub>2</sub> on host defense against respiratory infections on pp. 3-98 to 3-100. While I believe that this discussion is technically accurate, I find that it is unnecessary and distracting given the relative lack of discussion of the strength of the epidemiological evidence on long-term exposure to NO<sub>2</sub> and asthma and the total absence of such discussion regarding respiratory symptoms.

Agency Charge Question 2. What are the views of the Panel on the characterization of groups likely to be susceptible or vulnerable to NO<sub>2</sub> and the potential public health impact of NO<sub>2</sub> exposure?

Chapter 4 does an adequate job of discussing the public health significance of exposure to ambient NO<sub>2</sub>. The chapter Summary (4.5) indicates that persons with preexisting respiratory disease (especially asthma), children, and older adults may be more susceptible to the effects of NO<sub>2</sub> exposure. I agree with this assessment of the evidence. There is one statement in section 4.5, however, with which I am uncomfortable. The final sentence on p. 4-16 states that “evidence, albeit inconsistent, exists for a gender-age-based difference in susceptibility” regarding the effects of NO<sub>2</sub> on children with asthma. I find this statement to be too strong. While I agree that there is evidence that the incidence of asthma differs between boys and girls with age, the evidence for a gender-age-based difference in susceptibility to effects of NO<sub>2</sub> exposure in asthmatic children is too limited to draw any conclusions. I would delete this sentence.

The major problem I have with Chapter 5 is that the relatively strong evidence for an effect of long-term exposure to NO<sub>2</sub> on growth of lung function in children reported by both the Southern California Children’s Health Study and the Mexico City study (Rojas-Martinez et al.) is diluted by combining discussion of these studies with those on asthma and respiratory symptoms. In my opinion, the issue of the strength of evidence for a

long-term effect of NO<sub>2</sub> on growth of lung function needs to be dealt with more directly in the document, in both Chapters 3 and 5.

## SPECIFIC COMMENTS

I have a number of suggested wording revisions that I will submit later.

### Comments from Dr. Douglas Crawford-Brown

The primary question addressed in this review is in reference to Chapter 5, Integrative Summary and Conclusions, and focuses on the adequacy of the ISA (especially Chapter 5) to provide support for future exposure and policy assessments. I begin by noting that this second draft is a significant improvement on the first. The authors have dealt with the majority of my comments on the first draft, and have responded to the majority of comments made by the committee as a whole in our earlier review. There is beginning to emerge a vision for what the ISAs in general are intended to accomplish, so this is a positive step forward for the EPA staff.

On the primary issue of adequacy as a basis for exposure and policy assessments, my comments are of two kinds. First, Chapter 5 does indeed summarize the information from earlier chapters that would be relevant to exposure and policy assessments. The authors have done a good job of both accurately reflecting that earlier material and choosing the material that is most relevant. They have drawn appropriate conclusions from that earlier material, including assessments of the strengths and limitations of the conclusions. While I might still have some quibbles over the lack of formal assessment of the epistemic strength of conclusions, I fully agree with the judgments they have made as to whether the evidence for each category of effect is conclusive, supportive, inconclusive, etc. I don't think I could trace the judgments of the authors back to any specific reasoning they have done, because the only reference to that reasoning is a vague citation of the Hill criteria (criteria that seem to me woefully inadequate as a philosophical basis for rigorous assessments of epistemic status). But the judgments of the authors nonetheless appear to be the correct ones based on the available evidence.

The second issue is whether Chapter 5 not only summarizes accurately the information in earlier chapters (which it does) but also provides an adequate basis for exposure and policy assessments. Here I am less comfortable the chapter, or even the entire document, has met the target. It is certainly the case that both the document and the chapter point the reader in directions that will be important for exposure and policy assessments. The reader is guided to judgments on the appropriate sensitive subpopulations that should be considered; on the specific effects that should be considered; on issues related to the relationship between ambient monitoring results and personal exposures; and on the issue of whether epidemiological studies are confounded by exposures to the mix of air pollutants that normally accompany exposure to oxides of nitrogen. An assessor will, therefore, find much useful information in the document and in Chapter 5.

However, neither the chapter nor the document provides an adequate basis for performing exposure and policy assessments quantitatively. It is not possible from the document alone to see what the levels of exposure will actually be to any specific population in any specific geographic location. I realize this level of detail may not be what is intended by an ISA, but then I am left wondering what could be meant by asking whether the document provides an adequate basis for exposure assessment. The only way it could do that, in my mind, is to summarize the actual exposure information available, guiding the reader to understanding how well that information will allow accurate estimates of exposure to specific subpopulations in specific geographic areas. The ISA does not summarize exposure information, or even monitoring information, but instead summarizes the strengths and limitations of that information. The latter is an important goal, as it will place caveats on the use of more detailed information in later exposure assessments. But it leaves the task of assembling the information on concentrations, and converting these to estimates of exposure, to a later step in the assessment process. ***As a result, I am not comfortable that the ISA provides an adequate basis for subsequent exposure and policy assessments, but rather provides an adequate basis for these subsequent steps to understand the strengths and limitations of the available data. Having said that, this may be all that the ISAs are intended to accomplish, in which case the current document would be judged adequate.***

I also was looking for some sort of scientific statement as to how the information available ought to be used in conducting an exposure or policy assessment. This can be done without wandering into policy decisions. Are the authors suggesting that no exposure-response curve can be developed, and hence the assessments must be conducted in ways similar to non-cancer risk assessments (with a presumed threshold of effect)? Should an uncertainty factor be applied? Would the goal of an assessment be to estimate the number of people in the U.S. population exposed at levels above the threshold (or with a hazard quotient above 1)? Should there exposure be averaged over an hour, or 4 hours or a day or year? These are questions that will crucial in a future assessment and can be answered scientifically without getting into the policy realm, so I expected to see them here if the document is to provide an adequate basis for exposure and policy assessments.

I then have a series of specific issues to raise.

1. I continue to be less than convinced by the epistemic assessments performed, or at least the formal descriptions of the assessments. The reference on page 5-1 is to the Hill criteria (which are completely inadequate philosophically in my view) and to “other pertinent frameworks”. No reference is provided to these other frameworks, so I have no idea what they are. I realize the ISA is not a philosophical document, but some better guidance as to how the specific judgments of the quality of conclusions are developed is needed. There is no problem in stating that they are informed judgments based on some set of criteria (and then stating these criteria). Having said that, I do agree with the conclusions drawn and so will not push this point too strongly.

2. The fourth bullet on Page 5-1 asks whether new data affect the plausibility of judgments about oxides of nitrogen causing adverse health effects. I had expected to see some further qualification such as “adverse health effects at levels of exposure of at levels found in the environment”.

3. On Page 5-3, the bullets don't seem to follow at all from the opening sentences of the paragraphs. I agree with what is contained in the bullets, but they seem to me unconnected to the sentences that presumably introduce them. I have read this paragraph several times and just don't see the link.

4. In that same list of bullets, 6<sup>th</sup> bullet, the authors talk about artefacts. There is no explanation as to what an artefact means, how it affects exposure or policy assessments, what a positive artefact would be (presumably in contrast to a negative one), etc.

5. The listing of issues in section 5.2.2 is a good one, and is certainly useful to anyone conducting an exposure assessment. But I point back to my earlier comment that even if the issues raised here are the correct ones, it is not possible to take this and form a basis for an exposure assessment because specific data are not provided and summarized. I also expected a bullet suggesting how exposures SHOULD be estimated, rather than simply listing the limitations in any exposure measures. I would expect a recommendation as to the ratio of personal to ambient levels in specific settings, the time period over which exposure should be averaged, etc. I also expected some conclusion as to whether it is even valid to use ambient monitoring results, from the network set up for regulatory monitoring, as a starting point for estimating exposures.

6. The last paragraph in section 5.2.2 cautions against strict conclusions being drawn based on the epidemiological studies due to a lack of conclusive evidence that ambient and personal exposures are completely correlated. The implications of a lack of complete correlation depend on the differences in exposure levels between the different exposure categories in the study. The implications become more significant as the exposure groupings are closer together in exposure, in which case there can be significant misclassification and therefore bias towards the null (generally at least). I think a more nuanced conclusion here is needed.

7. I found the summary tables quite useful, so the authors are to be applauded for developing them in this coherent fashion.

8. In the sections on effects, it would be useful if the authors were to provide information as to whether an increase in exposure changes the fraction of people with effect, the frequency with which a subset of people develop the effect, the severity of the effect, etc. I realize that this distinction may not be so important in a regulatory decision (where one person getting the effect N times counts the same as N people getting the effect once), and that the available exposure-response information may not allow development of exposure-response curves, but some comment on this issue would be useful.

9. In the last paragraph (Page 5-22), the authors correctly conclude that it is plausible to believe that effects are occurring at levels of exposure below the current NAAQS. But the question here is “how far below?”. I should think that the science assessment (this document) would be the place where the EPA states clearly the answer to this crucial scientific issue – an issue that will greatly affect the policy assessment.

#### Comments from Dr. Terry Gordon

Charge Question 3. To what extent is the discussion and integration of evidence from the animal toxicology and controlled human exposure studies and epidemiologic studies technically sound, appropriately balanced, and clearly communicated?

The ISA document is a tremendous improvement over the first draft. There are appropriate-length discussions of key studies and the overall message of the ISA is clearly communicated. There are, perhaps, sections of the animal toxicology data that have too much detail given the high(er) concentrations of NO<sub>2</sub> used in some studies. As mentioned in past discussions by the Panel, animal toxicology studies should be included in the ISA only when they are of relevant concentrations.

What are the views of the Panel on the conclusions drawn in the draft ISA regarding the strength, consistency, coherence and plausibility of NO-related health effects?

The development of the guidelines to evaluate the strength, consistency, coherence, and plausibility of NO<sub>x</sub>-related health effects is critically important to the ISA. In a similar fashion, the use of structured language to describe relationships between exposure to NO<sub>x</sub> and adverse health effects is an important improvement to the ISA. The 5 descriptor categories appear well thought out and appropriate, but discourse on these categories may bring modifications to future ISAs if not this one.

Minor Comments:

Disclaimer page

Page ii – first line – First or second external draft?

Page ii, para 2 – typo for (4) before “to set....”?

Page 1-17, para 2 – Why would tox and cancer have a linear response? Many researchers say otherwise. What does dose-transitional mean?

Page 2-4, last line – ‘at low temperature’ is imprecise; does this mean ambient temperatures?

Page 2-21, figure 2.5-1 – Label for largest part of pie chart should be ‘In a residence’.

Page 3-6, table 3.1-1 – Under peripheral blood, ‘Total macrophages’ should be (re)moved as they are in the lower airways.

Page 3-8, para 3 – The 3 to 9 ppm 2 week study has little relevance to the toxicity of exposure to peak ambient levels of 0.05 or 0.1 ppm NO<sub>2</sub>. Also, it is unbalanced to give the details and results of this high dose study and then say that alveolar macrophages are a sensitive target for NO<sub>x</sub>’s effects and give no refs on macrophages and just refer the reader to the Annex.

Page 3-12, figure 3.1-1 – The figure legend needs more details (e.g., what are the + and – on the Y-axis for; what does the \* mean?; what do the 3h and 6h mean? Some of this is explained later in the legend of figure 3.1-2.

Page 3-22, first and second line – Needs a ref.

Page 3-22 – There is a lot of detail here for high dose (5 ppm and 20 ppm NO<sub>2</sub>) animal studies. Please consider cutting the long paragraph and summarizing in a couple of sentences.

Page 3-24 – Add ‘Nonspecific’ before ‘Airways’ in the title for consistency with the last 2 subtitles.

Page 3-25, para 2 – Delete ‘of’ before ‘as low as....’. Having stated that effects can occur at 0.26 ppm, why include the 1-sentence results of a 5 ppm study?

Page 3-44, para 4 – Elsewhere the Gong study had 18 subjects.

Page 3-47 – I may be reading this all wrong but some of the bars on the figure make no sense: on the preceding page, it says the Tolbert study had a 2% increase with a 95% CI of 0.5, 3.3. The figure shows a significant box with error bars totally dissimilar the reported 95% CI. The same goes for the Barnett (2005) study which says something different in the text (page 3-47) than is in the figure. Should all data be re-checked?

Page 3-59, second line – insert ‘for asthma’ after ‘.... or ED visits’.

Page 3-59, line 7 – Adding ‘other’ before ‘diseases’ would make this sentence clearer.

Page 3-62, lines 4, 5 – I’m not sure what is meant by ‘....NO<sub>2</sub> effects to be mediated by other pollutants or exposures;’

Pages 3-62, 63 – Although the physiology descriptions are good, the text has 15 lines describing HRV and then 5 lines to say there have been mixed results. The same goes for the repolarization section – 5 lines to describe what it is and then 1 line to say there was no effect. Condense?

Page 3-79, 2nd para – The statement regarding animal studies and mortality as an endpoint in acute studies is not correct. Several studies have looked at LC50 values for NO<sub>2</sub> in different species.

Page 3-90, para 3 – This paragraph on epidemiology appears to be misplaced in this tox section.

Page 3-96, para 1 – A mention of the higher doses used in animal tox studies could be added to qualify the last sentence.

Page 3-111, para 3 – Add ‘a’ before ‘sensitivity’.

Page 3-118, para 2 – Should the last line of this para read, 1.0 to 2.8, or 0 to 1.28?

Page 4-4, first line – Has this really been recapitulated in preceding sections of this chapter?

Page 4-8, para 1 – Adding a conclusion sentence for this section on asthma would be appropriate. The same goes for the other susceptibility sections.

Page 4-12, para 1 – The sentence states that in-vehicle concentrations are 2 to 3 times ‘ambient’. Does this mean that it’s higher in the vehicle than just outside the car? Or does it mean compared to non-traffic ambient levels?

Page 4-13, first line – Ponce 2003 or 2005?

Page 4-16, para 2 – This is a strange intro sentence for the Summary section.

Page 5-26 – The title for the table is unclear – Legend of Figure 5.3-1??

#### Comments from Dr. Dale Hattis

1. What are the views of the Panel on the characterization of the search strategy for identifying literature, criteria for study selection, the framework for scientific evaluation of studies and causality determination?

I used a few different methods to evaluate the results of the literature search. First I read through all of the references cited at the end, noting the frequency of 2006-2008 references, and selecting in particular titles that appeared highly relevant to the analysis of health hazards. From the large numbers of 2007 references and the occasional 2008 citation, it is clear that the authors have brought their literature searches up to date as of the present. I then retrieved the abstracts for about a dozen interesting recent references and checked how the revised ISA reported on them in the main text or tables. In every case I found that the description in the text or tables corresponded reasonably to the abstract.

#### Comments from Dr. Donna Kenski

*Charge Question 2: To what extent are the air quality characterizations and analyses clearly communicated, appropriately characterized, and relevant to the review of the primary NO<sub>2</sub> NAAQS? Are the properties of ambient oxides of nitrogen appropriately characterized, including spatial and temporal patterns and relationships between ambient oxides of nitrogen and human exposure? Does the information in Chapter 2 provide a sufficient atmospheric science and exposure basis for the evaluation of human health effects presented in later chapters?*

This second draft ISA is much improved from the first draft ISA and it was gratifying to see the panel's suggestions from the last review incorporated to a large extent. The discussions of measurement interference and its spatial and temporal variability were much better, as was the discussion of ambient concentrations of NO<sub>2</sub>. The paragraph on NO<sub>2</sub> increases in other countries was somewhat irrelevant (p. 2-13, lines 25-30) and could be deleted or moved to the section on policy-relevant background concentrations. Sec. 2.4.5 on concentrations of NO<sub>x</sub> species is still quite limited and I think more information on these various species and their interrelationships would be helpful. Data are admittedly sparse but there are more papers that could be usefully summarized. Two that I know of are:

*Continuous wet denuder measurements of atmospheric nitric and nitrous acids during the 1999 Atlanta Supersite*, Atmospheric Environment, Volume 37, Issues 9-10, March 2003, Pages 1351-1364

Zhang Genfa, Sjaak Slanina, C. Brad Boring, Piet A. C. Jongejan, Purnendu K. Dasgupta

*Measurements of gaseous HONO, HNO<sub>3</sub>, SO<sub>2</sub>, HCl, NH<sub>3</sub>, particulate sulfate and PM<sub>2.5</sub> in New York, NY*, Atmospheric Environment, Volume 37, Issue 20, June 2003, Pages 2825-2835,

Abdul Bari, Vincent Ferraro, Lloyd R. Wilson, Dan Luttinger, Liaquat Husain

Section 2.2.1 on sources of NO<sub>x</sub> still needs to be augmented with a *shortened* version of Annex Table AX2.6-1 (the old AX2-3) giving quantitative data on emissions contributions from major source categories. Fig. 2.2.1 is nice but just doesn't convey any quantitative sense of emissions.

The reorganized Sec. 2.5 on exposure is a much better integration of the relevant information.

In the previous CASAC review, we requested that chapter 2 in particular include section summaries, as was done so effectively in later chapters. It doesn't seem like this has been accomplished, but it would help tie up this chapter and make the conclusions drawn from it in Chapter 5 more obviously connected to the relevant sections.

Aside from these fairly minor tweaks, I found this section of the ISA satisfactory and I think (together with the supporting documentation in the annexes) that it provides a suitable basis for the Risk and Exposure Assessment.

Other comments:

Chapter 5 is great; it gave a fair and balanced presentation of the studies reviewed and the conclusions drawn. Very helpful.

In our review of the first draft ISA, we requested that EPA consider multipollutant approaches to managing air quality. While acknowledging that changing the traditional one-pollutant-at-a-time approach is likely to take considerable time and effort to implement, I'd like to encourage EPA to move in that direction with the ISA process.

The inclusion of data for multiple N species as well as some sulfate and oxidant chemistry was a step forward.

p. 2-17, Fig. 2.4-6d, title of figure should be Weekend, rather than Weekday

Comments from Dr. Steven Kleeberger

The document reads very well. I have very few comments that would not be considered only editorial- or style-related. However, I found the Figures 3.1-1 and 3.1-2 to be somewhat confusing. While the information presented is very useful, I would suggest that the data would be better presented in tabular form rather than figures. At first glance, the +/- on the y-axis suggests a degree or quantitation of positive or negative finding. It would be simpler simply to list the studies in a table with negative and positive categories.

Comments from Dr. Timothy Larson

General Comments:

I am limiting my comments to the exposure issues covered in Chapter 2 and parts of Chapter 6. In general, this document is much improved on the first draft and the authors should be commended. The issues raised by the committee upon review of the first draft have been addressed for the most part. I think the Chapter 6 summary of the topics in Chapter 2 is reasonable and consistent. The discussion of NO<sub>2</sub> infiltration seems a little long.

However, I disagree with the statement beginning on page 2-31 about street canyons being complicated and modeling their effects to be “highly problematic”. In fact, recent work has shown that both CFD and much simpler integral models provide reasonable predictions (c.f. Di Sabatino S, Buccolieri R, Pulvirenti B, et al. (2008) *Atmos. Environ.* 41 (37) , 8316-8329). In addition, there are a number of recent studies showing good prediction skill with these simpler models (c.f. Mensink C, Cosemans G (2008) *Env. Modeling & Software.* 23 (3) , 288-295; Berkowicz R, Ketzel M, Jensen SS, et al. (2008) *Env. Modeling & Software.* 23 (3) , 296-303; Ghenu A, Rosant JM, Sini JF (2008) *Env. Modeling & Software.* 23 (3) , 314-321). This seemingly minor point is in fact rather important to the interpretation of the spatial heterogeneity of NO<sub>2</sub> in built up urban areas. These studies and others (e.g. Vardoulakis S, Gonzalez-Flesca N, Fisher BEA, et al. (2005) *Atmos. Environ.* 39 (15) , 2725-2736) indicate that street canyons are similar in NO<sub>2</sub> concentrations to on-road values in otherwise open areas, i.e., enriched by about a factor of 2 above measurements taken away from the road in more open areas. This adds another dimension to the exposure assessment, namely the fact that pedestrians spend time walking in these canyons and having windows opening onto these canyons and can therefore experience exposures for equal or greater times than they do on roads in transit (they may not even own cars in dense urban areas).

Specific comments:

Page 2-25 line 26 and page 2-26 line 6: Text seems to arrive at different conclusions about passive samplers without comment.

Page 2-35 line 13: This sentence needs a reference that indoor pollution affects outdoor levels.

Page 2-49 line 6: Reference to Table 2.5-5, but table does not exist nor does Table 2.5-6.

Page 2-51 line 16: The NO<sub>2</sub> east-west spatial variation in greater Los Angeles varies broadly with distance from the coast due to well known meteorological and chemical phenomena. NO<sub>2</sub> levels in Riverside are determined in large part by pollution transported from upwind urban areas to the west.

Page 2-55 Table 2.5-9 is presented without comment.

#### Comments from Dr. Edward Postlethwait

1. In general the document shows considerable improvement over the initial draft and thus the EPA staff should be commended.
2. The issue of endogenous NO<sub>2</sub> generation appears not to have been addressed as a potential contributor to uncertainties with regard to data interpretation, thresholds, and assessing posited mechanisms of action. This is especially important for three of the posed questions in section 5.1 regarding how new information altered support for occurrence of health effects, at what levels of NO<sub>x</sub> do health effects of concern occur, and plausibility of adverse health effects.
3. On page 2-61, the section dealing with modeling NO<sub>2</sub> dose should be clarified with regard to NO<sub>2</sub> flux. The term “NO<sub>2</sub> flux to air-liquid interface” does not accurately describe the net movement of NO<sub>2</sub> from the intrapulmonary gas phase into the surface lining layer, or potentially epithelial cells. If NO<sub>2</sub> flux occurs as written, there is a disconnect between the sites of focal injury and mass transfer from the intrapulmonary gas phase.
4. To this reader, there is a consistent ambiguity in the way NO<sub>2</sub> thresholds are presented. As written, it is not especially clear in the document whether the lack of documented thresholds means that NO<sub>2</sub> related effects can be extrapolated to zero [NO<sub>2</sub>] in a linear fashion or that threshold concentrations have not been identified due to the numerous confounding factors. The document would be strengthened if this issue was revised throughout to unequivocally present a consistent interpretation.
5. It is somewhat curious that the document supports causal relationships between NO<sub>2</sub> exposure and acute but not chronic health effects. If NO<sub>2</sub> is able to induce short term effects at the denoted concentrations, one would anticipate that individuals residing in the same geographic locale would also experience long term impacts. Thus, it may be

useful to consider whether this represents an inconsistency or is due to any number of mitigating factors.

6. Per discussions from the initial review meeting, the document does not extensively link the potential occurrence of short term NO<sub>2</sub> spikes and health outcomes. A more thorough discussion in the summary chapter (Chapter 5) of the possible occurrence rates, NO<sub>2</sub> spike concentrations and thus exposures relative to the current long term average values, and observed effects would strengthen the document and provide the impetus to help support a shorter term standard if warranted.
- 7.

#### Comments from Dr. Armistead Russell

Again, like the first draft, the 2<sup>nd</sup> Draft ISA is an effective document for providing the information needed to conduct the risk and exposure assessments. It is substantially improved from the last draft. However, I was a bit disappointed that some pieces did not get changed between the 1<sup>st</sup> and 2<sup>nd</sup> drafts.

1. I still think there should be a table of sources in Section 2.2.1. There is no Table 2-3 in the Annex: it is Table 2.6-1. Having a table in the ISA might help OAQPS note that their commercial air craft estimate for Philadelphia does not look right. They should also note that being an elevated source also leads to the emissions being more dispersed. Indeed, the highest NO<sub>2</sub> levels found in cities are not from EGUs.
2. In the chemistry section, a bit more should be said about NO to NO<sub>2</sub> transformation as NO<sub>2</sub> is the apparent indicator species.
3. The measurement section is still a bit biased when highlighting the Mexico studies. Mexico City is a unique place, so I tend to downplay those results. Switzerland is more appropriate. There really should be some US-based studies.
4. The section on ambient concentrations presents an appropriate level of information on NO<sub>2</sub> levels in the US at the country/urban scales, but should provide more information on NO at the same time. I would treat them together, with much of the transport discussed as transport and decay of NO<sub>x</sub>. If one looks at the REA, an important issue is roadway levels. This is hardly treated here. This section should discuss roadway levels of NO<sub>2</sub> and NO, as opposed to covering it later in 2.5.4. A major missing component of ambient characterization is that of other pollutants. It is important to show the correlation between NO<sub>2</sub> and some other species (e.g., EC) and I think it belongs here as opposed to later in 2.5.8. It would seem to fit more naturally here.
5. 2-31. It is not apparent that a 15 m monitor will lead to an underestimate of what people are breathing given that in cities (e.g., NY) air inlets in to homes may be that high. Also, as noted in the REA, very few of the monitors are at 15m. Give a balanced presentation.
6. 2.5.4: This section is fine, but could go deeper to provide the detail that is ultimately needed in the REA. It should discuss the conversion of NO to NO<sub>2</sub> with information as to conversion rate and how quickly NO falls off and NO<sub>2</sub>

- rises. The current REA uses little from this section, instead deriving its own fit to the decay. It is also not apparent how the REA used the information about intrusion of the vehicles own exhaust in to the cabin.
7. 2.5.8. This is a very important section, and I think it does a reasonable job of providing the type of information needed to see how NO<sub>2</sub> correlates with other important species, including EC. One of the problems with having this section much later in the document is that they then go over points discussed earlier (e.g., diurnal variation).
  8. Conclusions:
    - a. P5-3, L19. In the body of the ISA, they never use 25%... even though I think the 50% value given earlier is extreme, whatever number chosen here should be supported in the ISA.
    - b. P5-3, L21-23. I would not add “are difficult to predict”. CMAQ can do so.
    - c. 5-3, L30: Precise is not the correct word here. Possibly accurate, but I would say that it may be accurate enough for the job at hand, so I would back off on this statement altogether.
    - d. P5-5, L7: While true, we find out in the ISA this is not a real problem since most monitors are not at 15m.

Something to think about for future ISA's: you should have a section discussing the models that might be appropriate for estimating ambient levels and, possibly, exposure. In this case, it would be AERMOD and APEX. This section should provide the model formulation, inputs and an evaluation of its capabilities. Modeling is discussed to some degree in the NO<sub>x</sub>-SO<sub>x</sub> SNAAQs ISA, which is good (though the committee wanted more evaluation).

#### Comments from Dr. Jonathen Samet

##### General Comments:

The second draft ISA for oxides of nitrogen (NO<sub>x</sub>) is improved and responsive in many respects to concerns raised by CASAC members, including myself, at the review of the first external review draft. The Agency's staff has attempted to be responsive in setting out a better framework for evidence identification, evaluation, and synthesis. I am hopeful that continued progress will be made to sharpen this process, in order to address issues that are still unsolved. Additionally, I think that the peer review process would be enhanced generally by the preparation of a note to CASAC that sets out the Agency's responses to major concerns raised by the CASAC reviewers. Such responses would be consistent with usual practice of peer review, and would provide a trail, documenting how comments made by CASAC were taken into account.

My responses to the charge questions from the Agency follow, and the attached table lists specific comments.

Charge Question 1:

In response to comments with regard to the Agency's approach to literature identification, study selection, and synthesis of the evidence, as well as causal inference, an extensive annex has been prepared that reviews a number of relevant frameworks. The background is a useful foundation for justifying the selected approach. The Agency has made a number of changes in Chapter 1 that are responsive to prior critiques. In particular, there is a description of literature selection, an approach to evaluating evidence for inferring causality is provided, and a reasonable set of descriptors of the strength of evidence for causation is offered.

On reading the draft ISA, there has been some impact throughout the document of having a better prepared first chapter and this framework. However, the sections that synthesize the evidence are still somewhat loosely written, and do not systematically apply the guidelines offered by Bradford Hill, and adopted by the Agency. (Note, that these should not be renamed as "decisive factors", a misnomer). Too often, sections that are offering judgments as to the strength of evidence use such language as "taken together or "integrating" without a more specific application of the criteria offered by Hill. Nonetheless, the new Chapter 1 and the approach set out represents an advance over the earlier draft. Discussion is needed as to whether the list developed by Hill should be replaced with the shorter set used in the Surgeon General's Report. Perhaps, a comparison could be made with a test case.

One issue that is left unaddressed is publication bias, a reasonable concern given that many studies address the health effects of air pollution and employed multivariable models to attempt to isolate the effects of particular pollutants. There must have been some tendency on the part of investigators to report positive associations. This topic needs to be mentioned, as the existence of publication bias would be difficult to set aside for some of the epidemiological studies.

Charge Question 3:

This charge question relates to the integration of evidence from the various lines of investigation. Of particular concern is the plausibility of effects observed in epidemiological studies in the context of animal and human toxicology. There is no doubt that high levels of NO<sub>x</sub> can injure the lung and other organs. For setting the NAAQS, the plausibility of effects at ambient concentrations is particularly relevant. I am concerned that the draft ISA has only partially addressed the plausibility of effects at current ambient concentrations and at those investigated in a number of the epidemiological studies. The ISA appropriately notes that both clinical and animal studies are carried out at doses well above those that are typical for population exposures. The document would be improved if more attention could be given to considering the relevance of mechanisms observed at higher levels to effects at ambient levels.

Charge Question 4:

Chapter 4, “Public Health Significance”, provides an overview of populations potentially at increased risk from exposure to NO<sub>x</sub>.

The listing of susceptible subgroups covers those of both particular relevance and of general relevance. The various groups identified are appropriate, although I have concern that EPA will follow the same template for all pollutants without giving sufficient consideration to the relevance of particular subgroups for particular pollutants.

#### Charge Question 5:

The revised document is greatly improved. It identifies health effects associated with exposure to nitrogen oxides and the strength of evidence supporting causality of associations. It makes an attempt to assess whether effects would be expected at current ambient concentrations. This is one of the weakest aspects of the current draft. Consider, for example, the conclusory language from page 5-22: “integrating across the epidemiologic human clinical and animal toxicological evidence presented above, we find that it is plausible that current exposures can result in adverse impacts to public health at ambient concentrations below for current NAAQS for NO<sub>2</sub>.” This sentence addresses the most critical matter around the need to revise the NAAQS: are there effects being observed at current ambient concentrations? The sentence addresses plausibility. Some statement with regard to the degree of certainty is needed, no matter how difficult it may be to characterize uncertainty.

#### Specific Comments:

Page #	Line #	Comment
1-7	5	Not certain this is true!
1-10	2	Along with considerations of plausibility
1-10	24	That may lead to confounding.
1-10	28	“...homogeneous groups with of the confounding...”
1-12	22	What does this mean?
1-13	1	Note, the following discussion refers to statistical models. The word “model” has many uses.
1-14	23	Should not label as “decisive factors”, these are guidelines.
1-18	8	Because of the experimental manipulation of exposure.
1-18	16	There should not be uncertainty about design and execution.
3-1	24	“obscured” What does this mean?
3-2	3	“partitioning the variance” Not the same as estimating an effect.
3-2	13	“effects observed at O-...” On what basis?
3-2	19	“health effects (and markers of injury) that...”
3-3	9	Could be true, but supporting references??
3-7	5	At what concentration?
3-10	1-2	This is a very sweeping claim. What does it mean?
3-17	4-7	Aren’t there more informative ways to show the data, e.g., as a forest plot.

Page #	Line #	Comment
3-19	29	In what way are these taken together?
3-20	29	Not the right place for research recommendations.
3-24	17-19	Speculation; should be deleted.
3-26	5	Isn't this inconsistent with an NO <sub>2</sub> effect?
3-40	4	Well-recognized
3-46	6	"pneumonia" is a LRI
3-46	19-22	"Collectively..." a conclusion—out of place
3-62	15-18	Where do the authors stand on this possibility?
3-101	1	Certainly, the possibility that traffic-related carcinogens are the etiologic agent needs to be raised.
3-106	11	Why confounding, implying underlying causation?
3-107	13	"(2000..." ) and earlier
4-5	29	Be careful not to overinterpret
5-1	28-29	?
5-22	21	"Integrating across the epidemiologic" How? What does this mean?

#### Comments from Dr. Richard Schlesinger

Overall, this is a very good document that clearly integrates the essential concepts required to make a judgment related to health effects from NO<sub>2</sub>.

p. 1-16, line 22. It may be better to change the first bullet to "sufficient to infer a definite causal relationship" since this would make it more consistent with the second bullet indicating a "likely" causal relationship. The way it is written, it is not clear what the difference is between "causal relationship" and "likely causal relationship."

p. 5-13, line 15. The 5 ppm exposure is quite high and its relevance to the document is not clear.

p. 5-16, lines 13-14. Does this sentence mean that the effects observed could have been due to NO<sub>2</sub> or are most likely due to confounder pollutants?

Table 5.3-1. This is an excellent table.

#### Comments from Dr. Christian Seigneur

*Charge question 2:*

*To what extent are the atmospheric chemistry and air quality characterizations clearly communicated, appropriately characterized, and relevant to the review of the primary NO<sub>2</sub> NAAQS?*

The second external review draft has been considerably improved in that regard. For example, Figure 2.2-1 is more accurate; the variability of the NO<sub>2</sub> fraction of NO<sub>x</sub> emissions is now correctly discussed, and the spatial variability of NO<sub>2</sub> concentrations near sources (in particular, roadways) is properly described.

*Are the properties of ambient oxides of nitrogen appropriately characterized, including spatial and temporal patterns and relationships between ambient oxides of nitrogen and human exposure?*

The spatial and temporal patterns of NO<sub>x</sub> and NO<sub>2</sub> are appropriately described. As mentioned above, the strong spatial gradients observed near roadways are discussed. The temporal variability of NO<sub>x</sub> and NO<sub>2</sub> concentrations is also well characterized with sufficient detail being provided in the Annex (e.g., diurnal and seasonal variability).

*Does the information in Chapter 2 provide a sufficient atmospheric science and exposure basis for the evaluation of human health effects presented in later chapters?*

There is one aspect of the link between atmospheric science and exposure/health effects that needs to be better discussed in the ISA. Currently, measurement error is highlighted early on in the document (Section 1, p-1-11) as important when adjusting for spurious associations. The difference between true and measured ambient concentrations is identified as one of the components of measurement error. Such measurement errors are then discussed in detail in Section 2.3 for outdoor measurements and in Section 2.5.2 for indoor measurements using passive samplers. Some estimates of measurement uncertainty are provided. For outdoor measurements with the Federal Reference Method (FRM), an average interference from NO<sub>z</sub> compounds of 22% is given for Mexico City data with peak interference of 50%. A comparative study in Switzerland is reported to have shown average errors of 10% in winter and 50% in summer. This section concludes that the interference is likely to be on the order of 10% or less in winter, but much larger in summer. It is also stated that the interference from NO<sub>z</sub> compounds is less significant near the emission sources (because there has not been enough time for NO<sub>z</sub> species formation). Passive samplers used in individual exposure studies are described in Section 2.5.2 as being within 10% of the FRM.

The epidemiological studies presented in Section 3 have used in most cases NO<sub>2</sub> measurements from FRM instruments or passive samplers. Therefore, one may imply that there may be significant uncertainties associated with the results of those studies because of the FRM interference errors discussed in Section 2. However, this aspect of the health effect uncertainty does not appear to be discussed. In reality, I anticipate that most NO<sub>2</sub> exposure occurs near emission sources where NO<sub>z</sub> concentrations are low and, therefore, interference error is small. Also, the seasonal variability of NO<sub>2</sub> concentration shows that in most cities higher concentrations occur in winter when the interference error is 10% or less. It may, therefore, be possible to consider that the NO<sub>2</sub> measurement error is not a major source of error in health effects studies. However, such a point needs to be made clearly in the ISA. One possibility is to provide a more definitive discussion of the implication of the interference of the FRM at the end of the introduction of Section

2.3 (p. 2-9). The fact that NO<sub>z</sub> species concentrations are low when NO<sub>2</sub> concentrations are high (near emission sources, in winter) and the reasons why (not sufficient time for reaction, low photochemical activity) should be explicitly stated. Then, the implications for the health effects studies need to be articulated clearly.

### **Annex 2.7.1. Chemistry-transport models**

This Annex section has been rewritten and a discussion of local-scale dispersion models has been added. This Annex section is overall well written and very useful. There is, however, one part, which requires some revision. On p. AX2-67 (at the end of the discussion of local-dispersion models), it is stated that (1) emissions from roadways are usually not in steady state and (2) that buoyant plume rise differs for point and line sources. The unwritten implication is that AERMOD, the model used in the exposure and risk assessment, would not be appropriate to simulate NO<sub>2</sub> concentrations from roadways because (1) it uses steady-state assumptions and (2) it is a point source model and does not treat, for example, vehicle-induced turbulence. The text then goes on to discuss CALPUFF, a non-steady-state model that can treat dispersion from surface sources. The implication is that CALPUFF is a better model to simulate NO<sub>2</sub> concentrations downwind of line sources such as roadways. This paragraph needs to be rewritten because (1) the limitations mentioned for AERMOD are not entirely correct and (2) the description of CALPUFF as a better model for line source dispersion is incorrect.

In my earlier comments on the Methods document, I raised some concerns regarding the use of AERMOD for line source modeling and asked why a line source model such as CALINE would not be used. EPA replied that CALINE was no longer supported by the developer and that AERMOD was an acceptable model because it could be adapted for line sources. Although further development of AERMOD to become a true roadway dispersion model is warranted, I agreed with EPA that, given the dispersion models currently available, AERMOD was an appropriate choice. The statement made in the Annex regarding the non-steady-state nature of roadway emissions is not relevant because AERMOD is used to calculate downwind concentrations within a few hundred meters from the roadway and the travel time will be less than one hour (i.e., the time typically used for meteorological and emission inputs). The statement regarding the inappropriate treatment of the initial plume dispersion characteristics for a roadway in AERMOD is misleading as CALPUFF does not treat such roadway initial plume dispersion either. Furthermore, one must note that the current CALPUFF versions that are publicly available (e.g., versions 5.8 and 6) include a coding error that leads to incorrect NO<sub>2</sub> concentrations near the source.

Therefore, the last two sentences of that paragraph (“In contrast, there are models that are non-steady-state...”) should be deleted and the limitations of AERMOD when applied to line sources such as roadways should be rewritten in a more objective light.

Comments from Dr. Elizabeth “Lianne” Sheppard

Preliminary pre-meeting comments:

Charge question 1:

Conceptually the framework outlined in Chapter 1 is on target. However it reads very much like a preliminary incomplete draft. Section 1.6 needs to be thoroughly revised. It has many of the right concepts mentioned, but often briefly, incompletely, or without good justification for the proposed modifications. For instance, Annex AX1 contains a thorough compilation of previous work on evidence classification, but details on the translation into Chapter 1 are absent – there is no evaluation, interpretation, or justification for the modifications proposed on page 1-16 of the ISA. As another example Table 1.6-1 is an adaptation from two previous documents, but the details of and reasons for the adaptation are not discussed in the Chapter or Annex.

Detailed specific comments: To be added.

Charge question 2: Air quality and exposure

The discussion of correlations is much improved although I would like to also see the formulas documented (e.g. in the annex) instead of just described. However, now that the discussion is clearer, it becomes even more questionable in my mind whether the comparisons are useful. Correlations are standardized quantities that depend on multiple features of the data. In a correlation, not only is the linear “relatedness” (covariance) of the two quantities important, but so is the variability of each. Thus two estimates of correlation could be very different just because one is restricted to a single season (with less variability) while the other captures data from an entire year. Since I expect this feature is extraneous to the interpretation goal, should it be part of the comparison? More work is needed to make the comparisons across studies, populations, pollutants, types of correlation, etc. really useful. That said, I was surprised to see an apparently reasonable summary of this information in Chapter 5.

Charge question 3: Integration of evidence about health effects and conclusions regarding health

I found the integration and presentation of evidence in Chapter 3 to be generally good. However there are a number of details in the presentation I think need to be clarified to support the intent of the chapter. It is also critical that staff continue to refine and improve the information presented in the Annex tables. (I recognize this represents a massive effort.) The organization of these tables has been much improved, but I did not find that they include much more pertinent information. In trying to discern my agreement with the interpretations in Chapter 3 I often found I wanted to evaluate more detail on a particular study but this wasn’t available in the annex. I was trying to answer questions such as “did this study properly adjust for confounding”, or “what is the seasonal variation in the data”. There are also some misleading features in the uniform approach to summarization (such as the change in a 4-point symptom scale in the

Chauhan et al (2003) study that appears to be reported as an OR in the appendix and does not include units in Chapter 3 (3-5 lines 21+)). Another important feature of the Chauhan et al study that is discussed in Chapter 3 but completely absent from the Annex summary is that it was conducted over a 13-month period. In looking ahead to future iterations of this ISA process, it is important to consider how to revise the approach. One suggestion is to prepare an annex to the annex with a much more thorough and less constricted summary of each study is available. Quite likely at least a page or more of abstracted information will be needed for each study, at least for the studies that end up providing the greatest weight of evidence for the inference. Finally, both for the current version and in future versions, better indexing and cross-referencing is needed so the supporting information can be found efficiently.

The conclusions brought forward into chapter 5 and summarized clearly in Table 5.3-1 looked generally appropriate. I ask staff and CASAC to consider whether it is worth also listing health outcomes that have not been studied in this table.

Charge question 4:

This appears to be appropriate.

Charge question 5:

Generally the integration and summarization of the evidence in Chapter 5 was quite good, even for cases where I quibble with the details in earlier chapters. The new framework for causality supports the goals of the ISA and provides clear-cut criteria for health endpoints to bring forward into the Risk and Exposure assessment. I would like to discuss whether this draft provides an adequate foundation for preparation of the ERA, particularly with respect to transparent use of the literature for policy.

Other general comments:

- There is still overuse of “statistically significant” in this document. This is binary summarization of the data that depends on multiple features including magnitude of the effect, variability, and sample size. It doesn’t reflect scientific meaning. To the degree possible replace focus on statistical significance with more meaningful quantities (e.g. effect estimates and confidence intervals).
- There continues to be a need to more thoroughly reference supporting information in the annex and make it easier for readers of the ISA to find this information.

#### Comments from Dr. George Thurston

In these pre-meeting comments, I will focus upon responding to my assigned questions for the ISA.

1. What are the views of the Panel on the characterization of the search strategy for

identifying literature, criteria for study selection, the framework for scientific evaluation of studies and causality determination?

In reading the NO<sub>x</sub> ISA document, I found that the epidemiological studies I was aware of were considered, and could see no gaps in the epidemiological literature, which has greatly strengthened the evidence for an association between acute NO<sub>x</sub> exposure and respiratory health effects. With regard to the toxicological studies, I felt that, considering the potential inter-species differences in vulnerability, that it was very appropriate for the document to include studies that used exposures on the order of 5 ppm (approximately 10 mg/m<sup>3</sup>) to be very appropriate in this document, as such studies (while not useful for the estimation of human dose-response estimation) may well provide insights into the potential mechanisms of damage that might be caused by NO<sub>x</sub>.

One area that was lacking was a more intensive consideration of the evidence of potential impact of the co-exposure of particles and NO<sub>x</sub>, both in the toxicology and epidemiology. While I am not as familiar with the toxicological literature, I note that an informative toxicological study by H.G. Boren ["Carbon as a carrier mechanism for irritant gases" *Archives of Environmental Health*, 8, 119-124]. In this paper, a short-term exposure to 47 mg/m<sup>3</sup> (25ppm) of NO<sub>2</sub> or inhalation of fine carbon particles exhibited no gross pathological effects in the mouse lung, but when the mice were exposed to carbon particles that had previously been exposed to NO<sub>2</sub>, the mice developed local destructive lesions, with loss of cells from the alveolar walls. While just one study with high levels of NO<sub>2</sub>, this study indicates that the co-presence of particles with NO<sub>2</sub> can enhance the effects of NO<sub>2</sub>. Since particles are always in co-exposure with NO<sub>2</sub> in the real world, this may provide an important pathway of effect, but one that is not considered by this document, despite my raising this concern early in the process. My question: are there more published toxicological studies considering this particle-NO<sub>x</sub> interaction mechanism? This seems well worth another intensive look through the entire literature with that focus in mind.

3a. To what extent is the discussion and integration of evidence from the animal toxicology and controlled human exposure studies and epidemiologic studies technically sound, appropriately balanced, and clearly communicated??  
?

This is where my above-noted concern really manifests itself. What we have is a wide gap that needs to be bridged between the controlled exposure studies and the epidemiological study results. The former show respiratory effects of NO<sub>2</sub> only down to about 200 ppb, while the latter routinely and robustly document significant NO<sub>2</sub>-respiratory associations down at ambient levels. How can that be the case? We previously confronted a similar situation in the 1980's, when our NYU-Harvard study's of children at summer camps documented significant lung function decrements among children to be associated with ozone exposures below 100 ppb, while the controlled exposure studies only showed effects down to 120 ppb. Subsequent more realistic controlled exposure studies (with exercise) later confirmed the epidemiology, and we now have a more protective ozone standard.

With regard to NO<sub>2</sub>, I suspect that ambient particles, always present in epidemiological studies, but not present in controlled NO<sub>x</sub> exposure studies, may provide the vector for the apparently enhanced effects of NO<sub>x</sub> in epidemiology vs. controlled studies, but the evidence for this possible avenue to justify the apparent discrepancy between the epidemiology and the controlled-exposure studies is not sufficiently explored in this report. In some cases it is noted in the ISA already (e.g., the fact noted that the APHEA study found greater PM effects in cities with higher NO<sub>2</sub> levels), but needs to be brought together to address this specific issue. Thus, while some of the evidence is already present, and more may be in literature not yet brought to bear (or not yet collected), it is important to identify in the ISA what we do and do not know about this potential mechanism of NO<sub>x</sub> effects, and about other possible factors that may be responsible for this apparent disparity between the levels of effects (e.g., that the most susceptible subjects may not be considered in controlled-exposure studies, or that exercise may be a factor, both of which are mentioned here and there in the ISA).

The way to bring this about in the document, I suggest, is to add a section to Chapter 5 where the questions asked at the start of the Integrative Summary (on page 5-1) are answered to the best we can at this time, and which identify areas of further needed investigation to answer the question more definitively. In particular, the most important question to be answered is:

- At what levels of nitrogen oxides exposure do health effects of concern occur?

Answering this particular question will expedite the addressing of the gap between the controlled-exposure results and the epidemiology results, as well as a comprehensive consideration as to why that might be (i.e., Particle-NO<sub>x</sub> interactions? Greater degrees of susceptibility in the general public? Exercise? etc.).

3b. What are the views of the Panel on the conclusions drawn in the draft ISA regarding the strength, consistency, coherence and plausibility of NO<sub>2</sub>-related health effects

While I generally agree with judgments reached by the EPA regarding the strength of the evidence regarding causality, there is a need to discuss the concentrations where the various effects are applicable, as discussed above.

Comments from Dr. Ronald Wyzga

Overall comments:

This draft is much improved over the previous version. The siting of monitors is discussed, and there is a much better understanding of reported ambient measurements represent. The review of the health literature is comprehensive and makes it easier to achieve a good overall understanding of the health consequences of NO<sub>x</sub> exposure. The summary of the health effects discussed in Chapter 5 is an excellent organizational tool that facilitates an understanding of the nature and consequences of ambient exposures to oxides of nitrogen. There are a few areas that need further clarification. These are presented below in more detailed comments.

Charge question 2: To what extent are the atmospheric chemistry and air quality characterizations clearly communicated, appropriately characterized, and relevant to the review of the primary NO<sub>2</sub> NAAQS? Are the properties of ambient oxides of nitrogen appropriately characterized, including spatial and temporal patterns and relationships between ambient oxides of nitrogen and human exposure? Does the information in Chapter 2 provide a sufficient atmospheric science and exposure basis for the evaluation of human health effects presented in alter chapters?

I have two suggestions here. The document refers to several studies undertaken overseas, several of which focus on distance to roadways as a factor “affecting indoor and outdoor NO<sub>2</sub> concentration and personal exposure”. To the extent that the overall nature of exposure could be quite different from that in the US given differences between the US and overseas sources in terms of fleet composition and extent of pollution control, there should be some note made about the geographic setting of these studies and whether the setting is typical of those found in the contemporary US. Any information on co-pollutants and concentrations would be particularly helpful.

Since the strength of the health argument for NO<sub>2</sub> health effects is tied to the observations that effects are found in studies which consider both indoor and outdoor exposures, it would be particularly interesting for this chapter to provide some additional information to inform this argument. For example, given the typical sources of indoor and outdoor NO<sub>x</sub>, how does the composition change with respect to the different oxides of nitrogen and are the co-pollutants the same or different in indoor and outdoor settings?

Charge question 3: To what extent is the discussion and integration of evidence from the animal toxicology and controlled human exposure studies and epidemiologic studies technically sound, appropriately balanced, and clearly communicated? What are the views of the Panel on the conclusions drawn in the draft ISA regarding the strength, consistency, coherence, and plausibility of NO<sub>2</sub>-related health effects?

One of the difficulties in interpreting the epidemiological study results is that it is often unclear whether health responses are due to traffic or to NO<sub>x</sub> per se, an ingredient of traffic-generated air pollution. To the extent possible, the document should attempt to indicate those studies where traffic is deemed to have a lesser influence on NO<sub>x</sub> exposure.

I would like to see a more rigorous examination/discussion of the co-pollutant issue; for example, which co-pollutants were discussed in which study; which were not. Are there any differences in measurement error, etc. I personally share some of the concerns raised by Brook et al. (2007) cited in the document.

In several places studies were not considered because they “did not inform”; this needs to be clarified.

Charge question 5: What are the Panel's views on the adequacy of this external review draft ISA to provide support for future exposure and policy assessments?

The current draft is a helpful document and a great improvement over the previous draft. With the appropriate consideration of the issues raised elsewhere in this review, the document would be an excellent resource for future exposure and policy assessments. I also believe that the document needs to tackle the issue of whether associations between NO<sub>2</sub> and health responses in epidemiological studies are a reflection of NO<sub>2</sub> exposures per se or is NO<sub>2</sub> an index surrogate for some other exposure associated with NO<sub>2</sub> sources. I believe the information is scattered throughout the document to help address this question, but an explicit and articulate consideration of this issue would greatly improve the document.

Specific comments:

Page 2-21: Figure 2.5-1 "residence"

Page 2-29, ll 1-3: Does this mean that local sources and near-source concentrations are not to be regulated? I think the wording needs to be changed here.

Page 2-42, ll. 5-6: statistical significance per se is not as informative as R<sup>2</sup>; if the sample size is large enough any non-zero correlation will be statistically significant.

Page 2-52, Table 2.5-7: The paper by Brook et al. (2007) cited later in the document should be mentioned here. I find it particularly noteworthy that NO<sub>2</sub> is highly associated with several organic compounds.

Page 3-12. Figure 3.1-1: This figure is helpful, but it should also indicate the time and concentrations of exposure of exposure in the subtable. See Figure 3.1-2. This would allow the reader to judge whether the application of Haber's law is appropriate and could facilitate the interpretation of results.

p. 3-26, ll 28- : Are there any co-pollutants associated with these exposures? Are they different from the outdoor studies?

pp. 3-51;3-52, Figures 3.1-10; 3.1-11: Where is Peel et al (cited elsewhere)?

p. 3-57, ll. 9-14: clarify what is meant by "did not inform"; if a study is not considered, it is important to understand why. Similar comments exist elsewhere; e.g., p. 3-58, ll 17-20.

