UV spectroscopic determination of the chlorine monoxide (ClO) / chlorine peroxide (ClOOCl) thermal equilibrium constant

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Abstract. The thermal equilibrium constant between the chlorine monoxide radical (ClO) and its dimer, chlorine peroxide (ClOOCl), was determined as a function of temperature between 228 – 301 K in a discharge flow apparatus using broadband UV absorption spectroscopy. A third law fit of the equilibrium values determined from the experimental data provides the expression: $K_{eq} = 2.16 \times 10^{-27} e^{(8528 \pm 25 \text{ K/T})} \text{ cm}^3 \text{ molecule}^{-1}$ (1σ uncertainty). A second law analysis of the data deviates minimally: $K_{eq} = (2.14 \pm 1.14) \times 10^{-27} e^{(8530 \pm 123 \text{ K/T})} \text{ cm}^3 \text{ molecule}^{-1}$. From the slope of the van’t Hoff plot in the third law analysis, the enthalpy of formation for ClOOCl is calculated, $\Delta H^\circ_f (298 \text{ K}) = 130.0 \pm 0.6 \text{ kJ mol}^{-1}$. The equilibrium constant results from this study suggest that the uncertainties in $K_{eq}$ recommended in the most recent (year 2015) NASA JPL Data Evaluation can be significantly reduced.

1 Introduction

Halogen-mediated catalytic processing of ozone accounts for the overwhelming majority of lower stratospheric ozone-loss processes in polar winter and spring (e.g., WMO, 2014; Wilmouth et al., 2018). Approximately half of this loss (Wohltmann et al., 2017) is resultant from the ClO dimer cycle (Molina and Molina, 1987), which occurs as a result of the highly perturbed physicochemical conditions of the polar vortices:

$$\text{ClO} + \text{ClO} + \text{M} \rightleftharpoons \text{ClOOCl} + \text{M} \quad (R1)$$

$$\text{ClOOCl} + h\nu \rightarrow \text{ClOO} + \text{Cl} \quad (R2)$$

$$\text{ClOO} + \text{M} \rightarrow \text{Cl} + \text{O}_2 + \text{M} \quad (R3)$$

$$2(\text{Cl} + \text{O}_3 \rightarrow \text{ClO} + \text{O}_2) \quad (R4)$$
Within this cycle, the equilibrium governing the partitioning of ClO and ClOOCl in reaction (R1) is defined as:

$$K_{eq} = \frac{[\text{ClOOCl}]}{[\text{ClO}]^2}$$

This thermal equilibrium is a key parameter that determines the nighttime partitioning of active chlorine in the winter-spring polar vortex. The value of $K_{eq}$ can also tune the efficiency of chlorine-mediated ozone destruction, particularly the radial extent of ozone loss within the warmer Arctic polar vortex. For example, Canty et al. (2016) quantified how small variations in $K_{eq}$ can modulate significant changes in the temperature at which photolysis of ClOOCl and thermal decomposition of ClOOCl occur at equal rates.

Although the partitioning between ClO and ClOOCl is highly important, relatively few laboratory measurements of $K_{eq}$ have been made, and there is significant disagreement between reported values. Accordingly, the uncertainty in $K_{eq}$ was large (e.g., ~75% at 200 K) as of the 2011 JPL compendium recommendation (Sander et al., 2011). The most recent 2015 JPL-recommended value of $K_{eq}$ was revised on the basis of a 2015 study by Hume et al. (2015), but the recommended uncertainties are still substantial—exceeding 50% at 200 K (Burkholder et al., 2015).

The preponderance of laboratory data from previous determinations of $K_{eq}$ was obtained at temperatures significantly warmer than the polar stratosphere ($T > 250$ K). Error in the extrapolation of these warm temperature data has often been cited to explain the lack of correspondence between values of $K_{eq}$ determined in the laboratory and those calculated from stratospheric observations (Avallone and Toohey, 2001; Stimpfle et al., 2004; von Hobe et al., 2005; Santee et al., 2010). The more recent results of Hume et al. (2015) are unique in that they were obtained at temperatures colder than other laboratory studies ($206$ K < $T < 250$ K), but their experimental method was compromised by secondary bimolecular reactions at warmer temperatures. In the present study, our spectroscopic data bridge the warmer temperatures where most laboratory determinations of $K_{eq}$ have been made to the colder temperature work of Hume et al. (2015), covering a broader temperature range than any previous study. The thermal equilibrium constant between ClO and ClOOCl was measured as a function of temperature ($228$ K < $T < 301$ K) by UV spectroscopy and is evaluated here in relation to prior determinations, observations, and compendial recommendations.

2 Experimental

All experiments were conducted in a discharge flow apparatus, as shown in Figure 1. Independently programmable thermal zones allow for the optimization of target chemistry as a function of flow velocity, temperature, and pressure. ClO is synthesized via the reaction of Cl atoms with O$_3$ (reaction R4). Cl is produced from a 1% Cl$_2$/He gas mixture, diluted further with UHP He and directed through a 45 W, 2.45 GHz microwave discharge. O$_3$ is produced via electric discharge of a 10% O$_2$/Ar source mixture and subsequently introduced 2.5 cm after the microwave cavity. Once formed, ClO readily dimerizes to form ClOOCl (reaction R1), particularly at higher concentrations and colder temperatures.
To facilitate dimerization of ClO, the gas mixture is cooled in a 20-cm long jacketed quartz cell immediately subsequent to the microwave discharge. This reaction cell (Figure 1) has an inner diameter of 1 cm and can be maintained at a temperature between 198 – 305 K via circulating chilled methanol (NESLAB Endocal ULT-80). The operation of this cell at cold temperatures additionally suppresses undesired chemistry, preventing the synthesis of side products such as OCIO per reaction R6, and subsequently Cl$_2$O$_3$ per reaction R7.

\[
\text{ClO} + \text{ClO} \rightarrow \text{OCIO} + \text{Cl} \quad \text{(R6)}
\]

\[
\text{ClO} + \text{OCIO} + \text{M} \rightarrow \text{Cl}_2\text{O}_3 + \text{M} \quad \text{(R7)}
\]

Following the reaction cell, the gas stream then passes through the cold trap zone, which is maintained at temperatures between 100 K and room temperature depending on the experiment. Cooling is accomplished by flowing N$_2$ gas through a copper coil immersed in liquid N$_2$ and then through an 18-cm long insulated aluminum jacket surrounding the 1-cm inner diameter quartz flow tube. Type K thermocouples (alumel/chromel) are affixed at three positions on the outside of the flow tube, opposite the cryogenic gas ports. These thermocouples are further insulated to ensure the recorded voltages correspond to the temperature of the quartz tube and not the temperature of the cryogenic gas. The coupling between the cold trap and the equilibrium cell is actually linear but is presented as a right angle in Figure 1 for graphical purposes.

The next section of the flow system in Figure 1, labeled equilibrium cell, is a jacketed 50-cm quartz tube of 1-cm inner diameter. This section is where the gases reach equilibrium prior to measurement in the absorption cell. The equilibrium cell and the absorption cell share a coupled circulating chilled methanol bath (NESLAB ULT-80) ensuring that the two cells are maintained at the same temperature. The equilibrium cell is isolated from the environment with two 10-mm blankets of aerogel insulation (Cryogel Z). Additionally, a flow of cryogenic N$_2$ passes through an insulated aluminum jacket surrounding the union between the equilibrium cell and the absorption cell. This N$_2$ is chilled by passing through a copper coil immersed in the reservoir of the circulating chiller servicing the reaction cell, and the flow is modulated to provide constant temperature as the gas mixture transits from the equilibrium cell to the detection axis. A 100 Ω thermistor is inserted into the gas stream at this location to verify the temperature.

Finally, the gas mixture enters the absorption cell, a 91.44-cm jacketed quartz tube with an inner diameter of 2.54 cm. This detection axis is oriented at a right angle to the equilibrium cell and is terminated with two quartz windows. A 100 Ω thermistor is positioned at the halfway point. Cryogenic circulating methanol provides for temperature control between 228 – 301 K. Two 10-mm blankets of aerogel insulation (Cryogel Z) provide thermal isolation from the environment. An exterior dry N$_2$ purge is employed to prevent window condensation.

The discharge reactor is operated at pressures between 100 – 333 mbar. Pressure is monitored with Baratron capacitance manometers. Carrier gas flow rates, ~1.0 – 1.8 L min$^{-1}$ depending on the experimental conditions, are metered via MKS mass flow controllers. Cl$_2$ flow rates are controlled via a needle valve, while O$_3$ addition is modulated using micrometer flow
control valves. Total system pressure and velocity are tuned using an integral bonnet needle valve. Residence times within the absorption cell range between ~1 – 11 seconds, depending on gas flow rates, system temperature, and pressure. A quarter-turn plug valve provides a bypass of the integral bonnet needle valve such that rapid pump down of the reactor and reignition of the plasma can be performed without disturbing pressure calibration during the course of experiments.

Data were acquired using a fiber-coupled Ocean Optics USB4000 UV-Vis spectrometer (~0.3 nm resolution) illuminated by a Hamamatsu L2D2 deuterium lamp. The need to correlate Baratron, thermocouple, and thermistor sensor readings with each UV spectrum required the in-house development of custom software. Drivers and libraries to operate the spectrometer and simultaneously interrogate analog sensors were written in Python 2.7 and, in combination with the Python-Seabreeze library, provided scriptable, automated control of nearly all aspects of the data acquisition system.

The deuterium lamp was allowed to warm up for at least one hour prior to data collection activity to reduce small variations in lamp output on experimental timescales. Dark spectra were acquired prior to any experiments on a daily basis. Background spectra were obtained with the microwave plasma extinguished and all gas flows of species that absorb in the region of 200 – 295 nm (e.g. O₃, Cl₂) off. For consistency, sample spectra were obtained exactly 100 seconds after the background spectra against which they were referenced. Each saved spectrum consists of the co-addition of 597 individual scans, the number of scans that could be obtained in exactly 3 minutes of acquisition time.

To aid in the selection of experimental conditions, a simulation of the discharge-flow reactor was constructed. A numerical integrator for chemical kinetics (written in Python 2.7 with NumPy and SciPy) for 18 chemical species and 45 relevant chemical reactions was informed by JPL Data Evaluation 15-10 kinetic rate constants (Burkholder et al., 2015) and coupled into a physical model of gas flows as a function of reactor geometry, temperature, and pressure. Temperature and pressure ranges were scanned to determine optimal conditions to ensure ClO–ClOOCl equilibrium within the real-world experiment. Because parameterized simulations carry inherent uncertainty, experimental conditions were selected at several pressures along the equilibrium asymptote ($K_{eq}$ vs $P$), and real-world experiments were performed at pressures above and below the identified value in order to confirm asymptotic equilibrium behavior. The kinetic model was only used to inform conditions for the experimental setup, but no results from the model were used in the determination of the reported equilibrium constants.

3 Results and Discussion

More than 136,000 background and sample spectra were obtained between the temperatures of 228 – 301 K at pressures ranging between 100 – 333 mbar. Typical initial concentrations spanned $2 \times 10^{13} – 4 \times 10^{14}$ molecules cm⁻³ for O₃ and $1 \times 10^{14} – 4 \times 10^{15}$ molecules cm⁻³ for Cl₂. Active chlorine (ClOₓ) concentrations were typically $1 \times 10^{13} – 1 \times 10^{14}$ molecules cm⁻³ with the microwave discharge on. These values were tuned according to the initial conditions prescribed by the model simulations, as described above. For example, as the target temperature of the experiment decreased, the system was operated at incrementally higher pressures to allow more time for equilibrium to be achieved. Higher temperature samples reached equilibrium more readily, so gas velocity was increased to limit the impact of enhanced rates of secondary chemistry on observed $K_{eq}$ values.
Multicomponent spectral curve fitting software packages were programmed in Python 2.7/LmFit (Newville et al., 2016) for the deconvolution of the UV absorption spectra of O₃, Cl₂, ClO, ClOOCI, OCIO, and Cl₂O₃. Reference cross sections were utilized as follows: For O₃ and OCIO, pure sample spectra were acquired and scaled to match the 2015 JPL-recommended cross sections of Molina and Molina (1986) and Kromminga et al. (2003), respectively. ClOOCI and Cl₂O₃ cross sections were obtained directly from the 2015 JPL data evaluation (Burkholder et al., 2015). Temperature-dependent cross sections of Cl₂ were obtained from Marić et al. (1993) and validated to match observed Cl₂ spectra along the experimental temperature range. Synthetic temperature-dependent cross sections from Marić and Burrows (1999) were used for ClO due to the broad temperature range over which the data are available.

The cross sections from Marić and Burrows (1999) were found to provide an excellent fit of experimentally obtained ClO at all relevant temperatures in this study and were validated against available laboratory-determined ClO cross sections from the literature. Our experimental spectra at 263 K fit using both the synthetic ClO cross sections of Marić and Burrows (1999) and the reported experimental cross sections of Trolier et al. (1990) at 263 K result in concentrations of ClO that differ by only 1.6%. Similarly, experimental samples at 300 K from this study fit to the room temperature, laboratory ClO cross sections of Simon et al. (1990) and Sander and Friedl (1988) have excellent correspondence with the synthetic cross sections of Marić and Burrows (1999): 1.9% deviation in ClO concentration in comparison with Simon et al. (1990) and 2.7% deviation in comparison with Sander and Friedl (1988). The resolution of our experimental spectra was degraded to match the lower resolution data of Trolier et al. (1990) for this comparison, while the cross sections of Sander and Friedl (1988) and Simon et al. (1990), which were published at higher resolution than provided by our spectrometer, were degraded to our spectral resolution.

For all reported experiments, spectra were fit between 235 – 295 nm in order to capture the peak absorbances of both ClO and ClOOCI. Figure 2 provides example fits at (a) 230 K and (b) 300 K. In both cases, the residual (offset 0.14 A.U. for clarity) is observed to be flat and minimally structured, and this holds true for all spectral fits reported in this work. In those places where the residual is structured, the structure is primarily resultant from mismatch in instrument function between reference and experimental spectrometers. Once the ClO and ClOOCI concentrations are determined from the spectral fit, the value of $K_{eq}$ at the relevant temperature is calculated per equation (1).

A total of 87 independent experimental observations of $K_{eq}$ were acquired. The results are plotted in Figure 3 as a function of inverse temperature and are enumerated in Table 1. Data were considered independent when experimental conditions were modified between sample acquisitions (e.g., change in initial gas concentrations, pressure, temperature, carrier gas flow rates, or microwave discharge power). Data points that were not independent were reduced to a single independent data point by coaddition. Samples at colder temperatures were subjected to more repeated evaluations to improve accuracy under the low ClO conditions.

The temperature dependence of $K_{eq}$ can be related as an Arrhenius expression, per equation (2), with free parameters $A$ and $B$.

$$K_{eq} = A e^{(B/T)}$$  \hspace{1cm} (2)
In a third law fit, the prefactor $A$ is fixed to a prescribed value. For this work, we employ the JPL Data Evaluation recommended $A$ parameter value of $2.16 \times 10^{-27}$ cm$^3$ molecule$^{-1}$, which is the most recent literature evaluation of this constant (Burkholder et al., 2015). A third law fit of our $K_{eq}$ data yields a $B$ parameter value of 8528 K. This result was obtained by an ordinary least-squares fit of the independent measurements of $K_{eq}$ (Table 1). The fit is shown in Figure 3 as the black trace. Error from the fitting process was quantified via bootstrapping with 2000 resamplings of the binned $K_{eq}$ results, which establishes a fit error interval of $\pm$ 5.0 K. This method only accounts for the fit error and does not take into account other potential sources of experimental error, as discussed below.

The precision of repeated measurements conducted at the same experimental conditions demonstrated a temperature dependence due to ClO or ClOOCl concentrations approaching their experimental limit of quantification (ClO limiting measurements at colder temperatures and ClOOCl at warmer temperatures). To assess the significance of the lower precision at temperatures above 290 K and below 250 K, weighted least-squares third law fits were performed on sub-sampled data populations. An analysis of $K_{eq}$ results obtained between 250 – 301 K, 229 – 291 K, and 250 – 291 K resulted in $B$ parameters of 8527 K, 8529 K, and 8528 K, respectively, which deviate minimally from the parameter of 8528 K obtained from a fit of the entire temperature range. The extrapolated value of $K_{eq}$ at 200 K obtained from these sub-sampled datasets varies by $< 1\%$, while at 180 K the spread of the maximal deviation in $K_{eq}$ between subsets is $< 2\%$. An estimate of error from temperature-dependent precision is $\pm 2$ K about the $B$ parameter. The precision of repeated measurements conducted at the same temperature but varied flow rates and pressures did not statistically deviate from the precision from temperature-dependence alone.

Variation in the reference cross section of one component in a multicomponent fit may impact the quality of fit for the other spectral components. It is the trend in the literature to not explicitly include uncertainties from the reference cross sections during assignation of error for $K_{eq}$; however, the choice of synthetic cross sections for ClO is considered further here. As discussed, the synthetic temperature-dependent ClO cross sections prepared by Marić and Burrows (1999) were employed for the determination of [ClO] in this study, and the ClO concentrations from fits using these synthetic cross sections differed from ClO concentrations determined using experimentally derived reference standards at most by 2.7%. To capture the uncertainty of this error, all ClO concentrations used to derive $K_{eq}$ were scaled by $\pm 2.7\%$ and then fit by a weighted least-squares third law analysis, producing an estimated error in $B$ due to ClO cross section selection of $\pm 13$ K.

The selection of the wavelength window used for the spectral fit also produced variations in derived concentrations, and consequently $K_{eq}$, but the impact was not substantial. Ultimately, estimating our uncertainty on the $B$ parameter from all known sources of potential error yielded comparable, but smaller, values than simply assigning the error interval such that it fully encompassed 95% of the individual $K_{eq}$ results. Accordingly, we assign error intervals ($1\sigma$) to our $B$ parameter of $\pm 25$ K. Systematic errors arising from experimental design and post-processing technique are estimated to contribute errors that sum to a total smaller than this boundary. Our resulting $K_{eq}$ expression from the third law fit is shown in equation (3).

$$K_{eq} = 2.16 \times 10^{-27} e^{(8528 \pm 25 K/T)} \text{ cm}^3 \text{ molecule}^{-1}$$

(3)
Figure 4 provides a comparison of our results (purple circles) to equilibrium constants determined from other laboratory studies. In this and subsequent figures, our data were binned in 5 K intervals, with each $K_{\text{eq}}$ point plotted at the average temperature of the data collected within the binning interval. The presentation of binned data is for improved figure clarity only; all data analyses were performed with the 87 independent experiments (Table 1). Notably in Figure 4, the $K_{\text{eq}}$ values reported in this work are typically smaller than prior evaluations of $K_{\text{eq}}$ at warmer temperatures but match well with the colder temperature observations of Hume et al. (2015) for those data points with overlap (228 – 250 K). Moreover, our third law fit matches well with the results of Hume et al. (2015) when extrapolated to 200 K.

The ratio of $K_{\text{eq}}$ values from prior laboratory studies relative to $K_{\text{eq}}$ calculated from equation (3) is shown in Figure 5. The $1\sigma$ ($B \pm 25$ K) and $2\sigma$ ($B \pm 50$ K) error bounds from this work are plotted as shaded gray tones. The $K_{\text{eq}}$ values from previous laboratory studies were derived using either UV absorption spectroscopy of equilibrium mixtures of ClO and ClOOCl or by determination of the individual forward and/or reverse kinetic rates of dimerization, reaction (R1). Experimental data from these previous studies are shown as circles and triangles, respectively. As evident in Figure 5, there is much greater variation in the determinations of $K_{\text{eq}}$ using kinetics methods.

The experiments of Cox and Hayman (1988) and Hume et al. (2015) were performed by UV analysis. The results of Hume et al. (2015) lie almost entirely within our $1\sigma$ error interval. Though the individual results of Cox and Hayman exhibit significant scatter and some measurements exceed the $2\sigma$ error reported here, an ordinary least-squares third law fit of their results using the JPL-recommended $A$ parameter (Burkholder et al., 2015) remains within our $1\sigma$ error boundaries.

Nickolaisen et al. (1994) and Ferracci and Rowley (2010) used flash photolysis/UV absorption spectroscopy to determine the kinetic rates of the individual reactions in order to determine $K_{\text{eq}}$. Though these two studies agree with each other in trend and magnitude, they both exhibit significant departures from our results, possibly due to secondary reactions given the high concentrations of ClO and Cl$_2$O employed in those studies.

Bröske and Zabel (2006) investigated the kinetics of the ClOOCl dissociation reaction and estimated $K_{\text{eq}}$ values using JPL 2002 kinetics (Sander et al., 2003) for the forward reaction (R1). A reanalysis of their results using JPL 2015 kinetics (Burkholder et al., 2015) is plotted as binned averages in Figure 5 (orange triangles, $P < 30$ mbar; brown triangles, $P > 30$ mbar). A reanalysis of the high-pressure results of Bröske and Zabel (2006) also provides a third law fit ($K_{\text{eq}} = 2.16 \times 10^{-27} \times 10^{(8498 K/T)}$ cm$^3$ molecule$^{-1}$ that resides within our $2\sigma$ error limits. The discrepancy between the experiments of Bröske and Zabel (2006) conducted at higher pressures and lower pressures is discussed in depth in their work.

Horowitz et al. (1994) examined the loss rate of ClO while monitoring the kinetics and branching ratio of the ClO + ClO reaction and provide a single-point estimate of $K_{\text{eq}}$ at 285 K that is within $1\sigma$ of the value determined at that temperature in this work. However, the $K_{\text{eq}}$ values of Boakes et al. (2005) using flash photolysis/UV absorption spectroscopy, Ellermann et al. (1995) using pulsed radiolysis/UV absorption spectroscopy, and Plenge et al. (2005) via mass spectrometric determination of the ClO-OCl bond strength, all lie outside the $2\sigma$ error limits from this work.

Figure 6 provides a comparison between observational determinations of $K_{\text{eq}}$ in the atmosphere and an extrapolation of $K_{\text{eq}}$ from this work to 190 K. The determination of $K_{\text{eq}}$ by Avallone and Toohey (2001), an analysis of AASE I and AASE II data in which mixing ratios of ClOOCl were inferred from total Cl$_p$ mass conservation rather than directly measured, agree within error
with the results of this work. Similarly, a determination of \( K_{eq} \) by Santee et al. (2010), informed by ClO mixing ratios retrieved via Aura MLS satellite data and ClOCl mixing ratios calculated from stratospheric modeling, lies in substantial agreement with the \( K_{eq} \) expression derived here. All observed ratios of \([\text{ClOCl}] / [\text{ClO}]^2\) in the Arctic stratosphere for the nighttime ER-2 flight of 3 Feb 2000 during the SOLVE/THESEO campaign (Stimpfle et al., 2004) also agree within the combined 2\( \sigma \)

measurement uncertainties (SOLVE/THESEO measurement uncertainties not plotted for clarity). The \( K_{eq} \) expression from von Hobe et al. (2005), which was based on observations in Arctic winter 2003, deviates substantially from \( K_{eq} \) determined in this work; however, von Hobe et al. (2007) postulate that those previous observations of ClO and ClOCl may not have been in equilibrium and that the ClOCl measurements may have been biased low.

A thermodynamic representation of parameters \( A \) and \( B \) [equation (2)] can be obtained from a manipulation of the van’t Hoff equation. The prefactor \( A \) encodes the standard entropy of reaction change per equation (4), in which the superscript indicates a standard state of one bar, \( R' \) is the gas constant (83.145 cm\(^3\) bar mol\(^{-1}\) K\(^{-1}\)), \( N_A \) is Avogadro’s constant (6.0221 \times 10\(^{23}\) molecules mol\(^{-1}\)), \( R \) is the gas constant in energy units (8.3145 J mol\(^{-1}\) K\(^{-1}\)), \( e \) is Euler’s number, and \( T \) is system temperature.

\[
\Delta S^\circ(298 \text{ K}) = R \ln \left( \frac{N_A A}{e R' T} \right)
\]

The exponential argument \( B \) relates the change in standard enthalpy of reaction as shown in equation (5), with \( R \) and \( T \) as defined above.

\[
\Delta H^\circ(298 \text{ K}) = -R(T + B)
\]

Evaluating equation (5) with our derived value of \( B = 8528 \text{ K} \) results in a \( \Delta H^\circ(298 \text{ K}) \) of -73.4 \pm 0.6 kJ mol\(^{-1}\) for reaction (R1). The uncertainty estimate on this value was obtained by combining the previously determined uncertainty on our \( B \) parameter (\( \pm 25 \text{ K} \)) with estimated uncertainties in the reference cross sections for ClOCl [\( \pm 17\% \) variation near the peak cross section at 248 nm, as reported by Lien et al. (2009), Papanastasiou et al. (2009), and Wilmouth et al. (2009)] and for ClO [\( \pm 3\% \) variation in the peak of the absorption continuum at 264 nm between Sander and Friedl (1988), Simon et al. (1990), and Trolier et al. (1990)] to produce a possible range in \( B \) of 8453 K to 8604 K, as determined from scaled, third law least-squares fits.

Combining our \( \Delta H^\circ(298 \text{ K}) \) value for reaction (R1) with the JPL-recommended \( \Delta H^\circ_f(298 \text{ K}) \) for ClO of 101.681 \pm 0.040 kJ mol\(^{-1}\) (Burkholder et al., 2015) yields a \( \Delta H^\circ_f(298 \text{ K}) \) for ClOCl of 130.0 \pm 0.6 kJ mol\(^{-1}\). This result is in excellent agreement with the JPL-recommended value of 130.1 \pm 1 kJ mol\(^{-1}\).

A least-squares second law fit of \( K_{eq} \), in which both \( A \) and \( B \) are free parameters, yields a determination of \( K_{eq} \) as shown in equation (6).

\[
K_{eq} = (2.14 \pm 1.14) \times 10^{-27} e^{(6530 \pm 123 \text{ K}/T)} \text{ cm}^3 \text{ molecule}^{-1}
\]
An application of equation (4) to the second law prefactor of $2.14 \times 10^{-27}$ cm$^3$ molecule$^{-1}$ produces $\Delta S^\circ(298 \text{ K}) = -147.8 \pm 3.6$ J mol$^{-1}$ K$^{-1}$ for reaction (R1), which agrees with the JPL-recommended value of -147.0 J mol$^{-1}$ K$^{-1}$ [calculated from the $S$ values for ClO and ClOOCl in Table 6-2 of Burkholder et al. (2015)]. The value of $\Delta H^\circ(298 \text{ K})$ of $-73.4 \pm 1.0$ kJ mol$^{-1}$ for reaction (R1) from the second law analysis is in excellent agreement with our results from the third law analysis.

Notably, the equilibrium constant results obtained in this work agree in trend and magnitude with the recently reported $K_{eq}$ values of Hume et al. (2015). This excellent correspondence is illustrated in Figure 7, in which a least-squares third law fit (with each study weighted equally) is presented for a combined data set containing the results of this work and the work of Hume et al. (2015) in ratio to the current JPL recommendation (Burkholder et al., 2015). The combined works span a temperature range of 206 – 301 K, and the resulting $K_{eq}$ is $2.16 \times 10^{-27} \times e^{(8533 \text{ K}/T)}$ cm$^3$ molecule$^{-1}$. This expression deviates from the JPL-recommended $K_{eq}$ value at 200 K by 2%. For illustrative purposes, the uncertainty bounds calculated from this work and the bounds recommended by the current JPL evaluation (Burkholder et al., 2015) are also plotted. The JPL uncertainty, which was scaled to include the warm temperature results of Cox and Hayman (1988) and Nickolaisen et al. (1994), greatly exceeds the scatter of the individual $K_{eq}$ values from the combined dataset of this work and Hume et al. (2015). Our results suggest that the uncertainties in the current JPL recommendation for $K_{eq}$ can be significantly reduced.

4 Conclusions

The thermal equilibrium governing the association of ClO and dissociation of ClOOCl was investigated in a custom-built discharge-flow reactor by UV spectroscopy between the temperatures of 228 – 301 K. The selected temperature range allowed us to bridge the warmer temperature regime where nearly all previous laboratory studies of $K_{eq}$ have been performed and the recent colder temperature work of Hume et al. (2015). A third law fit of our $K_{eq}$ results deviates from some prior laboratory studies but demonstrates excellent agreement with the work of Hume et al. (2015) and with the currently recommended parameters in the JPL compendium (Burkholder et al., 2015). The agreement between our third law and second law analyses lends further confidence to the results reported herein. Our calculated enthalpy of formation for ClOOCl from the slope of the van’t Hoff plot is in excellent agreement with the recommended value (Burkholder et al., 2015).

The current JPL-recommended error bounds for the ClO-ClOOCl equilibrium constant are large (Burkholder et al., 2015), exceeding 50% at 200 K. The excellent correspondence between the $K_{eq}$ results from this work and Hume et al. (2015) lends confidence to the established parameterization of the JPL data evaluation (Burkholder et al., 2015), suggesting that prescribed error intervals for this reaction can be reduced.

Author contributions. JEK and DMW designed, constructed, and operated the experiment and analyzed and reported the data described in this work.
**Competing interests.** The authors declare no competing financial interest.

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Figure 1. Schematic of the discharge-flow absorbance experiment. Dilute chlorine gas in helium flows through a microwave discharge to form Cl radicals. Dilute ozone in nitrogen is then injected to produce ClO radicals. Self-reaction of ClO occurs in the cold reaction cell to form ClOOCl. When utilized, the cold trap provides for halogen oxide purification. ClO/ClOOCl equilibrium is established in the equilibrium cell, which is held at the same temperature as the absorption cell. The gas mixture is then characterized via UV spectroscopy in the absorption cell using software developed in-house.
Figure 2. Deconvolution of experimental absorbance spectra to component contributions. (a) Experimental spectrum obtained at 230 K. (b) Experimental spectrum obtained at 300 K. In each panel, the raw absorbance spectrum appears as a black line, and the fitted components and residual from the fit are shown. Fitted concentrations (molecules cm$^{-3}$) are given for various component gases in the legend. The residual is offset by 0.14 A.U. for clarity. OCIO and Cl$_2$O$_3$ contributions are small, especially at low temperatures, and are omitted from the figure.
Figure 3. Experimental $K_{eq}$ values as a function of $1000/T$. The black trace is a third law fit of the orange circles, $K_{eq} = 2.16 \times 10^{-27} e^{(6528 \text{ K}/T)} \text{ cm}^3 \text{ molecule}^{-1}$. Individual values are enumerated in Table 1.
**Figure 4.** $K_{eq}$ values as a function of $1000/T$. Data from this work are plotted as purple circles; for figure clarity, multiple samples (Table 1) are binned and averaged in 5 K intervals. The black trace is the third law fit as determined in Figure 3, $K_{eq} = 2.16 \times 10^{-27} e^{(8528 \, K/T)}$ cm$^3$ molecule$^{-1}$. The colored markers are $K_{eq}$ values from prior laboratory studies as reported in the literature.
Figure 5. Ratio of $K_{eq}$ values from prior laboratory studies to $K_{eq}$ determined in this work ($K_{eq} = 2.16 \times 10^{-27} e^{(8528 K/T)} \text{ cm}^3 \text{ molecule}^{-1}$) as a function of $1000/T$. Circles indicate studies in which $K_{eq}$ was measured with UV spectroscopy of equilibrium mixtures, while triangles indicate works in which $K_{eq}$ was determined from individual reaction kinetic rates. Dark gray shading encompasses the total estimated 1σ error from this study and light shading encompasses the 2σ error.
Figure 6. Comparison of extrapolated $K_{eq}$ values from the third law fit in this work ($K_{eq} = 2.16 \times 10^{-27} e^{(8528 K/T)}$ cm$^3$ molecule$^{-1}$) to atmospheric observations. $K_{eq}$ (solid black) and error boundaries (gray shaded regions) determined in this study are extrapolated to the temperature range of 190 – 203 K. Expressions for $K_{eq}$ derived from previous atmospheric measurements are presented as dashed lines. Observations of [ClOCl] / [ClO]$^2$ from the nighttime ER-2 flight on 3 Feb 2000 in the SOLVE/THESAO mission out of Kiruna, Sweden are indicated as orange diamonds.
Figure 7. Ratio of a weighted third law fit of $K_{eq}$ (black dashed line) determined from a combination of this work (circles) and Hume et al. (2015) (triangles) to the JPL compendium recommended value (Burkholder et al., 2015). Error intervals as reported in this work (darker blue = 1σ, lighter blue = 2σ) and as recommended by JPL-2015 (red cross hatch).
### Table 1. Experimental conditions and $K_{eq}$ values

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*Temperature reported to five significant figures for purposes of fitting.