Reviewer 2

We thank the Reviewer for their constructive comments on our manuscript. Please find our responses below highlight in yellow, and changes to the manuscript in **bold**.

"In this paper, the authors present a CRDS for HCl and evaluate its performance with an appropriate set of laboratory evaluations. They also report measurements of ambient HCl and compare these with another measurement technique; this comparison is done extremely well.

The findings are within the scope of AMT, substantive conclusions are reached about the instrument performance and field comparison, and the paper is well written. The instrument itself is not particularly novel since an extremely similar instrument was presented in AMT almost a decade ago (Hagen et al., 2014). This instrument does seem better than that one in some ways (mainly in the 30 s precision of 6 pptv, which is quite impressive), so I suppose publishing on this instrument is justified.

Still, there are some pieces that are missing or need revision. I discuss these in 'Broader Comments.' I'm calling these minor, but they do approach the threshold between 'minor' and 'major' revisions. I also have some decidedly minor suggestions, which follow as 'Specific Comments.' I recommend accepting for publication after these aspects are addressed."

Broader Comments:

1) The introduction should have a slighty more comprehensive discussion of HCl measurement techniques. There have been many cavity-enhanced HCl instruments developed in the past decade or so that are not mentioned here but should be. Los Gatos Research and Tiger Optics have already developed commercial, cavity-enhanced HCl spectrometers. Hagen et al., 2014 and Wilkerson et al., 2021 have also published on cavity-enhanced HCl instruments. Hagen's instrument is even a CRDS that measures HCl via its first overtone—the same as the instrument discussed here. So you should also explain how this instrument compares and contrasts with Hagen's since they are so similar.

Response: We thank the Reviewer for this suggestion to better represent spectroscopic techniques in the introduction and have added the following to the text.

"Spectroscopic techniques offer distinct advantages over some previous methods. Spectroscopic techniques for measuring atmospheric HCl reported by Hagen et al. (Hagen et al., 2014) and Wilkerson et al. (2021) have shown the precedent for fast time response, as well as sensitive, selective, and robust detection. The portability and fast time response for instruments is of great importance for spatial resolution and is therefore a key factor for field deployment." (P5, L90-L95)

2) There is no CRDS instrument schematic in this CRDS instrument paper. I understand that proprietary issues may limit how specific this can be, but please include at least a rough schematic of the instrument layout as either a main figure or supplemental figure.

Response: We have added a simple schematic of the CRDS to the supplement.

"The Picarro G2108 Hydrogen Chloride Gas Analyzer system was used for all analyses (see Figure S1, Dawe et al., 2019, www.picarro.com)." (P6, L119-120)



Figure S1. Simple schematic for the Picarro CRDS.

3) You state that tropospheric HCl mixing ratios vary between 10-1000 pptv. Yet, the data in Fig. 2 extends all the way up to 45 ppbv HCl and doesn't go below 10 ppbv HCl. You really need to be evaluating these effects closer to relevant, atmospheric levels of HCl. Ideally, you would dip below ppbv levels since those are the most relevant (as you further demonstrate in Fig. 3). Hagen et al., 2014 goes slightly below 1 ppbv HCl when they produced their correlation line (Fig. 8); they proposed an instrument extremely similar to this one and published that discussion in this very same journal (same with Wilkerson et al., 2021). There is, therefore, a clear precedent for a more rigorous evaluation than this. Having said that, I appreciate that laboratory assessments of sub-ppbv levels for HCl can get quite difficult. At bare minimum, you should remove the data above 30 ppbv and add at least two iterations of this experiment below 10 ppbv HCl.

Response: We appreciate the Reviewer's concern here. The levels we have generated are higher than typical atmospheric mixing ratios but validate the linearity and accuracy of the instrument in this range. As the Reviewer notes, it is challenging to generate an accurate HCl source at low mixing ratios. Thus, we have chosen to explore the lower HCl ranges using a field intercomparison. We have added text to clarify:

"We compared the CRDS analyzer-measured HCl with the gas standard mixing ratios provided by an IC-certified PD under dry conditions and observed a close to 1:1 correlation. We explored 5 mixing ratios 12, 16, 21, 32, and 45 ppbv (Figure 2). These levels are higher than have been observed in the ambient atmosphere but demonstrate good signal response linearity." (P15, L287-L290) "To practically validate the CRDS under real-world conditions **and atmospherically relevant mixing ratios**, an ambient intercomparison was performed over a period of 7 days (4–11 April 2019)." (P16, L302-L304)

Specific Comments:

Line 24: Recent reports of response times for HCl instruments are usually presented as 90% response times. Is that the case with this range? If not, please change this to a 90% response time range, so your instrument can be more easily compared with other recently reported ones (or at least explicitly call your current range a X% response time, where X is whatever this reported range corresponds to).

Response: We chose to use the more conservative τ_2 estimate on our response time to reflect challenges with inlet effects. However, we do present the 90% response times in Table 2. We have also added the minimum 90 % response time to the abstract:

"**The minimum 90% response time was 10 seconds and the equilibrated** response time for the tested inlet was 2–6 minutes under the most and least optimal conditions, respectively." (P2, L24-L25)

Line 29: You should probably drop 'response time' from this list. First, a response time of 2-6 minutes is quite worse than recent HCl instruments discussed in scientific literature (Hagen et al., 2014; Wilkerson et al., 2021) and a bit worse than commercial HCl instruments (e.g. LGR's HCl analyzer is ~1 minute: http://www.lgrinc.com/documents/HCl_Datasheet2019.pdf). Second, you don't display response times in Table 1 where evidence of a favorable response time compared to other instruments would presumably be presented to the reader.

Response: It is difficult to compare response times for instruments that measure these high surfaceactive gases, as it often reflects the response of the whole system (i.e. instrument and inlet). Under optimal low humidity conditions, we can achieve a 90% response time of 10 seconds (see Table 2). Without in-depth knowledge of the inlets, flow rates, RH, etc., response times of different instruments cannot be directly compared. We have added text to clarify this point:

"Major differences in the surface area between our instrument and the instruments to which we compare are likely to cause τ_2 differences. Our method employs the use of three filters that increase the gas to surface interactions, and therefore increase our equilibrated response time τ_2 ." (P21, L396-L399)

Line 49: Add just a couple sentences to explain where you would expect to see atmospheric HCl levels of 10 and 1000 pptv. For example, my understanding is that atmospheric HCl is elevated near oceans because sea salt spray provides a large source of chlorine to the atmosphere. (And if I'm wrong, then that's more evidence to support adding a couple sentences to briefly explain this to readers).

Response: The following text has been added:

"Elevated levels of HCl are typically found near marine environments polluted with NO_x; where reactions involving the chloride in sea spray aerosols can be a major source of chlorine to the troposphere (Crisp et al., 2014; Finlayson-Pitts et al., 1989; Haskins et al., 2018; Wang et al., 2019)." (P3, L50-L52)

Line 118: The first overtone has a very low intensity compared to the fundamental transition for HCl, and the fundamental transition is by far the most common transition used in HCl spectrometers (there are also ro-vibrational lines in the fundamental transition that are spectroscopically isolated from other atmospherically relevant species like H2O). You should clarify the language 'relatively high intensity' so the reader knows what this intensity is relative to.

Response: Text added for clarity:

"The first overtone (2-0 absorption band) of HCl is easily discernable from other absorbing species (e.g. H₂O, CH₄), has a relatively high intensity **compared to the fundamental absorption transition**, and is accessible to near-infrared (IR) diode laser light sources." (P6, L125-L129)

Line 234: The 30-second precision of 6 pptv is great—surprisingly great. Hagen et al. 2014 discussed a CRDS HCl instrument that also engaged a line in HCl's first overtone. This is pretty similar to your setup. Yet, their 60 s precision is 20 pptv. You should include a compelling explanation in this paper as to how your precision is so much better than any other HCl instrument, especially one that is so similar to yours.

Response: We agree that the performance of the commercial device is quite good. Unfortunately, we cannot compare the two instruments without detailed knowledge of the specific technologies. The commercial device is proprietary, and so we can only offer the references describing the methods as already presented in the Hagen et al. (2014) and Crosson (2008) references.

Line 234: Your response time is 2-6 minutes, but the precisions you report skip from 30 seconds to 1 hour. Can you provide a precision for an averaging time that is within range of or only slightly above your response time (e.g. 5 minutes or 10 minutes)?

Response: The Reviewer raises an excellent point. We have added precision for a 5-minute averaging time. *Also note that we corrected the reported value for the 2 second LOD and precision, which we noticed was mistakenly reported.

"The LODs determined in the CRDS measurements for 2 second, 30 second, **5 minute**, and 1 hour integration times were **66**, 18, **5**, and 2 pptv, respectively. Similarly, precision was determined from the Allan-Werle deviation in the blank over the same 10 hours of zero air sampling. Precision in a 2 second, 30 second, **5 minute**, and 1 hour integration time was **22**, 6, **2**, and 0.8 pptv, respectively." (P11, L240-L244)

Line 240: Change to, "**This** CRDS has many advantages compared to…" since you are comparing this instrument to another CRDS in Table 1.

Response: Changed as suggested.

Line 253: Again, this response time is not that fast compared to many recent HCl instruments (see earlier comment). Can you better contextualize this assertion?

Response: As described above, we cannot directly address this. A shared inlet intercomparison of HCl instruments would be beneficial to better understand the relative roles of inlets and detectors in determining instrument response times.

Line 261: Why does the Iodide Cl-HR-ToF-MS have a higher LOD (30 pptv) than precision (53.3 pptv)?

Response: We thank the Reviewer for pointing out this error. We have corrected the reported precision to match the LOD in the 30 sec integration.



Line 261: You allude to two recent cavity-enhanced HCl instruments in the main text that were published on in AMT (Hagen et al., 2014; Wilkerson et al., 2021), but only one is present in the table. There are also other commercial, cavity-enhanced instruments beside the one in this paper (LGR and Tiger Optics). If possible, can you add these missing ones to your table? (I say 'If possible' because their descriptions may be missing too much information to warrant inclusion).

Response: We have added data from Wilkerson et al. (2021) to the table. We have chosen to include only peer-reviewed data in Table 1. In our experience, specifications reported on data sheets are not held to the same rigour as peer-reviewed publications.

Off-axis integrated cavity output spectromet er (OA- ICOS)	78 pptv (30 sec)	<mark><11%</mark>	<mark>26 pptv</mark> (30 sec)	<mark>1 s</mark>	NR	NR	<mark>(Wilkers</mark> on et al., 2021)
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Line 261: Hagen et al., 2014 has an accuracy <10%, not >10% (>10% reads as if the accuracy is at best 10% but could be infinitely worse). I believe that the Tandem mist chamber and IC-CD is the same situation (>25% when it should be <25%). Please correct these or just report the number without a '<' sign.

Response: We thank the Reviewer for noting this mistake. It has been corrected.

Near-IR CRDS	60 pptv ^a (1 min)	<mark><10%</mark>	<mark>20 pptv</mark> (1 min)	<mark><15 s</mark>	NR	NR	<mark>(Hagen</mark> et al., 2014)
Tandem mist chamber and IC-CD	48 pptv ^c	<mark><25 %</mark>	<mark>24 pptv^c</mark>	<mark>2 h</mark>	NR	NR	(Keene et al., 2007, 2009)

Line 304: I think you mean to refer to Figure 3a.

Response: The numbering in references has been corrected throughout.

Line 425: Some of these references successfully used a silicon coating on their instrument, as you state on Line 422. Yet, your focus for the rest of this paragraph is on all the unpleasantness associated with fluorinated materials. What's your evaluation of silicon coatings, which lack all of these disadvantages? Please include that (even if it's just a few sentences).

Response: We thank the reviewer for their comment and have added the following text:

"Silicon coatings on all plumbed surfaces have been successfully used for atmospheric HCl measurements (Wilkerson et al., 2021), and recommended for applications where PFA use is impractical. Although a direct comparison has not been conducted for HCl, PFA inlet material has been reported to yield better response times than silicon coatings for nitric acid (Neuman et al., 1999). Differences in instrument configurations and applications may warrant the use of different inlet materials and coatings for successful measurement of atmospheric HCl." (P24, L449-L455)

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