**Comments from the editors**

We thank the Editors and Reviewers for their thoughtful and constructive suggestions. We have revised the manuscript in response to their comments, as detailed below.

All page/line/reference numbers refer to the clean revised manuscript.

**Please take particular note of reviewer comments below on justifying and describing the use of the Bayesian hierarchical model, on determination and clarification of confounder control, and the lack of control for potentially important factors such as BMI, smoking, and indoor air quality.**

We have addressed this, as detailed throughout relevant responses to Reviewers below. In particular, please see responses to Major Comments 4, 6, 10, 15 and Minor Comments 3, 5, 8 for Reviewer 1, and responses to Major Comments 1, 2, 4, 6, 7, 8, 9, 10, 12 and Minor Comments 14, 18, 19, 21, 22 for Reviewer 2.

**Second, please ensure that the summary of main findings are fully supported by the results, e.g., take into account and explain sensitivity analyses in eFigure 1 that show weaker single-pollutant estimates for EC than those from the main models.**

Our main model demonstrated that there was a potentially positive association between EC and ALS diagnosis, which contained traffic-related pollutants (EC, NOx, CO) and non-EC PM2.5, as well as adjusting implicitly by matching cases and controls on age, sex, year of birth, vital status, and explicitly by including terms in the model for SES, civil status, residence, place of birth.

Because the main models contain more terms which are highly correlated, it was not surprising that the estimated standard errors would be larger than single-pollutant models. Furthermore, because of the high correlations among traffic-related pollutants, the point estimates are also expected to somewhat vary between analyses. However, because of the consistency of the sign of the EC association throughout (i.e., whether in single-pollutant or multi-pollutant models), we have suggested that EC may be a driver of the relationship in the revised manuscript (P. 14, Lines 298-309):

*We found that EC had the largest-in-magnitude independent association with ALS diagnosis, while associations with NOx and CO were negative with credible intervals overlapping the null, and smaller in magnitude. Sensitivity analyses demonstrated that for single pollutant models, the association for EC was smaller than for our main multi-pollutant model, which took into account the variance-covariance structure of traffic-related pollutants. Overall conclusions for the association between EC and ALS diagnosis would have been similar from the single- or multi-pollutant models. The inconsistent associations for NOx and CO in the multi- and single-pollutant models suggest that the model may have had limited success identifying each individual pollutant’s association with ALS diagnosis due to the high level of collinearity of traffic-related pollutants. Nevertheless, the consistency of the sign of the central estimate of EC in all models suggests that EC may be a driver of the ALS and traffic-related pollutant association, though further analysis is required.*

**Finally, please ensure consistency in results presentation throughout the abstract and main text; e.g., the abstract presents posterior probabilities that are not included in the results section.**

We have ensured that there is consistency in results presentation throughout the abstract and main text, e.g., (P. 13, Lines 273-278):

*For 5-year average pollutant concentrations, we observed the largest overall association for the individual SD increase in EC (11.5%; 95% CrI: -1.0%, 25.6% per 0.42 µg/m3; 96.3% posterior probability of positive association) (Figure 2). SD increases were associated with a decrease in odds of ALS diagnosis in NOx (-4.6%; 95% CrI: -18.1%, 8.9% per 20 µg/m3; 27.8% posterior probability of positive association) and CO (-3.2%; 95% CrI: -14.4%, 10.0% per 106 µg/m3; 26.7% posterior probability of positive association).*

**Comments from reviewers**

**Reviewer #1: In this manuscript, the authors investigate the association between long-term traffic related air pollution and amyotrophic lateral sclerosis (ALS) in Denmark. The authors used Bayesian hierarchical modeling to estimate joint and individual effects of traffic related air pollutants (NOx, CO, EC) on odds of developing ALS. The major finding of this paper is that 5-year average EC concentration was individually associated with ALS. The study found no overall or joint association of traffic related pollution with ALS. This is an important topic and using methods to estimate joint effects of correlated pollutants is a necessary next step in air pollution studies. This paper was well done, though authors should explain certain analysis choices more clearly and provide more detailed results and discussion.**

We thank the Reviewer for the thoughtful and constructive suggestions. We have responded point-by-point to the Reviewer’s questions and comments below.

**Major Comments**

**1) Methods, page 5, line 46: Why did authors only include patients that were at least 20 years old at diagnosis? This choice needs to be explained/motivated.**

Cases in younger patients (i.e., younger than 20 years old) would have been at a much greater chance of misclassification, with a very high likelihood that a case identified in such a young person is an error in diagnosis coding (Trabjerg et al. 2020). Further, juvenile ALS cases have been explained to much larger degree by genetic mutations (Mathis et al. 2019). We have added to the explanation of the 20-year-old limit in the revised manuscript (P. 4, Lines 85-88):

*We only included patients who were at least 20 years old when diagnosed because (i) cases younger than 20 years old were at a greater chance of misclassification, since ALS has been predominantly diagnosed in older adults,46 and (ii) the very few juvenile ALS cases have been explained to a much larger degree by genetic mutations (~40%).47*

**2) Methods, page 6, line 51: Please include more detail about the spatio-temporal air pollution modeling system.**

We have now added more detail about the spatio-temporal air pollution modelling system in revised manuscript (PP. 5-6, Lines 111-120):

*We obtained predictions on monthly concentrations of nitrogen oxides (NOx), carbon monoxide (CO), elemental carbon (EC), and fine particles (PM2.5) (as well as ozone, O3, for a sensitivity analysis, usually negatively correlated with other pollutants due to its chemistry51), at residential addresses of study participants from the validated spatio-temporal air pollution modelling system DEHM-UBM-AirGIS that provides full space and time coverage over the study period, described in detail elsewhere.52–55 In brief, DEHM-UBM-AirGIS is a human exposure modelling system for traffic pollution, developed for application in Danish air pollution epidemiological studies. The modelling system is able to generate street configuration and traffic data based on digital maps and national databases, which enables estimation of air quality levels at a large number of addresses in an automatic and effective way.*

We have additionally provided metrics of predictive accuracy for each of the used pollutants in the revised manuscript (P. 6, Lines 121-124):

*The models have good predictive accuracy, with average monthly correlations between measured and modelled results of 0.85 for NOx, 0.91 for CO, 0.92 for O3, 0.79 for EC, and 0.83 for annual concentrations of PM2.5.52,55*

**3) Methods, page 7, line 17: Please provide more details on how the 1-, 5-, and 10-year averages were created for air pollution exposures. Was a weighted average created based on how long the participant lived in one location?**

We have given more details on how the 1-, 5-, and 10-year averages were created for the air pollution exposures in the revised manuscript (P. 6, Lines 129-133):

*Based on the residential history of each case or control, we calculated 1-, 5-, and 10-year average exposure to each pollutant ending at one year before the index date, as diagnosis has been shown previously to occur at a median of 12 months after symptoms onset.60 Specifically, each case or control average value (1-, 5- or 10-year) was calculated as the mean of all concentrations recorded across time at the recorded addresses within each time window.*

**4) Methods, page 7, line 41: How did the authors determine confounding variables?**

We matched cases and controls on age, sex, year of birth, and vital status, as ALS prevalence varies according to these characteristics. We also accounted for socioeconomic status (SES), civil status, last reported place of residence, and place of birth. SES influences many lifestyle factors, such as obesity, and is associated with ALS diagnosis in Denmark (Dickerson et al. 2018). Civil status was included due to the influence that a spouse has on visiting a family physician (Bucher et al. 2019). Last reported place of residence was included to account for various local environmental and behavioral stressors, such heavy metals, which may have an influence on ALS prevalence (Oskarsson, Horton, and Mitsumoto 2015). Place of birth was included to account for the variety of childhood exposures, which vary by location, which may have an impact on the probability of developing ALS (Norman et al. 2013). Ultimately, we were limited by what was available in the Danish Civil Registration System.

We have added these details to the revised manuscript (PP. 7-8, Lines 141-159):

*We included a set of covariates based on as close as possible to index date to account for potential confounding bias, including household socioeconomic status (SES) based on last-reported job title at index date; civil status at index date, last reported place of residence at index date, and place of birth. We used a five-category individual-level SES definition developed by the Danish Institute of Social Sciences, based on job titles from income tax forms, which has been associated with ALS diagnosis in Denmark,61 as well as how quickly one is identified as having ALS in the Danish Civil Registration System.62 Group 1 (highest status) includes corporate managers and academics; group 2: proprietors, managers of small businesses and teachers; group 3: technicians and nurses; group 4: skilled workers; and group 5: unspecialized workers, such as entry-level positions within food and retail environments. We also included a group for participants whose job title was unknown (group 9). For each married participant, we used the higher of the couple’s individual SES categories, when available. We also used information on civil status (never married, married, divorced, widowed) due to the influence that a spouse may have on visiting a family physician,63 last reported place of residence from postcode (Greater Copenhagen, big cities of Denmark, rest of Denmark, Greenland) to account for various local environmental and behavioral stressors,7 and place of birth (Greater Copenhagen, big cities of Denmark, rest of Denmark, Greenland, foreign, unknown) to adjust for other potential family-specific, location-specific, and early-life confounders, which may have an impact on the probability of developing ALS.64 Ultimately, we were limited by what was available in the Danish Civil Registration System.62*

**5) Methods, page 8, line 50: Please include more motivation and reasoning for ozone sensitivity analysis.**

There is evidence from other studies that ozone concentrations are associated with many different adverse health outcomes (Nuvolone, Petri, and Voller 2018). However, ozone is highly negatively correlated with traffic-related air pollutants, which is also the case in our study, as can be seen in Figure 1 in the main manuscript. Adding an extra pollutant which is highly-correlated with the others may have posed challenges to the statistical inference of the model, which is why we included it in the sensitivity analysis rather than the main analysis.

Nevertheless, our sensitivity analyses demonstrated that inclusion of ozone did not noticeably change our results (eFigure 1). Our conclusions from the main analysis, thus, would not have changed had we included ozone.

We have clarified why we included ozone in the sensitivity analysis in the revised manuscript (P. 9, Lines 197-199):

*In a sensitivity analysis, we included O3 in the model, as O3 concentrations have been associated with many adverse health outcomes,69 and were negatively correlated with traffic-related pollutants […]*

**6) Methods, page 9-10: When discussing priors used for the Bayesian model why are weakly-informative priors given to non-EC PM2.5, but non-informative priors are given to other parameters? Please give more detail when justifying use of priors.**

We have edited the description of priors used in the Bayesian model to clarify that weakly-informative priors were given to all of the pollutant coefficients, with the other pollutant coefficients (i.e., not that of non-EC PM2.5) given the same prior via the hierarchical structure of the traffic-related pollutant concentrations. The revised description can be found in the revised manuscript (P. 10, Lines 220-223):

*We used weakly-informative priors so that data drove parameter estimation. Hyper-priors for coefficients on non-EC and covariates were N(0,10); for and we used Half-Cauchy(0,10), as recommended by Gelman, Polson and Scott as a weakly-informative prior;71,72 was defined by the weakly-informative prior LKJCorr(1).73*

**7) Results, page 11: Why are only 5-year average exposures presented in the results section? Please provide more justification or present the 1- and 10-year data as well in the main tables/figures.**

We provide the 5-year average exposures as the main results here because a 5-year average exposure is a balance between most recent exposure as well as long-term concentration. We have clarified this by adding more justification in the revised manuscript (P. 12, Lines 262-263):

*For the main results, we present 5-year average exposure associations as a balance between representation of most recent exposure as well as long-term concentration.*

**8) Results, page 12: Please provide more discussion of the protective effect of NOX and CO. How does this effect the null joint effect of NOX, CO, and EC?**

While the point estimates of the association of ALS and NOx and CO are less than 0% change, the credible intervals widely overlap with the null. Nevertheless, we were also surprised by this result. We discuss further why this might be in the Discussion of the revised manuscript (P. 14, Lines 303-309):

*Overall conclusions for the association between EC and ALS diagnosis would have been similar from the single- or multi-pollutant models. The inconsistent associations for NOx and CO in the multi- and single-pollutant models suggest that the model may have had limited success identifying each individual pollutant’s association with ALS diagnosis due to the high level of collinearity of traffic-related pollutants. Nevertheless, the consistency of the sign of the central estimate of EC in all models suggests that EC may be a driver of the ALS and traffic-related pollutant association, though further analysis is required.*

**9) Results, page 12, line 46: Please provide correlation coefficient for ozone and other pollutants.**

We now provide the correlation coefficient for ozone and the other pollutants in the revised manuscript (P. 13, Lines 270-271):

*O3 was negatively correlated with other pollutants, ranging from -0.54 to -0.89.*

**10) Results, page 13, line 9: Can the authors also provide posterior probability for the null?**

The posterior probability is the amount of the marginal posterior distribution estimate of the coefficient of interest which is above the null. Therefore a 50% probability means it is as likely as not that the marginal estimate is null. We clarify this in the text with a further exposition of what various posterior probabilities mean (P. 11, Lines 237-241):

*To calculate the probability that an association estimate was greater than null, we used the 4,000 samples of the posterior distribution and took the proportion of samples which were above the null. A 50% probability means that it is as likely as not that the marginal estimate is null, a probability closer to 100% indicates that the association is more likely to be truly positive, with closer to 0% indicating more likely to be truly negative.*

**11) Results page 13, line 22: "(eFigure 1) resulted in positive associations for each of EC, NOx, CO, with positive associations for non-EC PM2.5 in all but the model with EC." Is eFigure 1 the correct figure? eFigure 1 shows protective effects for all but EC.**

We have clarified that we are only referring to the single-pollutant models D, E and F in eFigure 1 in the revised manuscript (P. 13, Lines 284-287):

*Single-pollutant models for each traffic-related pollutant adjusting for non-EC PM2.5 (eFigure 1; single traffic-related pollutant models D, E and F) resulted in positive associations for each of EC, NOx, CO, with positive associations for non-EC PM2.5 in all but the model with EC.*

**12) Discussion, page 13, line 38: Authors state that they found an average increase in concentration of traffic-related pollutants was associated with and increase in odds of ALS. Though only EC showed a positive association and joint effect was null?**

We have clarified that we are referring to the joint association here (P. 14, Lines 296-298):

*In the largest case-control study of ALS and traffic-related air pollution to date, we found that a joint increase in average concentrations of traffic-related pollutants was potentially associated with an increase in odds of ALS diagnosis, with the clearest results for EC.*

**13) Discussion, page 14, line 48: If EC and NOX are so highly correlated why are their associations with ALS so different?**

While EC and NOx are highly-correlated (0.94-0.96), they are not perfectly correlated. Our Bayesian hierarchical model structure allowed us to incorporate their high correlation while also leveraging the differences to make inferences about their single-pollutant associations with ALS diagnosis. With this high collinearity in mind, however, it is reassuring that single-pollutant model associations for EC and NOx (eFigure 1; Models D and F) give similar results. We discuss that while the EC results were consistent, the NOx results were not in the revised manuscript, indicating that while this would indicate that EC is potentially driving the association, further work is necessary (P. 14, Lines 304-309):

*The inconsistent associations for NOx and CO in the multi- and single-pollutant models suggest that the model may have had limited success identifying each individual pollutant’s association with ALS diagnosis due to the high level of collinearity of traffic-related pollutants. Nevertheless, the consistency of the sign of the central estimate of EC in all models suggests that EC may be a driver of the ALS and traffic-related pollutant association, though further analysis is required.*

**14) Discussion, page 14, line 58: the 1-year estimate may be the most robust to exposure misclassification, provide more justification for why it may be the most relevant exposure window.**

We understand why the Reviewer suggests that the 1-year estimate may be the most robust to exposure misclassification. However, for every case and control, we have addresses geocoded over time for each year during our study period. This would minimize the potential for exposure misclassification for earlier years in the study period compared with later years. Therefore, we do not think that the 1-year estimate may necessarily be the most robust to exposure misclassification.

We have added more justification for why the 1-year average concentrations may be the most relevant exposure window in the revised manuscript (PP. 15-16, Lines 333-339):

*EC exposure was more strongly associated with 1-year than for 5-/10-year average concentrations, which may indicate that the previous year of exposure may be the most relevant exposure window relevant to traffic-related exposures and ALS; this is biologically plausible, as this critical exposure window would be at the pre-symptomatic stage of underlying ALS progression, where traffic-related pollution exposure may add to the ongoing cellular or molecular process of the disease, to the point where the body can no longer compensate and subsequently enters the clinical phase.82–84*

**15) Please include justification for use of the Bayesian hierarchical model, as opposed to other mixture modeling methods (Bayesian kernel machine regression, etc.) that are more established.**

Bayesian Kernel Machine Regression is not currently appropriate for matched case-control studies. Further, we required a Bayesian model to be able to take into account variance-covariance structure of the traffic-related pollutants, as well as being able to optimally capture uncertainty. We have clarified this in the future research paragraph in the revised manuscript (P. 17, Lines 364-367):

*Other mixture model methods, such as Bayesian Kernel Machine Regression (BKMR)88 might be useful in further exploring the robustness of joint associations in a different framework, though BKMR was not appropriate for our particular research question, since BKMR is not currently appropriate for case-control studies.*

**16) Figure 1. Please provide figure for 1-, and 10-year exposure estimates.**

On Figure 1, we now provide Spearman correlations for 1-, 5-, and 10-year pollutant concentrations, as seen below:

****

**Figure 1**. Spearmancorrelation of 1,- 5-, and 10-year average pollutant concentrations.

**17) Table 1. SES group 9, why were unemployed and unclassified grouped together?**

This should have been stated as ‘unknown’ here and we have corrected this in the revised manuscript (P. 7, Lines 150-151):

*We also included a group for participants whose job title was unknown (group 9).*

**Minor Comments**

**1) Throughout the paper there is inconsistent use of the abbreviation SD for standard deviation.**

We have now introduced the abbreviation for the term standard deviation (SD), which is also in the list of Abbreviations. We now consistently use this throughout the revised manuscript e.g., (P. 13, Lines 273-275):

*For 5-year average pollutant concentrations, we observed the largest overall association for the individual SD increase in EC (11.5%; 95% CrI: -1.0%, 25.6% per 0.42 µg/m3; 96.3% posterior probability of positive association) (Figure 2).*

**2) Introduction, page 4, line 26: Unclear why this sentence is a contradiction.**

We have added the word ‘also’ in the subclause of the sentence to demonstrate that not only should air pollution be studied in relation to respiratory- and cardiovascular-related outcomes, but also with nervous system-related outcomes and neurodegeneration in the revised manuscript (P. 3, Lines 53-56):

*Although air pollution is commonly studied in association with respiratory- and cardiovascular-related outcomes, e.g., references 9–14, epidemiologic and toxicological studies also support several plausible biological mechanisms in association with the nervous system and neurodegeneration, e.g., references 15–34.*

**3) Methods, page 8 line 42: Authors say non-EC PM2.5 adjust for other air pollutants from other sources. This would only adjust for PM2.5 from other sources.**

This Reviewer is correct. We have also clarified that PM2.5 is also an indicator of overall air pollution mixture, and that adjusting by PM2.5 will also indicate whether pollution from other sources not explicitly quantified might also have associations with ALS. We have clarified this in the revised manuscript (P. 9, Lines 192-194):

*If other sources of air pollution are associated with ALS, then including non-EC PM2.5 adjusts for PM2.5 from other sources,67 as well as indicating whether pollution from other sources not explicitly quantified might also have associations with ALS.*

**4) Methods, page 9, line 16: Model quantifies log-odds of one standard deviation increase. Please add explanation of why you chose one standard deviation increase instead of interquartile range.**

Presenting effect estimates for a standardized increase in pollutant concentrations is preferrable when examining multiple pollutants and interested in the overall effect, because we combined concentration associations via the traffic terms in the model. Both standard deviation (SD) and interquartile range (IQR) are measures of the spread of values, which can be equivalently used to present effect estimates for an increase in standardized pollutant concentrations. There is no inherent benefit to picking one or the other in this case, as the role of dividing by both measures of spread is to normalize concentrations. We have added that both are equivalent ways of standardizing in the revised manuscript (P. 9, Lines 187-192):

*, , , the pollutant-specific coefficients (log-odds) per standard deviation (SD) increase in concentration of , , , respectively, scaled by their respective SDs and centered at their means, with each a pollutant-specific association adjusted by other terms in the model and the rest as coefficients for subject-specific covariates. Interquartile Range (IQR) could equivalently be used to scale pollutant concentrations.*

**5) Results, page 12, line 4: Joint association of which pollutants?**

We have clarified which pollutants in the revised manuscript (P. 13, Lines 278-280):

*The joint association of traffic-related pollutants (EC, NOx, CO) was 2.3% (95% CrI: -3.3%, 7.7%), with an 77.8% posterior probability of a positive association.*

**6) Results, page 13, line 17: "10-year average exposure results were attenuated versions of the 1- and 5-year results." Wording could be improved.**

We have expanded upon this phrase to be more descriptive in the revised manuscript (P. 13, Lines 283-284):

*Compared to the 1- and 5-year results, the 10-year average exposure results were attenuated, as associations tended further to the null.*

**7) Results overall: eFigure 2 is not mentioned.**

We now mention eFigure 2 in the revised manuscript (PP. 13-14, Lines 291-293):

*A map of average concentration of included pollutants (NOx, EC, PM2.5, CO, O3) across Denmark for a representative year (2000; middle of study period 1989-2013) is also available in eFigure 2.*

**8) Discussion, page 14, lines 29-39: If BMI is not a confounder this is unnecessary to include.**

We have deleted these lines in the revised manuscript.

**9) Figure 2. Please make the point estimate dots bigger.**

We have made these points bigger in Figure 2 in the revised manuscript, copied below for convenience:



**Figure 2.** Percentage change in odds of ALS diagnosis per 1-, 5- and 10-year average standard deviation (SD) increase for each pollutant. Results are from the Bayesian hierarchical model including each of EC, NOx, CO, and non-EC PM2.5 together, and were additionally adjusted by age, sex, year of birth, vital status, socioeconomic status, civil status, last reported place of residence, and place of birth.

**Reviewer #2: Comments pasted below. The uploaded review contains a figure.**

**This paper describes a study of the effect of traffic-related air pollution exposures on amyotrophic lateral sclerosis (ALS). The study, set in a very large healthcare administrative database in Denmark, has several strengths, including exposure assessments that span decades and an attempt to account for a latent period between disease onset and clinical diagnosis. The report lacks clarity and detail on important aspects of the investigation, encompassing the target estimands, validity of the ALS measurement, and sources of confounding. My concerns follow:**

We thank the Reviewer for the thoughtful and constructive suggestions. We have responded point-by-point to the Reviewer’s questions and comments below.

**MAJOR  
  
1. The interpretive distinction between the 3 estimands pursued in this study was unclear. E.g., from the abstract, "… the overall and joint association for the three traffic-related pollutants (NOx, CO, and EC), as well as pollutant-specific associations." What does "overall" mean exactly, and how does "overall" differ from "joint"? Does "joint" include interactions? Does "pollutant-specific" reflect adjustment for other pollutants? For example, how would you express the parameter estimate from each of these in words that are true to the underlying mathematics?**

We distinguished between the “joint” association of the three pollutants (i.e., total percentage change in odds of ALS diagnosis with increase in each of EC, NOx, CO), and “overall” association of the three pollutants (i.e., average percentage change in odds of ALS diagnosis from each of EC, NOx, CO). We have stated this in the revised manuscript (P. 8, Lines 171-176):

*We employed a Bayesian hierarchical formulation because it enables estimates of (a) independent pollutant-outcome associations, (b) a joint association of the three pollutants (i.e., total percentage change in odds of ALS diagnosis with increase in each of EC, NOx, CO), and (c) an overall average traffic association (i.e., average percentage change in odds of ALS diagnosis from each of EC, NOx, CO), while accounting for the variance-covariance structure between the highly-correlated exposures and their coefficients.66*

The Reviewer is correct that the “pollutant-specific” associations reflect adjustment not only for other pollutants, but also for other covariates to account for potential confounding bias, described in more detail in the revised manuscript (P. 9, Lines 187-191):

*, , , the pollutant-specific coefficients (log-odds) per standard deviation (SD) increase in concentration of , , , respectively, scaled by their respective SDs and centered at their means, with each a pollutant-specific association adjusted by other terms in the model and the rest as coefficients for subject-specific covariates. […]*

**2. Related to #1, the joint association is described in these terms: "This sum quantifies the association (log-odds) with ALS of a one-SD increase in the three pollutants simultaneously." Although it is mathematically possible to compute this, how well does an increment of SD in all 3 pollutants match up with the joint distribution of these pollutants in the population? Do the concentrations vary at about the same pace? i.e., can you identify locations (or location-periods) that are 1-SD apart (or 0.5 or 0.1 SD apart, etc.) on all 3 pollutants?**

As described in the revised manuscript, the pollutants are highly correlated (PP. 12-13, Lines 265-271):

*The 5-year average traffic-related pollutant concentrations were 27 µg/m3 for NOx (SD=20 µg/m3), 238 µg/m3 for CO (SD=106 µg/m3) and 0.85 µg/m3 for EC (SD=0.42 µg/m3) (Table 2). Figure 1 shows Spearman correlations between pollutants for 1-, 5-, and 10-year average exposures. Traffic-related pollutants (NOx, CO, EC) were highly correlated in cases, controls and overall, ranging from correlations of 0.91 to 0.96. Otherwise, non-EC PM2.5 was most highly correlated with CO, ranging from 0.67 to 0.7. O3 was negatively correlated with other pollutants, ranging from -0.54 to -0.89.*

This would imply that the increase in one pollutant by 1-SD would result in a similar relative increase in the other related pollutants. Therefore, each of the three traffic-related pollutants each increasing simultaneously by 1-SD is a physically-plausible scenario.

**3. Introduction: the literature review seemed cursory (e.g., "… epidemiologic and toxicological studies support several plausible biological mechanisms in association with the nervous system and neurodegeneration.15-34"). I recommend citing systematic reviews (e.g., Integrated Science Assessments from the US EPA) or using "e.g." before some of the citations.**

As per the Reviewer’s suggestion we have added “e.g.” before some of the example citations in the Introduction here (P. 3, Lines 53-60):

*Although air pollution is commonly studied in association with respiratory- and cardiovascular-related outcomes, e.g., references 9–14, epidemiologic and toxicological studies also support several plausible biological mechanisms in association with the nervous system and neurodegeneration, e.g., references 15–34. Ambient air pollution, especially urban air pollution, is a ubiquitous exposure that has been associated with several other neurodegenerative disorders, e.g., references 16–21,35,36. and is consistently linked to systemic inflammation,22–24 oxidative stress,25–28 and neuroinflammation,15,29 all of which, in turn, have been reported as key pathways to ALS pathogenesis, e.g., references 30–34.*

**4. Introduction/Methods: the use of ozone in this investigation was confusing. The Introduction states, "Using three air pollutants commonly used in health studies as traffic-related emissions tracers—nitrogen oxides (NOx), carbon monoxide (CO), and elemental carbon (EC)— as well as fine particles (PM2.5) and ozone (O3), we aimed to assess whether exposure to (a) each individual air pollutant is independently associated with ALS diagnosis …," which loosely implied that ozone was a traffic-related pollutant of interest. Later, in the Methods, O3 is described as being part of "a sensitivity analysis, usually negatively correlated with other pollutants due to its chemistry." This requires more explanation. Was exposure to O3 not expected to be relevant to ALS? Are the predicted O3 concentrations inaccurate?**

Our study aim was to investigate whether each traffic-related air pollutant (EC, NOx, CO), individually, jointly and overall, was associated with ALS diagnosis, while also fully adjusting for other pollutants (PM2.5 from other sources and O3) and other relevant covariates. To clarify that our main focus was the traffic-related pollutants, we have edited the last paragraph of the Introduction in the revised manuscript (P. 4, Lines 68-72):

*Using three air pollutants commonly used in health studies as traffic-related emissions tracers—nitrogen oxides (NOx), carbon monoxide (CO), and elemental carbon (EC)— we aimed to assess whether exposure to (a) each individual air pollutant is independently associated with ALS diagnosis, and estimate their (b) joint and (c) overall traffic-related emissions associations.*

While there is no existing evidence, to our knowledge, to assess whether O3 concentrations may impact ALS diagnosis, we included a sensitivity analysis to examine whether including it made a difference to the main set of results, which it did not. We explain this in the revised manuscript (P. 9, Lines 197-198):

*In a sensitivity analysis, we included O3 in the model, as O3 concentrations have been associated with many adverse health outcomes,69 […]*

The models we have used for air pollution prediction, DEHM-UBM-AirGIS, have good predictive accuracy for all pollutants, including predicted concentrations of O3, with average monthly correlations between measured and modelled results quoted in the revised manuscript (P. 13, Lines 289-291):

*Results from variations of the main model in the sensitivity analyses were robust to prior choices, inclusion of O3, and inclusion of parish-level SES (eFigure 1).*

**5. Methods: The validity of the registry for identifying ALS cases requires more detail, as it is a fundamental aspect of this investigation. Was the validation against an in-person assessment? a neurologist's in-depth review of medical records? Did the validation compare date of diagnosis with the date of symptom onset? What were the quantified indices of accuracy? As appropriate, it could be useful to mention the potential influence of outcome misclassification (or lack thereof) on the findings and mention determinants of misclassification if known.**

We had claimed the validity of obtaining ALS diagnosis data from the Danish National Patient Register based on previous work from some of our co-authors, including the senior author of this current study (Kioumourtzoglou et al. 2015). In that work, a specialist ALS neurologist reviewed medical records of registry and mortality data for factors that may have been slightly related to agreement of ALS diagnosis. This previous study found that the use of hospital discharge and death certificate records are a valuable and highly reliable tool for ALS epidemiologic studies such as our current one. We have added detail to the revised manuscript (PP. 4-5, Lines 88-91):

*In our validation study, Register data for ALS ascertainment were highly reliable; working with a specialist ALS neurologist to review medical records and comparing to death certificates and hospital discharges, the Danish National Patient Register was found to have an overall predictive value for ALS of 82%.48*

We have added to the Limitations in the Discussion that while the Danish National Patient Register was highly reliable for ALS ascertainment, outcome misclassification may still have been possible (P. 16, Lines 356-357):

*While a previous study found that ALS ascertainment from the Danish National Patient Register was highly reliable,48 outcome misclassification cannot be ruled out, […]*

While the studied quoted above did not examine date of system onset compared with date of diagnosis, another study in Ireland found that diagnosis of ALS occurs at a median of 12 months after symptoms onset (Galvin et al. 2017), which we have stipulated in the revised manuscript (P. 6, Lines 129-131):

*Based on the residential history of each case or control, we calculated 1-, 5-, and 10-year average exposure to each pollutant ending at one year before the index date, as diagnosis has been shown previously to occur at a median of 12 months after symptoms onset.60*

Nevertheless, we have added that this may also be a limitation of the work in the Discussion of the revised manuscript (P. 16, Lines 356-358):

*While a previous study found that ALS ascertainment from the Danish National Patient Register was highly reliable,48 outcome misclassification cannot be ruled out, nor can the possibility that date of diagnosis and symptom onset were irregularly aligned.*

**6. Methods: "… we removed the EC concentration from the total PM2.5 mass concentration …." How did you "remove" EC? Subtraction? Using residuals from a regression model?**

Our pollutant model for PM2.5, DEHM-UBM-AirGIS (Khan et al. 2019; Jørgen Brandt et al. 2001; J Brandt et al. 2003; Frohn et al. 2021), constructed PM2.5 concentrations by adding from specific species of pollutants, one of which was EC. We were therefore able to subtract the EC concentration from the PM2.5 concentration to obtain non-EC PM2.5 concentrations. We have added a clarification to the revised manuscript (P. 6, Lines 124-127):

*Because traffic is a major source of PM2.5 and EC one of the main PM2.5 components in urban environments,59 we removed the EC concentration from the total PM2.5 mass concentration (non-EC PM2.5) by subtraction, to avoid overadjustment when including both in the models simultaneously.* **7. Methods, adjustment for SES/occupation: was the goal to adjust for individual-level education and/or occupation (and its attendant exposures) or to adjust for overall SES in the household? E.g., "For each married participant, we used the higher of the couple's individual SES categories." This seems to be getting at household SES. Co-habitation among unmarried couples is common in Denmark. How was this addressed?**[**https://academic.oup.com/ije/article/42/2/559/737789**](https://academic.oup.com/ije/article/42/2/559/737789)**Was it important to capture information differently from previously married people? This should come down to the construct you are trying to measure. (The SES-ALS paper cited was about individual-level occupation.**

Our goal in adjusting for SES was to adjust for occupational exposure as well as household SES, as this would be potentially related to how quickly one is identified as having ALS in the registry system and to the predicted air pollution concentrations at their residence. Related to this, when compared with previous marriage information, current household SES may have had more influence on the speed at which ALS diagnosis could have occurred. In any case, we have a category of ‘divorced’ people in our civil status adjustment (P. 7, Lines 151-153):

*For each married participant, we used the higher of the couple’s individual SES categories, when available. We also used information on civil status (never married, married, divorced, widowed) […]*

The Reviewer is correct to point out that co-habitation is common among couples in Denmark, which would not be captured by our analysis. We have no detail in our dataset which would allow this to be adjusted for. We have added this as a Limitation in the Discussion of the revised manuscript (PP. 16-17, Lines 358-361):

*While our analysis adjusted for marital status and household SES, many couples in Denmark cohabitate. This would not be captured by our analysis, and ALS diagnosis in relation to cohabitation status should be further investigated.87*

**8. Methods, timing of covariates: for covariates whose values could vary over time (e.g., occupation, civil status, parish-level SES), what was the timing of these covariates relative to the exposure and outcome periods?**

All covariates for cases and controls were constructed similarly, using exact dates and matching to index date where the data allowed. Specifically, civil status was assessed at index date; last reported place of residence was the last residence the case or control had reported at the index date; parish-level SES is based on last reported address at index date; employment was based on last self-reported job title; SES was obtained based on the last self-reported job title, which for older ages, during which most the ALS cases in our study occurred, was not usually obtained at the index date, which we have clarified in the revised manuscript (P. 7, Lines 141-144):

*We included a set of covariates based on as close as possible to index date to account for potential confounding bias, including household socioeconomic status (SES) based on last-reported job title at index date; civil status at index date, last reported place of residence at index date, and place of birth.*

**9. Methods: calendar time as a course of confounding? Given that air pollutant concentrations and other determinants of ALS may have changed over time, might calendar time be a source of confounding?**

In a cohort study, this could certainty be the case. However, ours was a case-control study. Given that our cases and controls were matched on age and year of birth (among other characteristics) the controls would have been identified at a point very close in time to the associated controls. Therefore, we do not hold that calendar time could have been a source of confounding. We have clarified this in the revised manuscript (P. 16, Lines 342-343):

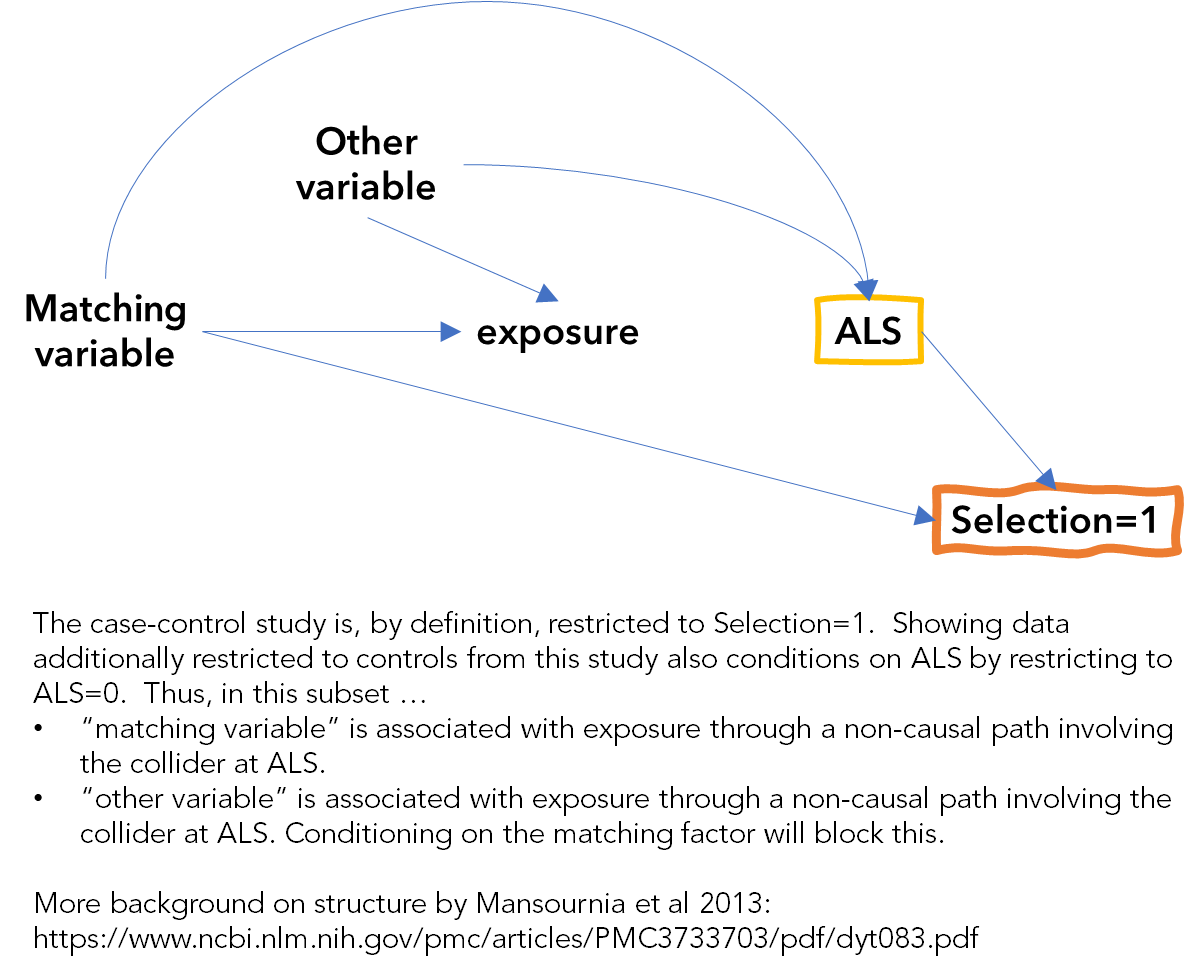
*We do not expect that calendar time was a potential source of confounding, as the controls were matched on age and year of birth.*

**10. Discussion: potential that smoking and/or BMI could be sources of confounding. The paper states, "… to induce confounding bias, any unaccounted-for variable would have to influence both ALS diagnosis and air pollution. BMI, previously associated with ALS, would not confound the association between traffic-related air pollution and ALS, as pollutant concentrations are derived independently from BMI distribution. Any BMI-air pollution association in our study, thus, would be via SES." Through complex social and economic mechanisms, the association of air pollution exposure with any given factor (e.g., BMI) can vary across populations. For example, in some study populations, areas where smoking is more common have higher concentrations of traffic-related air pollutants; in other settings, the pattern is reversed; and still others, there is little association. How is it known that BMI is not associated with exposure in this study population? Furthermore, how is it known that any such association in this study population would operate through SES? Analogous questions could be posed about smoking, as well. In the absence of firm answers to these questions, it could be useful to conduct a quantitative bias analysis, particularly as the estimated effect sizes are small.**

We agree with the Reviewer that there are complex social and economic mechanisms through which BMI may well be related to pollution levels in Denmark. However, to be a potential source of confounding, BMI would have to be causing the variation in pollution levels. There is no evidence that we know of that would suggest that BMI drives variation in pollution levels, and the way that the exposures were predicted did not in any way take into account individual-level BMI data. Rather, SES predicts where one lives, and also one’s BMI, and where one lives is a driver of air pollution levels, which warranted adjusting for in our analysis. We have blocked the path via SES if there were such an association.

Because Reviewer 1 asked us to remove most of this section, we have largely deleted and summarized the original statements.

**11. Table 1/Table 2: Distribution of covariates by exposure level. To provide more information on correlates of exposure in the underlying population, conditional on the matching factors, it would be helpful to provide a table showing the co-distribution of key covariates and air pollutant exposure among the controls. This could be a challenging proposition, though. The matching scheme means that the associations of matching factors with exposure may be distorted so long as there are other common causes of exposure and ALS. Thus, it is not clear whether, without extensive additional exploration, it would be possible to show meaningful co-distributions of exposure with the matching variables. However, conditional on the matching variables, it may be informative to show associations of other covariates with exposure. See example DAG below.**



We thank the Reviewer for this comment. We now provide summaries of 5-year average pollutant concentrations of controls by socioeconomic status, civil status, last reported place of residence, and place of birth in eTables 1-4, with eTable 1 below as an example:

**eTable 1.** Summary of 5-year average pollutant concentrations of controls by socioeconomic status (all in μg/m3).

| Pollutant | Overall, N = 19,2981 | Group 1 (Highest), N = 1,8861 | Group 2, N = 2,3401 | Group 3, N = 3,5751 | Group 4, N = 5,5221 | Group 5 (Lowest), N = 3,7021 | Group 9 (Unknown), N = 2,2731 |
| --- | --- | --- | --- | --- | --- | --- | --- |
| **NOX** | 27 (20) | 29 (20) | 27 (20) | 25 (17) | 27 (20) | 27 (19) | 30 (23) |
| **CO** | 237 (105) | 244 (103) | 233 (104) | 225 (89) | 237 (104) | 237 (102) | 258 (130) |
| **EC** | 0.85 (0.42) | 0.89 (0.42) | 0.84 (0.42) | 0.79 (0.37) | 0.85 (0.42) | 0.84 (0.41) | 0.92 (0.48) |
| **non-EC PM2.5** | 11.76 (2.37) | 11.75 (2.21) | 11.54 (2.24) | 11.58 (2.30) | 11.69 (2.34) | 11.93 (2.43) | 12.13 (2.62) |
| **O3** | 52.0 (6.0) | 51.1 (5.9) | 52.0 (5.9) | 53.0 (5.6) | 51.9 (5.9) | 52.2 (5.9) | 50.7 (6.4) |
| 1Mean (SD) | | | | | | | |

**12. Methods/Bayesian hierarchical approach: I am not particularly fluent in Bayesian methods, so will leave to others to evaluate the particulars of this approach. Nonetheless, if possible but without offering an entire course in Bayesian methods, it would be helpful for readers like me to see a clearer justification for choosing this method over, say, conventional conditional logistic regression (some of this clarification might tie into clarifying the target estimands mentioned in #1), and motivation or intuition for some of the specific steps and interpretation of posterior probability. For example, in writing, "We placed a hierarchy on the traffic-specific pollutant terms in the model," does this mean that you have assumed that one pollutant emanates from another? Also, does it make sense to compute credible intervals for the posterior probabilities?**

The main advantage of the Bayesian hierarchical structure we have utilized in our analysis is that the variance-covariance structure of the traffic-related pollutants (EC, NOx, CO) can be incorporated into the model, enabling an estimate of each individual pollutant’s association with ALS diagnosis, as well as joint (i.e., percentage change in odds of ALS diagnosis with increase in each of EC, NOx, CO) and overall (i.e., average percentage change in odds of ALS diagnosis from each of EC, NOx, CO) associations. The Reviewer is correct that this enables the model to account for the fact that the traffic-related pollutants originate from common sources. We have added this description in the revised manuscript (P. 10, Lines 209-211):

*We placed a hierarchy on the traffic-specific pollutant terms in the model to account for the fact that the traffic-related pollutants, EC, NOx, CO, originate from common sources and primarily traffic in urban environments:*

The posterior probabilities are generated from the full posterior probability distributions of the marginal estimates for each pollutant-specific, overall, and joint association. To calculate the posterior probability that an effect estimate was greater than null, we took a large amount of draws from these full distributions (4,000 in our case) and estimated the proportion of samples which were above zero (null association). To clarify how to interpret this value, for which there is no credible interval, we have added a description to the revised manuscript (P. 11, Lines 237-241):

*To calculate the probability that an association estimate was greater than null, we used the 4,000 samples of the posterior distribution and took the proportion of samples which were above the null. A 50% probability means that it is as likely as not that the marginal estimate is null, a probability closer to 100% indicates that the association is more likely to be truly positive, with closer to 0% indicating more likely to be truly negative.*

**MINOR  
  
13. Abstract: "For a standard deviation (SD) increase in 5-year average…." For more context, please provide the SD for each pollutant.**

We now provide the SD values for each pollutant in the Abstract of the revised manuscript (P. 1, Lines 15-19):

*For a standard deviation (SD) increase in 5-year average concentrations, EC (SD=0.42µg/m3) was potentially individually associated with an increase in odds (11.5%; 95% credible interval[CrI]:-1.0%,25.6%), with decreases individually for NOx (SD=20µg/m3) (-4.6%;95%CrI-18.1%,8.9%) and CO (SD=106µg/m3) (-3.2%;95%CrI-14.4%,10.0%) and a null association of non-EC PM2.5 (SD=2.37µg/m3) (0.7%;95%CrI-9.2%,12.4%).* **14. How PM2.5 was used in this investigation was presented with some ambiguity. The introduction states, "Using three air pollutants commonly used in health studies as traffic-related emissions tracers—nitrogen oxides (NOx), carbon monoxide (CO), and elemental carbon (EC)— as well as fine particles (PM2.5) and ozone (O3), we aimed to assess whether exposure to (a) each individual air pollutant is independently associated with ALS diagnosis," suggesting that PM2.5 was being considered as a traffic-related air pollutant (similar to the aforementioned situation for ozone). PM2.5 is not necessarily traffic-related, as the authors later state, but here it appears to be one of the primary exposures of interest.**

To avoid ambiguity, which the Reviewer has correctly pointed out, we have removed reference to PM2.5 and ozone at this point in the Introduction in the revised manuscript (P. 4, Lines 68-72):

*Using three air pollutants commonly used in health studies as traffic-related emissions tracers—nitrogen oxides (NOx), carbon monoxide (CO), and elemental carbon (EC)— we aimed to assess whether exposure to (a) each individual air pollutant is independently associated with ALS diagnosis, and estimate their (b) joint and (c) overall traffic-related emissions associations.*

**15. A curiosity: the relative difference in odds (percentage difference in odds) is effectively an arithmetic variation on the odds ratio. Was there a particular reason the authors opted for the percentage difference expression?**

The Reviewer is correct, and we opted for percentage difference purely for wider interpretability, i.e., so a non-specialist epidemiologist might be able to read through the results and understand the numerical output.

**16. Abstract: Given the results, it was a surprise to see this conclusion in the abstract "Our results indicate a potential positive association between ALS diagnosis and pollutants, particularly for EC." Perhaps this ties into clarifying the contribution of the Bayesian approach?**

We attempted to make our Conclusions in the original Abstract indicative but not conclusive, which is the reason behind using the phrase ‘potential association’. We have further clarified with the clear statement that our results are inconclusive in the revised manuscript (P. 2, Lines 24-26):

***Conclusions:*** *A potential positive association between ALS diagnosis and pollutants, particularly for EC, though results are inconclusive. Further work is needed to understand the role of traffic-related air pollution on ALS pathogenesis.*

**17. Methods/Index date: Please state earlier that the date of diagnosis as indicated in the database is the index date. I.e., "We identified ALS cases based on their International Classification of Diseases (ICD) discharge diagnoses, …, using the date of the first relevant code as the diagnosis date. This was the index date."**

We have done this (P. 4, Lines 82-84):

*We identified ALS cases based on their International Classification of Diseases (ICD) discharge diagnoses, i.e., ICD-8 code 348.0 (ALS) until 1993 and ICD-10 code G12.2 (motor neuron disease) thereafter, using the date of the first relevant code as the diagnosis date.*

**18. Methods/matching scheme: what was the degree of match sought for age and year of birth (within months, years?).**

The matching was made by age, sex, single year of birth, and vital status. We have clarified that matching by finer scale than year of birth was not possible in the revised manuscript (P. 8, Lines 169-170):

*Matching by finer scale than year of birth was not possible.*

**19. Methods/study design: The control-sampling scheme seems to follow a risk-set matching pattern, so cases could serve as controls. If that is correct, could state that. It also means that computed ORs are estimates of IRs.**

The Reviewer is correct. We have stated this in the revised manuscript (P. 5, Lines 97-99):

*Controls were alive and free of diagnosed ALS at the ALS diagnosis date of the matched case (index date). The control-sampling scheme followed a risk-set matching pattern, so cases could have served as controls before diagnosis of ALS.50*

We have also added clarification that (pg. 112) to be non-cases at the time of occurrences, odds ratios are IRs in the revised manuscript (P. 11, Lines 230-233):

*We present all results as percentage change in odds of ALS diagnosis per SD increase in pollutant concentration (calculated via e.g., , etc. obtained in the modelling process). Due to the risk-set matching pattern of our case-control study, odds ratios are also equivalently incidence ratios (IRs).65*

**20. Methods/occupational classes: these are likely official terms of the DK government, but they are not very descriptive and "unskilled" is somewhat derogatory. Although extensive detail is not needed, a little more would be informative.**

We accept the Reviewer’s point that the ‘unskilled’ by itself as a term is derogatory, so have replaced with ‘unspecialized’, and have provided some context in the revised manuscript (P. 7, Lines 147-150):

*Group 1 (highest status) includes corporate managers and academics; group 2: proprietors, managers of small businesses and teachers; group 3: technicians and nurses; group 4: skilled workers; and group 5: unspecialized workers, such as entry-level positions within food and retail environments.*

**21. Discussion: "If other sources of air pollution are associated with ALS, then including non-EC PM2.5 adjusts for other air pollutants from other sources." Is it known that air pollutants that fall outside of PM2.5 (most obviously, anything in the coarse fraction of PM10) are not related to ALS risk?**

We have corrected this sentence to indicate that we are talking about other sources of PM2.5 in the revised manuscript (P. 9, Lines 192-194):

*If other sources of air pollution are associated with ALS, then including non-EC PM2.5 adjusts for PM2.5 from other sources,67 as well as indicating whether pollution from other sources not explicitly quantified might also have associations with ALS.*

**22. This phrasing was unexpected: "The conditional approach automatically accounts for matching factors (age, sex, year of birth, vital status) …." What is meant by "automatically accounts for"?**

We have clarified the language in the revised manuscript (P. 8, Lines 167-169):

*The conditional approach examines contrasts within matched strata, i.e., groupings of case and matched controls, implicitly adjusting for matching factors (age, sex, year of birth, vital status) within each matched stratum.65*

**23. Discussion: Please take care to avoid relying on null hypothesis significance testing to interpret the findings. See the journal's guidance here:**[**https://edmgr.ovid.com/epid/accounts/ifauth.htm/**](https://edmgr.ovid.com/epid/accounts/ifauth.htm/)**In addition, the American Statistical Association issued a strong critique of significance testing (**[**https://www.amstat.org/asa/files/pdfs/P-ValueStatement.pdf**](https://www.amstat.org/asa/files/pdfs/P-ValueStatement.pdf)**), and additional cogent arguments along these lines have been issued elsewhere, including Nature. (**[**https://www.nature.com/articles/d41586-019-00857-9**](https://www.nature.com/articles/d41586-019-00857-9)**)**

We agree wholeheartedly with the Reviewer that testing significance as the sole mechanism to deciding whether results are non-null should be avoided, as it focuses purely on the extreme part of the distribution and is prone to dismissing potentially important conclusions from results.

One of the main benefits of our Bayesian approach is that each marginal estimate’s distribution probability mass is described fully, and therefore one is able to examine how much of the probability mass is above a null association, which in our view, and many statisticians’ views, is a more ‘natural’ way of describing the confidence that a result is non-null.

We have avoided any reference to p-values, as they do not have a place in Bayesian analysis, and have now avoided any use of words related to the significance of a result in the revised manuscript, e.g., (P. 14, Lines 296-300):

*In the largest case-control study of ALS and traffic-related air pollution to date, we found that a joint increase in average concentrations of traffic-related pollutants was potentially associated with an increase in odds of ALS diagnosis, with the clearest results for EC. We found that EC had the largest-in-magnitude independent association with ALS diagnosis, while associations with NOx and CO were negative with credible intervals overlapping the null, and smaller in magnitude.*

**\* \* \* \* \***

**Preparing a revision**

**1. For estimates of causal effects, we strongly discourage the use of categorized P-values and language referring to statistical significance, including whether a confidence interval covers the null. We prefer instead interval estimation, which conveys the precision of the estimate with respect to sampling variability. We are more open to testing with respect to modeling decisions, such as for tests of interaction and for tests for trend.**

We have avoided p-values throughout.

**2. We do not permit acronyms unless they are generally recognized by epidemiologists (e.g. HIV is okay, but LVA is not). When in doubt, we recommend that you spell out.**

We have been careful to introduce acronyms where used.

**3. Please do not include uninformative precision (excessive decimal places). For example, percents should be rounded to nn%, n.n%, or 0.0n% and risk ratios should be rounded to nn, n.n, or 0.nn unless clarity of the presentation and the sample size justify**

**more significant digits.**

We have done this.

**4. Please be sure to include explicit information about approval of human subjects research by an independent review board. If no such review was required, include an explicit statement about why the requirement for review was waived.**

We have done this in the manuscript (P. 5, Lines 107-108):

*This study was approved by the Institutional Review Board Committee at the Columbia University and the Danish Data Protection Agency.*

**5. Do not include public health policy recommendations in Brief Reports or Original Articles that present new research findings.**

We have not included any public health policy recommendations.

**6. Data appearing in the abstract must also be cited in the main text, not just in tables or figures.**

We have done this.

**7. Resubmissions must adhere to word limits. The word limits for main text (generally the introduction, methods, results, and discussion) are 1500 words for Brief Reports (plus 150 words for its abstract), 4000 words for Original Articles (plus 250 words for its abstract), 5000 words for reviews (plus 250 words for its abstract), 2000 words for Commentaries (no abstract), 600 words for Research Letters (no abstract), and 400 words for Letters to the Editor (no abstract).**

We have done this, with an Abstract of 250 words and an Original Article of 3,896 words in the revised manuscript.

**8. We advise that total word counts for Original Articles should not exceed 7500 words and for Brief Reports should not exceed 3500 words. The total word count includes main text (introduction, methods, results, and discussion), bibliography, figure legends, tables, and figures (250 words per figure, including each figure in a panel). The title page, abstract, acknowledgments, and funding information do not count in the total word count.**

We have adhered to this, with a total word count of 6,383 words in the revised manuscript.

**9. Figure labels: Make font size as large as possible, so as to be legible when figures are reduced for publication (typically one column [8.5cm] in width).**

We have made the Figure labels large and legible.

**10. Footnotes to tables and figures should use superscript lowercase letters to link content to the footnote, not symbols or numerals.**

The footnote in Table 1 uses a superscript lowercase letter.

**11. Do not use parenthetical phrases like “(data not shown), (results not shown), or (available from the authors upon request).” In these circumstances, the data or results should be provided in Supplementary Digital Content.**

We have avoided any use of these phrases.

**12. Additional details regarding submission requirements can be found in the Instructions for Authors, which are posted at**[**http://edmgr.ovid.com/epid/accounts/ifauth.htm**](http://edmgr.ovid.com/epid/accounts/ifauth.htm)**.  
  
Preparing for resubmission**

**13. Prepare a response document for the Editor that responds point-by-point to the reviewers' comments (presenting each comment followed by your response). Give the page number where revised text can be found and, where practical, paste revised text directly into the reply document.**

We have done this.

**14. Submit versions of the manuscript with and without your changes displayed.**

We have submitted clean and tracked versions of the revised manuscript.

**15. Supplementary Digital Content should be submitted as a single PDF file, and you should use our convention - e.g. eFigure 1, eAppendix 2 - to label and refer to online content.**

We have done this.

**16. Authors should submit copies of any closely related manuscripts (published, in press, or under review).**

We do not have similar manuscripts to submit at this time.

**17. Please revisit information about page charges and color printing charges available in the Instructions for Authors, which are posted at**[**http://edmgr.ovid.com/epid/accounts/ifauth.htm**](http://edmgr.ovid.com/epid/accounts/ifauth.htm)**.**

We acknowledge the charges on the link provided.

**18. We request that the complete revised manuscript (with all tables and figures) be completed by 05 May 2022. If you are not able to meet this deadline, please notify the editorial office.**

We have submitted before 5th May 2022.

**Resubmitting via Editorial Manager**

**19. Log-in to Editorial Manager as an author using the credentials above.**

**20. Click on the "Submissions Needing Revision" link.**

**21. To view the previous decision letter and reviewer comments, please click the blue decision term listed under the View Decision menu.**

**22. If you would like to download the previous manuscript to make revisions, click on "Download Files" under the Action menu.**

**23. To begin the resubmission: Click "Submit Revision" under the Action menu.**

**24. Proof each screen to ensure the information is still correct (the Title, Authors, etc.), then click Next at the bottom of each page.**

**25. On the Attach Files screen, select each previous submission item that you would like to carry forward to the resubmission.**

**26. Upload the revised versions of the main text (with and without tracked changes), and order them with the highlighted version first.**

**27. Upload the point-by-point reply to review.**

**28. When you are finished uploading, please click Next.**

**29. Click "Build PDF for My Approval."**

**30. Click "Go to Submissions Waiting for Author’s Approval."**

**31. Wait for the PDF to build. When it has been built, you will see the link "View Submission" in the Action menu. Click "View Submission," and open the manuscript to proof your work.**

**32. If you find problems with the manuscript, click "Edit Submission" from the Action menu. Make the required changes, and begin again at the file uploads.  
33. Once the submission is complete and acceptable, click "Approve Submission" from the Action menu.**

**34. If you have difficulty with these procedures, you may send questions to**[**timothy.lash@epidemiology-journal.com**](mailto:timothy.lash@epidemiology-journal.com)**.**

Thank you for the resubmission instructions. We have followed them.

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