

Reviewer#2

Overview -

The authors report measurements relevant to the problem of CCN activation. Their data set contains activated ratios (AR(D)) at six supersaturations ($S \sim 0.07, 0.1, 0.2, 0.4$ and 0.8%), an assessment of organic aerosol mass fraction (integrated over all particle sizes), ambient aerosol particle size distribution (PSD), and ambient CCN concentration. Comparing measurements made at two locations (Xinzhou in 2014 and Xianghe in 2013), the authors find a difference in the shape of the AR(D) and a difference in the 50% activation dry diameter. Some of the Xinzhou AR's plateau at a value that is significantly less than unity.

What I have summarized (these results are in Figures 1 and 2) is compelling, easy to understand and worthy of publication. However, the presentation that follows (Figures 4, 5, 6, 7 and 8) is difficult to understand. Most perplexing is the use of terminology "estimated Nccn", "calculated Nccn", "predicted Nccn" and "modeled Nccn" to describe quantities derived from the measurements. As I discuss below, these things need to be described.

Re: the terminologies like "estimated, calculated or predicted ..." in the paper has been unified.

There another issue:

I am surprised the two methods (estimated and observed in Figure 5) do not agree better. I note that there is a bias and that the correlation coefficients (R^2) is rather small. Is it possible that coincidence in either the SMPS/CN or in the CCN contribute to this poor agreement? Related to this, Rose et al. (2008) say: "For the calibration experiments, the number concentration of monodisperse aerosol particles was kept below $3 \times 10^3 \text{ cm}^{-3}$ to avoid counting errors caused by coincidence." The other possibility is that using a campaign-averaged AR(D) may have contributed the discrepancy. Perhaps there are other reasons. The authors should address why the two methods compared in Figure 5 correlate so poorly.

Re: the Figure 5 was with the aim to investigate the influence of the variation in size distribution on N_{CCN} , the actual measured CCN efficiency spectrum is multiplied by a campaign averaged particle number size distribution (PSD) (assuming not changed along with the time), which yields the CCN size distribution. This is then integrated over the whole size range (14-600 nm) to estimate N_{CCN} . But actually, the PSD changed significantly and showed temporal variations, which is the main reason for the poor correlations exhibited in Figure 5. That shouldn't be caused by the instruments. However, after careful reconsideration and revision of the paper, this section just was removed (as proposed by the first reviewer), and the new version will be focused on detection the

sensitivity of fraction of organics as well as the oxidization level of organics on estimating N_{CCN} .

Summary -

The manuscript needs to be reworked.

Specific Comments -

Abstract

Isn't there is a contradiction between the statements 1) variation with PSD showed a poor correlation and 2) the PSD played a dominant role?

Re: revised. Please see the above response.

P16146L13-16

Here you say there is little pollution from cars or industry, but in the next sentence there is mention of plumes from Xinzhou. This needs clarification.

Re: thanks for the comments. It has been clarified in the revised version.

(Page 6-7, Lines 81-89)

P16147 L10 These "relative deviations" are because of particle loss in the nafion dryer? Also, it is not clear what the "kinetic limitations" are.

Re: the relative deviations are due to the influence of dehydration-related particle mobility changes inside DMA. According to Mikhailov et al., 2009, void fractions as well as residual water in dried aerosol particles that are not water-free (due to kinetic limitations of drying or stable hydrate formation) should be taken into account in Kohler model

calculations of hygroscopic growth and CCN activation.

First, if the dehydration processes (efflorescence, restructuring, or desorption) inside DMA are completed within 0.1 s, then the resulting changes in particle mobility diameter should be fully captured with deviations <1%. And also, kinetically limited dehydration processes that lead to progressive changes of particle mobility on a time scale of 0.1–10 s should significantly influence the particle sizing (deviations >1%) and lead to a broadening of the measured size distributions. Dehydration processes progressing on time scales >10 s should have no effect on particle sizing (no change of mobility diameter and no broadening of size distribution).

P16147L25

It is the inner diameter, not the outer diameter, that is relevant.

Re: it is just the outer diameter.

P16147L29

2.5 mm ?

Re: 2.5 um, revised.

P16148L19

“..temperature stability was zero.” I don’t understand what you are referring to here.

Re: here it means the data is invalid if the “temperature stability” was flagged as “0”. And for the valid data, the “temperature stability” was flagged as “1”. The sentence has been revised.

P16148L21

Here you define the “aerosol number (CN) size distribution spectrum.”

How is this different from the PSD mentioned on P16147L1 and on P16148L24?

Re: that’s exactly the same thing. We have corrected all of them to particle number size distribution in the revised paper.

P16148L27

Since Figure 1 shows efficiency curves for both sites, I don’t understand why the “campaign-averaged” efficiency was used for this. Do you mean the campaign-average for the Xianghe site? To me it is unclear what the campaign is (both Xinzhou in 2014 and Xianghe in 2013; or just Xinzhou in 2014)?

Re: in Figure 1, the campaign averaged CCN efficiency spectra observed at Xianghe in summer 2013 are showed with black lines, and the campaign averaged CCN efficiency spectra observed at Xinzhou site in summer 2014 are showed with red lines.

P16152L6

It is the “CCN activity”, not the “aerosol activity”, that is the focus here.

Re: corrected.

P16153L2

Transportation -> transport

Re: corrected.

P16153L26

Here is a relevant reference.

Snider, J.R., and, S.Guibert, J.-L. Brenguier and J.-P.Putaud, Aerosol activation in marine stratocumulus clouds: Part – II Köhler and parcel theory closure studies, J. Geophys. Res., 108, doi:10.1029/2002JD002692, 2003

Re: the reference has been added.

P16154L9-9

I don't agree with your statement that the Ji and Shaw (1998) and/or the Twomey (1959) ($N_{ccn} = CSk$) parameterizations assume uniform aerosol composition. I also do not agree with your assertion that these parameterizations do not take into account variation in CCN loading. For example, you could have taken an observed CCN spectrum (N_{ccn} versus S) and fitted that as a function of S (e.g., $N_{ccn} = CSk$). That spectrum accounts for the PSD and composition. The issue you mention is relevant only if you do not have a measured CCN spectrum and you proceed, in a model, with a generic "C" and a generic "k".

Re: yes, the reviewer is right. The paragraph has been rewritten. Some new discussion has been revised and added (see Section 4.3, Page14-15,

Lines 268-284).

Figure 1

Symbols and line color (black and gray) are used to designate results obtained at the two locations (Xinzhou and Xianghe). The plot would be much more easily understood if you present with a better scheme for delineating the two stratifications (location and supersaturation).

Re: revised. Thanks for the comments.

Figure 2 – Case #1

Isn't the behavior seen here ($AR < 1$ at large diameter) indicative of an externally-mixed aerosol, for example one containing pure soot and hygroscopic particles? Isn't that worth mentioning/discussing? Isn't external mixing a complicating factor when it comes to converting the measurements to a CCN spectrum?

Re: Yes, the reviewer proposed the point exactly. We have mentioned this thing in the revision (see Page 13, lines 244-247), but because we didn't measure the mixing state of the particles, we say the low maximum activation fraction (<1) is just an indication of externally-mixed particle, as the reviewer suggested. Of course, the mixing state of the particles would greatly impact the CCN activation. We are now working on a paper, which will concentrate on discussing this issue by using the size-resolved chemical composition data measured by AMS.

P15154L7

What are “bulk ARs”?

Re: bulk ARs means the ratio of N_{CCN} to N_{CN} , which is calculated from the total CN and CCN number concentrations. Thus, we called it bulk ARs.

For the size-resolved CCN measurements, we can get size-resolved ARs from size-resolved CCN and CN number concentrations.

P16154L8

Reference to the specific section of Pruppacher and Klett is needed.

Pruppacher and Klett is a huge book.

Re: revised.

P16154L23

In Figure 4 there is no obvious indication of negative correlation in the plotted data, or in the fit lines (all of these have positive slope). If you provided the “R”, not R^2 (text) and if you defined the “R” (Pearson product moment), that would solve this problem. Also, the Figure legend shows $R^2 < 0$, which is mathematically impossible. All of this needs work.

Re: the Figure 4 has been replotted. For the cases with $R^2 < 0$, that is because that we used the lognormal axis here.

P16154L25-26

I find this statement of a “CN size distribution” confusing. Isn’t this the PSD define earlier. If that is the case you should use a consistent definition throughout. If a distinction is needed, you

should define/distinguish these early in the manuscript.

Re: "CN size distribution" here is just the PSD. Thanks a lot for the comments. This issue has been revised.

P16155L1

This is confusing. If you multiply the CCN efficiency spectrum by the (CN) size distribution, you get a "CCN distribution", but that is only valid for the selected S. I feel more clarity is needed here.

Re: revised. (see Pages 16, lines 306-313 and see Pages 18, lines 358-364)

P16155L2 What you are calling the "estimate" is the summed product of AR(D) and PSD(D)? Right? By "Observation" you are talking about the direct measurement of the ambient $N_{ccn}(S)$ made _without_ the DMA in front of the CCN instrument. I did not see mention of the ambient $N_{ccn}(S)$ measurement (without the DMA in front) in Section 2.1.

Re: We used a Scanning Mobility Particle Sizer (SMPS), combined with a Droplet Measurement Technologies-Cloud Condensation Nuclei Counter (DMT-CCNc) (Lance et al., 2006), for size-resolved CCN measurements as well as particle number size distribution (PSD) measurements. The SMPS is just the DMA.

To estimate N_{CCN} , estimated CCN size distributions at the five supersaturations were calculated by multiplying the campaign-averaged CCN efficiency spectrum with the actually measured PSD. The estimated

N_{CCN} at the five supersaturations was then calculated by integrating the estimated CCN size distribution over the whole size range.

The measured CCN size distributions are integrated to produce the observed N_{CCN} .

P16155L5-6

This is a big jump. Why would the predicted CCN (you meant estimated, or is “predicted” different?) be influenced by PSD more than it is already? I note that the estimated is the summed product of $AR(D)$ and $PSD(D)$?

Re: in the early version, we discussed the influence of variations of PSD on CCN estimation. However, the part in the revised manuscript has been removed. (See above comments and response)

P16155L8

Here you are using “calculated” CCN? Is this different from “estimated” CCN and “predicted” CCN? If you mean “estimated CCN”, it is not clear how the organic mass fraction is being used in these calculations.

Re: all of them have been changed to “estimated” CCN. The section was revised and some more details regarding to how the organic fraction is being used in these calculations have been included. (see Pages 15-16, lines 292-305)

P16155L19

I would reword this because the Figure 7 shows how the difference

(estimated minus observed) varies with chi-org. The latter is the independent variable. There are other places in the manuscript where “sensitivity” is used. I would change the word order in some of these instances too. E.g., P16145L26. There are other places too.

Re: revised. Because in the revised version, the sensitivity of oxidation level (using f_{44} , the fraction of m/z 44 in total organics, as an indicator) of organics to estimation of N_{CCN} is also examined. The section is thus rewritten (see Section 4.4, and Figure 5 and Figure 6).

Figure 5

These are single-parameter (slope) fits, I think. Hence, the “linear” used here is a special case of linear. This comment applies to Figures 6 and 8 too.

Re: revised.

P16158L24

Mr. Mcribb's. Isn't this Maureen Cribb?

Re: yes. Thanks a lot for the careful check. The sentence in acknowledgement was removed.