Coupled Air Quality and Boundary-Layer Meteorology in Western U.S. Basins during Winter
Design and Rationale for a Comprehensive Study


ABSTRACT: Wintertime episodes of high aerosol concentrations occur frequently in urban and agricultural basins and valleys worldwide. These episodes often arise following development of persistent cold-air pools (PCAPs) that limit mixing and modify chemistry. While field campaigns targeting either basin meteorology or wintertime pollution chemistry have been conducted, coupling between interconnected chemical and meteorological processes remains an insufficiently studied research area. Gaps in understanding the coupled chemical–meteorological interactions that drive high-pollution events make identification of the most effective air-basin specific emission control strategies challenging. To address this, a September 2019 workshop occurred with the goal of planning a future research campaign to investigate air quality in western U.S. basins. Approximately 120 people participated, representing 50 institutions and five countries. Workshop participants outlined the rationale and design for a comprehensive wintertime study that would couple atmospheric chemistry and boundary layer and complex-terrain meteorology within western U.S. basins. Participants concluded the study should focus on two regions with contrasting aerosol chemistry: three populated valleys within Utah (Salt Lake, Utah, and Cache Valleys) and the San Joaquin Valley in California. This paper describes the scientific rationale for a campaign that will acquire chemical and meteorological datasets using airborne platforms with extensive range, coupled to surface-based measurements focusing on sampling within the near-surface boundary layer, and transport and mixing processes within this layer, with high vertical resolution at a number of representative sites. No prior wintertime basin-focused campaign has provided the breadth of observations necessary to characterize the meteorological–chemical linkages outlined here, nor to validate complex processes within coupled atmosphere–chemistry models.

KEYWORDS: Chemistry, atmospheric; Greenhouse gases; Aircraft observations; Field experiments; Mountain meteorology

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Winter episodes of high aerosol concentrations occur frequently in urban and agricultural basins and valleys across the globe [e.g., Yamuna Basin, India (Tiwari and Kulshrestha 2019); Tokyo Basin, Japan (Osada et al. 2019); Taiyuan Basin, China (Miao et al. 2018); and the San Joaquin and Salt Lake Basins, United States (Whiteman et al. 2014; Zhang et al. 2020)]. These episodes may last from several days to several weeks and often arise due to the development of persistent cold-air pools (PCAPs), within which lateral and vertical mixing are inhibited due to sheltering by surrounding topography and a stable temperature profile (Dorninger et al. 2011; Reeves et al. 2011; Lareau et al. 2013; Sheridan et al. 2014; Holmes et al. 2015; Sun and Holmes 2019; Ivey et al. 2019; Sun et al. 2020). While a number of field campaigns targeting either wintertime basin meteorology (e.g., Lareau et al. 2013; McCaffrey et al. 2019) or wintertime pollution chemistry (e.g., Brown et al. 2013; Franchin et al. 2018; Young et al. 2016) have been conducted in the United States, only a few of these campaigns have explicitly
considered coupling between interconnected chemical and meteorological processes (e.g., Baasandorj et al. 2017; Prabhakar et al. 2017; Salvador et al. 2021). The upcoming Alaskan Pollution and Chemical Analysis (ALPACA) is specifically targeting this knowledge gap in cold and dark conditions.

Current gaps in our understanding of the coupled chemical–meteorological interactions that result in high-pollution events in many basins worldwide make identification of the most effective air-basin specific emission control strategies challenging. Meteorological processes (thermodynamic, radiative, and dynamical) influence both pollution accumulation, dispersion, and transport and aerosol pollution chemistry, while chemical processes in turn influence radiative transfer, cloud formation, and mixing processes. Figure 1 presents a graphical illustration of some of the coupled chemical–meteorological processes that occur in basins. Some key meteorological processes that control the formation, duration, and breakdown of PCAPs include synoptic drivers such as high pressure and associated subsidence, which can precipitate elevated thermal inversions; warm air advection aloft and large-scale winds and turbulent mixing, alongside local drivers such as the surface energy and radiation budget, which strongly influence formation and dissipation of surface-based thermal inversions; characteristics of the underlying surface (e.g., snow cover, water, urban or non-urban landscape); low clouds and fog; and local boundary layer flows near the surface. In turn, the location and types of urban emissions, aerosol formation and growth processes, and chemical cycling processes influence and are influenced by the ambient meteorology (Fig. 1). The unique basin topography (e.g., slope, how enclosed the basin is, and the size of basin) also play an important role in modulating the rate of pollutant buildup and vertical profiles of temperature and moisture. The interactions between these numerous meteorological processes regulating the frequency, location, and speed of chemical processes in PCAPs, and complex wintertime chemistry, has not yet been observed with sufficient detail to provide satisfactory understanding of these complex pollution episodes and their evolution in time and space

Ammonium nitrate ($\text{NH}_4\text{NO}_3$) is a major component of aerosol mass in many polluted boundary layers during wintertime episodes (e.g., Womack et al. 2019; Fu et al. 2020; Kelly et al. 2018; Kim et al. 2014; Aksoyoglu et al. 2017; Schaap et al. 2004). Organic aerosol (OA) can also contribute substantially (Chen et al. 2018). In some basins, the $\text{NH}_4\text{NO}_3$ contribution increases with the total aerosol loading while in others OA dominates at high loadings, often with differences between day and night. Understanding the factors that govern the evolution of pollution within a basin is critical for implementing effective emission control policies. Precursor emissions and chemical transformations leading to $\text{NH}_4\text{NO}_3$ formation are an area of ongoing research, as are a number of processes controlling OA production and loss. While OA formation has been extensively studied in warm seasons with large emissions of volatile organic
compounds (VOCs, precursors to secondary organic aerosol) often from biogenic sources, characterization for winter urban environments is lacking. Slower photochemistry occurs in the winter, and biogenic and evaporative emissions tend to decrease with temperature, suggesting that these emissions play a smaller role in wintertime OA formation. However, recent evidence of rapid and widespread OA formation (Shah et al. 2019; Schroder et al. 2018) indicate that urban wintertime OA formation is an important contributor to PM$_{2.5}$ mass. Sulfate is a small fraction of PM$_{2.5}$ mass during winter in basins such as the Salt Lake Valley but is more important elsewhere (Wang et al. 2016). Mechanisms leading to winter sulfate oxidation are a topic of current interest for which detailed studies in the western United States may serve as a test bed. Consequently, it is imperative to better understand the mechanisms that drive local nitrate, organic and sulfate formation, alongside the emission sources of key precursor gases—e.g., nitrogen oxides, ammonia, SO$_2$, and VOCs. This is particularly important given recent findings showing the importance of urban VOC emissions from evaporative sources relative to those from fuel use (McDonald et al. 2018). Process-level understanding also requires investigation of radical cycling involving VOCs and NO$_x$ in the winter and its relationship to high pollutant levels in stagnant boundary layers.

In recent decades, overall aerosol concentrations have declined in the United States owing to emissions changes driven by regulatory policies (Bennett et al. 2019). However, a recent study has conclusively demonstrated that particulate levels across the United States are associated with mortality impacts and loss of life expectancy, with the highest rates observed within the San Joaquin Valley in California (Bennett et al. 2019). Furthermore, particulate matter with diameters less than 2.5 microns (PM$_{2.5}$) within major urban areas across the United States has declined more slowly in winter than in summer, and average winter levels are now higher in winter than in summer (Chan et al. 1994). These trends are particularly pronounced in basins across the western United States, where wintertime aerosol concentrations regularly approach or exceed regulatory standards (Green et al. 2015). Gaps in understanding the coupled chemical–meteorological interactions that result in high-pollution events may preclude identifying the most effective emission control strategy for a given air basin. With this challenge at the forefront, a workshop took place in September 2019 at the University of Utah with the goal of planning a future winter research campaign to investigate mountain basins of the western United States. With funding from the National Science Foundation (NSF) Atmospheric Chemistry Program and the National Oceanic and Atmospheric Administration (NOAA) Atmospheric Chemistry, Climate and Carbon Cycle Program, the workshop brought together ~120 air quality experts and meteorologists from across the globe, representing 50 institutions and five countries.

As summarized in this article, the workshop outlined the rationale and design for a comprehensive study that couples atmospheric chemistry and meteorology for wintertime poor air quality episodes in mountain basins across the western United States. The campaign framework is laid out in the next section. The existing uncertainties and opportunities for this campaign are summarized in the following sections, organized by science subthemes. Finally, an integrated perspective on measurements and modeling is presented, along with next steps, in the final section.

**Design of the research study**

The two western U.S. regions with the most severe winter aerosol pollution are northern Utah (comprised of the Salt Lake, Utah, and Cache Valleys) and the San Joaquin Valley (SJV) in California. The workshop concluded that while winter meteorology and a mix of urban and rural emissions affect basins across the western United States (see Fig. 2), a comprehensive study should focus on these two regions, which exhibit contrasting aerosol chemistry, basin topography, climate, agricultural practices, and meteorological–chemical interactions. In the Salt Lake City area, ammonium nitrate is the major contributor to wintertime PM, whereas
in the SJV primary and secondary organic aerosol (SOA) are also major contributors (Baasandorj et al. 2017; Lurmann et al. 2006; McDuffie et al. 2019; Chow and Watson 2002). Furthermore, historical databases and previous studies provide context for an air quality study in both of these regions, as PM regulatory standard exceedances in wintertime are common to both. Finally, strong science capacity is available in these areas to conduct this research. Figure 2 shows a long-term record of \( \text{PM}_{2.5} \) at the Hawthorne monitoring site in Salt Lake City, the SJV (Bakersfield), and, for comparison, a site in California’s South Coast basin (Riverside–Ruidous). The decreasing trend in \( \text{PM}_{2.5} \) in the South Coast basin is visually apparent from Fig. 2, in contrast to the SJV and Salt Lake City sites, which still regularly experience regulatory exceedances during winter, despite modestly decreasing \( \text{PM}_{2.5} \) trends (Green et al. 2015).

The onset and evolution of wintertime pollution episodes in the SJV of California differs in many respects from those in the colder, smaller and frequently snow-covered Intermountain West basins. Additionally, western U.S. mountain valleys show significant variability in emissions. Emissions of nitrogen oxides (\( \text{NO}_x \)), VOCs, and ammonia (\( \text{NH}_3 \)) are highly dependent on the nature of the agricultural and industrial sectors in each valley (Kelly et al. 2013; Wang et al. 2015), and their chemical transformations are dependent on many meteorological and topographical factors (Green et al. 2015; Wang et al. 2015; Kleeman et al. 2005; Pusede et al. 2016).

The proposed field program focuses on understanding how variations in coupled meteorological and chemical processes contribute to the production, transformation, cycling, and destruction of chemical species in each locale. This paper highlights the need for acquisition of chemically and meteorologically comprehensive datasets using airborne platforms with sufficient range, coupled to extensive surface-based measurements that provide continuous, chemically detailed data at ground level where human exposure occurs. Sampling of the boundary layer at high vertical resolution will provide representative profiles that reflect regional topographical, meteorological, and emissions variability. The chemical measurements will be combined with comprehensive meteorological measurements to characterize the influences
of a wide range of meteorological and land surface processes and parameters (e.g., transport and mixing, surface albedo) and topography on chemical processes during the PCAP episode.

**Understanding the coupling between “cold-air pool” meteorology and air quality**

The complex cold-air pool basin meteorology that impacts pollutant dispersion and air pollution chemistry remains an active area of research worldwide (Giovannini et al. 2021). Many of the meteorological processes and their effects on air pollution transport and chemistry are still not well documented, understood, or adequately modeled despite having contributed to poor air quality in western U.S. basins for over a century (Fig. 1: see Lareau et al. 2013; Giovannini et al. 2021; Lighthall and Capitman 2007). Figure 1 illustrates the coupling of chemical and meteorological processes wherein the atmospheric state impacts the atmospheric chemistry and, in some cases, results in feedbacks between the two (e.g., aerosols affect cloud properties and lifetime and can absorb shortwave solar radiation, which leads to alteration of the vertical temperature profile and hence pollutant vertical transport). Transport and mixing processes, insolation, and microphysical cloud processes all affect the type and extent of different chemical processes. In turn, weaker feedbacks between aerosol loading and cloud chemistry can have significant impact—via radiative feedbacks (e.g., longwave energy transfer back to the surface from low clouds, net decreases in incoming shortwave radiation into the PCAP due to reflection of the shortwave solar energy back to space at cloud top)—on the vertical stratification and thus vertical mixing processes within the PCAPs.

The complexity of interactions between topographical and other physical characteristics of basins, meteorological, and chemical processes introduced in Fig. 1 leads to pollution with characteristics, severity, and duration differing both in time and space and across different basins. For example, variations in the vertical temperature profile and magnitude of turbulence and mixing from solar heating or wind shear can dramatically impact the amount of entrainment/dilution of pollutants and vertical layering of pollutant precursors within the basin atmosphere.

The workshop identified four critical couplings processes between meteorology and chemistry requiring investigation as part of the proposed field program: 1) surface fluxes of energy and momentum and chemistry; 2) moisture and fog, and heterogeneous, multiphase, and aqueous-phase chemistry; 3) PCAP vertical thermodynamic profiles and vertical chemical profiles, and 4) meteorology and chemistry associated with thermally and dynamically forced exchange processes. No prior wintertime field campaign has provided the breadth of observations necessary to analyze the meteorological–chemical linkages outlined here or to validate complex processes within coupled atmosphere–chemistry models.

A critical tool for both basic research and air pollution control strategies are coupled meteorological and chemical models. Development of useful guidance for the research and regulatory communities requires improvements to both meteorological and chemical components of these models (Giovanni et al. 2021). To improve meteorological model simulations of PCAPs, considerable work is underway or has been conducted in recent years (e.g., Saide et al. 2011; Lareau and Horel 2015; Ahmadov et al. 2015; Saide et al. 2016; Tran et al. 2018; Sun and Holmes 2019; Kelly et al. 2018; Sun et al. 2020). Some of the key meteorological processes that are very difficult to model in stable wintertime boundary layers include vertical temperature and humidity structure, cloudiness, turbulent mixing, and boundary layer flows (Baklanov et al. 2011; Holmes et al. 2015). Improvements in and testing of the representations of heterogeneous chemical processes (Holmes et al. 2019; S. S. Brown et al. 2006), links between aerosol phase and gas–particle equilibration and partitioning (Shiraiwa et al. 2013b; Zaveri et al. 2018), secondary organic aerosol life cycles (Cappa et al. 2016), and organic compound oxidation pathways (Bianchi et al. 2019), especially as they occur within colder wintertime environments, is needed to accurately
understand and predict both aerosol and gas-phase abundance and composition, discussed further later on.

The coupled meteorological and chemical models parameterize vertical mixing using relationships between turbulent fluxes and mean thermodynamic profiles that have significant impacts on predicted atmospheric composition. It is therefore paramount that meteorological (e.g., temperature and wind speeds to quantify mixing and transport, relative humidity and downwelling radiance to estimate liquid water path and boundary layer moisture) and chemical vertical profile measurements be collocated with observations of the radiative and turbulent components of the surface energy balance. This approach will allow a detailed quantification of mass, moisture, heat, and chemical budgets within basins. It is also important that variations in the surface energy balance across the basins be well captured using flux sites and satellite data. Important land surface parameters to observe include the depth and age of snow (which can impact the amount of reflected solar radiation and rates of photochemical reactions), and soil temperature and moisture (which can impact fog formation and vertical mixing processes, both which impact chemistry).

Sufficient data, both temporally and spatially, are needed to ensure that spatial gradients are captured across the basins of interest to form a three-dimensional representation of the atmospheric state for both chemical and meteorological properties. Chemical processes driven by mixing also need to be resolved alongside the meteorological measurements for turbulence, mixing, and transport, to allow linking of these two processes. A holistic, interdisciplinary, and multiagency approach will be used in this study where existing infrastructure such as National Weather Service daily rawinsonde launches, weather stations from public and private sectors available from MesoWest (Horel et al. 2002), wind sodars, lidar profilers, and ceilometers will be supplemented with instrumentation dedicated to this field study.

One study design approach is a process-focused deployment, targeting regions of interest within a basin to investigate and quantify coupled chemical–meteorological processes. Three examples of targeted meteorological processes and their impacts on the chemistry include 1) interbasin exchange, 2) sidewall- and canyon-flow transport, and 3) vertical layering and exchange (Fig. 1).

In addition to using frequent rawinsonde launches and vertical-profiling wind lidars, sodars, ceilometers, and acoustic-sounding systems at fixed locations, use of mobile vertical profilers and tall buildings as instrument platforms can help resolve the vertical structure of the basin atmosphere. Mobile ground-based systems measuring both meteorology and chemistry, such as the California Air Resources Board (CARB) Mobile Measurement Platform (Park et al. 2011) or the Utah Transit Authority Light-Rail Transit Express (TRAX) air quality light-rail train (Mitchell et al. 2018a,b; Mendoza et al. 2019), will fill gaps in the fixed-instrumentation networks. Mobile ground-based systems driven up canyons and slopes can provide measurements along topographic sidewalls, and drones can obtain shallow but highly resolved vertical profiles in areas with limited topography. Coupled meteorological data (including turbulence and the surface energy balance) collected on towers and high buildings at multiple locations within the basins would also provide valuable observations in the lowest 100 m of the PCAP atmosphere. A potential exists for deploying airborne Doppler wind lidar to provide similar vertical wind-flow information along a 2D transect, during times and in locations where fog and low stratus do not impede operations. Lidar-based vertical wind profiles at basin boundaries would greatly assist in quantifying background pollution and meteorological parameters as well as transport processes within a single basin and between adjacent basins.

The field deployment will also include modeling teams, using tools that span the scales from large-eddy simulations (meters resolution) to regional coupled meteorological and chemical
models (kilometers resolution), to forecast PCAP events and contribute to flight planning; this will help ensure the measurements address the modelers’ needs. Complementing research-grade models and forecasts, operational air quality forecasts used to warn the public about pollution events will be evaluated to identify deficiencies and accelerate the transition of any improvements from research to operations.

**Emissions of relevant short-lived pollutants**

Within basins that experience adverse air quality in winter, direct emissions of aerosols are of interest, as well as emissions of short-lived gases, including reactive nitrogen species (e.g., $\text{NO}_x$ and $\text{NH}_3$) and intermediate-volatility/volatile organic compounds (I/VOCs; e.g., hydrocarbons, amines, oxidized and reduced sulfur compounds, and oxygenated organic compounds). Such emissions are largely from urban, agriculture, and biomass burning sources. Here, residential wood combustion is considered an urban emissions source, and biomass burning is limited to wildland fires (wildfires and prescribed burns). Different sources emit at different elevations above the surface, with consequent implications for the fate of emitted compounds and their impacts on surface air quality. For example, emissions from vehicles occur at the surface, whereas emissions from some industrial sources or the power sector may occur at elevations tens to hundreds of meters above the surface.

Mobile sources and residential wood combustion contribute significantly to primary and secondary aerosols in Salt Lake City in wintertime (e.g., Kelly et al. 2013). Similar sources contribute to elevated aerosol concentrations in the SJV; in addition to local transportation and residential wood combustion, cooking was identified as significant source of aerosols, particularly in the evening (Young et al. 2016). While emissions reductions from passenger vehicles have led to a continuing decline in on-road vehicle contributions to $\text{NO}_x$ and aerosols (Dallman and Harley 2010; Bishop et al. 2012), the contributions of older and off-road vehicles to $\text{NO}_x$ and aerosols are not well quantified. In addition, the temperature dependence of mobile source emissions is not well constrained for real-world conditions. Previous in-use laboratory testing has shown that lower ambient temperatures can lead to higher gaseous and particulate mass emissions from gasoline and diesel vehicles, and to changes in chemical composition (Zielińska et al. 2004). In addition, vehicle operation specific to winter conditions, such as changes in the frequency of hot versus cold starts and an increase in idling, can affect mobile source emissions and ambient concentrations. Finally, $\text{NH}_3$ emissions from selective catalytic reduction systems may increase in winter, given the increase in $\text{NH}_3$-forming deposits with decreases in ambient temperature (Strots et al. 2010).

Reductions in mobile source I/VOC emissions spanning several decades (e.g., Warneke et al. 2012) have led to an increase in the relative importance of other, understudied sources of I/VOCs including cooking and volatile chemical products (VCPs) (McDonald et al. 2018). A large fraction of VCP emissions occur inside buildings. More restrictive air exchange between buildings and ambient air during the winter to conserve energy may affect VCP emissions (Pagonis et al. 2019); currently, there are few constraints regarding the seasonality in VCP emissions. Evaporative emissions of transportation fuels generally decrease as ambient temperature decreases (Harley et al. 1992; Rubin et al. 2006), which also may be observed for VCPs.

Regarding residential wood burning, in areas affected by poor wintertime air quality, efforts have been made to reduce solid fuel combustion (wood, wood pellets, coal, etc.) through implementation of burn restrictions when poor air quality is forecast, and incentives to exchange wood-burning devices for natural gas heating options. Such strategies appear to be reducing the prevalence of solid-fuel burning, including wood (Kotchenruther 2020); however, it has been demonstrated that primary and secondary aerosols originating from residential wood burning persist, even when solid-fuel burning is prohibited (Kotchenruther 2020). In addition,
solid fuels used in wood stoves are poor in nitrogen, which affects the emissions of nitrogen-containing VOCs, NO\textsubscript{x}, and NH\textsubscript{3} (Coggon et al. 2016).

Agricultural emissions have been directly linked to atmospheric aerosol formation (Bauer et al. 2016), mainly via the dominant contribution of agriculture to NH\textsubscript{3} emissions both globally and in the study region (Bouwman et al. 1997; Paulot 2014). Uncertainties in quantifying the contribution of NH\textsubscript{3} to aerosols in winter are largely associated with parameterizations of emissions as a function of temperature. NH\textsubscript{3} emissions from manure vary nonlinearly with temperature, with higher temperatures increasing volatility (e.g., Hempel et al. 2016). For animals in heated housing, there is concern that process-based NH\textsubscript{3} predictions for ambient temperature may underpredict winter emissions (Moravek et al. 2019). In addition to NH\textsubscript{3}, several recent studies have pointed to the importance of fertilized agricultural soils as a source of NO\textsubscript{x} (Almaraz et al. 2018; Trousdell et al. 2016), which could have a particularly large impact in California where crop fertilization occurs in the winter months.

A number of individual agricultural VOCs have been identified, and include aliphatic and aromatic hydrocarbons, phenols, alcohols, aldehydes, ketones, esters, carboxylic acids, terpenes, heterocyclic compounds, amines, and reduced sulfur compounds (e.g., Blunden et al. 2005; Filipy et al. 2006; Shaw et al. 2007; Turan et al. 2007; Trabue et al. 2008; Rumsey et al. 2012; Rumsey and Aneja 2014). While past studies suggested that, relative to other anthropogenic sources, agricultural VOCs do not contribute significantly to \textit{O}_3 production, recent studies indicate their role has been overlooked (Pusede et al. 2014; Parrish et al. 2017; Trousdell et al. 2016). Several of these identified compound classes are known aerosol precursors, particularly amines and reduced sulfur compounds. Amines have been reported as a significant component of the carbonaceous fraction of aerosol observed during cold wintertime inversion conditions in western agricultural valleys, in Utah (Silva et al. 2008), and in Washington (Bottenus et al. 2018). Reduced sulfur compounds (e.g., dimethylsulfide, dimethyldisulfide, and methanethiol) have received less attention (Trabue et al. 2008; Rumsey et al. 2014), but agricultural contributions of such compounds can be significant (Shaw et al. 2007).

In certain states and regions, there may be unique agricultural sectors unaccounted for in emissions inventories and air quality models. For example, the state of Utah is the number two producer of mink pelts in the United States, but such emissions are lacking in the National Emissions Inventory because there are no data available. Also, rapid changes in agricultural source emissions may occur owing to the recent expansion of anaerobic digesters. California alone has dozens of digesters under construction for the dairy industry and they are being installed in other states as well. Few studies have been conducted to see how the digester processes might impact non–greenhouse gas (GHG) emissions.

Wildland fire emissions likely contribute to the elevated PM\textsubscript{2.5} concentrations observed during the winter months. Fire activity in winter has become more prominent in the western United States. For example, the Thomas Fire in California, which started in early December, became one of the largest wildfires recorded in California (Kolden and Abatzoglou 2018). Higher summer temperatures and decreased precipitation in the fall could potentially continue to extend the western United States fire season into the winter (Guzman-Morales and Gershunov 2019). Transported emissions from these late fall/early winter wildfires in the western United States could then contribute to enhanced PM\textsubscript{2.5} concentrations in the SLC region.

\textbf{Winter atmospheric chemical cycles}

Chemical transformations and subsequent phase partitioning of primary air pollutants such as NO\textsubscript{x}, VOCs, and reduced nitrogen are responsible for the conversion of these primary emissions into aerosols. While the daily integrated solar radiation available to drive winter photochemistry is generally lower than summer, multiphase chemical reactions that are
prevalent in winter can lead to photolabile radical sources that drive aerosol formation. The nature and magnitude of radical sources impacts the sensitivity of chemical oxidation cycles to primary emissions of NO\textsubscript{x} and VOCs, leading to fundamental shifts in the prevalent chemical regime during winter. The shallow, stable boundary layer associated with PCAPs tends to concentrate primary emissions, further shifting the sensitivities of oxidation cycles. This section examines some of the features of winter chemical oxidation cycles in polluted boundary layers and identifies major uncertainties.

Snow cover is conducive to the formation and intensification of PCAP conditions due to enhanced radiative cooling at the snow surface that strengthens the low-level wintertime stable layer. In the SLV, for example, PM\textsubscript{2.5} exceedances are observed 4 times as frequently on snow covered days compared to those without snow cover (Whiteman et al. 2014). The effect of snow cover is complex, since it changes both the boundary layer dynamics and atmospheric chemistry. Reflection of solar radiation by snow increases actinic flux and photochemical reaction rates. Surface snowpacks also directly impact atmospheric composition by serving as a sink/reservoir for atmospheric trace gases and particles upon deposition and as a source from reactions on and within the snow grain surface (Grannas et al. 2007). For example, nitrate deposited on the snowpack undergoes photolysis to produce NO\textsubscript{x} and HONO released to the air above (Chen et al. 2019; Honrath et al. 2000; Michoud et al. 2015; Zatko et al. 2016). Snowpack photochemistry also contributes to near-surface OH through the production and subsequent photolysis of H\textsubscript{2}O\textsubscript{2} and carbonyls, including formaldehyde, acetaldehyde, and acetone (Couch et al. 2000). Dinitrogen pentoxide (N\textsubscript{2}O\textsubscript{5}) reactions on saline snow grains can result in CINO\textsubscript{2} formation, with the snowpack serving as a net source or sink of CINO\textsubscript{2} depending on temperature (Wang et al. 2020).

The presence of large surface area in the particulate phase and in fogs also increases heterogeneous reaction rates during winter, altering levels of several criteria air pollutants including ozone (O\textsubscript{3}), nitrogen dioxide (NO\textsubscript{2}), SO\textsubscript{2}, and particulate matter (Sarwar et al. 2012; Sarwar et al. 2014). One important heterogeneous reaction known to contribute to accumulation of aerosol mass is that of N\textsubscript{2}O\textsubscript{5} on deliquesced aerosol particles. N\textsubscript{2}O\textsubscript{5} chemistry occurs primarily during darkness, and is favored at lower temperatures and elevated NO\textsubscript{x} levels, which are characteristic of urban areas during winter (Chang et al. 2011). This reaction regulates the wintertime NO\textsubscript{x} lifetime and leads to the formation of nitric acid (HNO\textsubscript{3}) (Chang et al. 2011), which contributes to particulate nitrate levels in the presence of ammonia (McDuffie et al. 2019). Nitryl chloride (CINO\textsubscript{2}), a photolytic source of chlorine radicals (Cl\textsuperscript{-}) and NO\textsubscript{2}, is produced from N\textsubscript{2}O\textsubscript{5} where chloride-containing particles are present (Osthoff et al. 2008). Atomic chlorine is highly reactive and initiates VOC oxidation cycles and SOA formation (Wang and Hildebrandt Ruiz 2018). The fate of the reactive uptake of N\textsubscript{2}O\textsubscript{5} and subsequent impacts on oxidant and PM concentrations depend on individual particle surface composition (Gaston and Thornton 2016; McNamara et al. 2020), which remains poorly constrained.

A second important wintertime heterogeneous and multiphase reaction is the production of nitrous acid (HONO), a radical reservoir whose photolysis may be a large OH source that affects urban air quality (e.g., Fu et al. 2019). Numerous heterogeneous and multiphase HONO sources have been proposed, such as photochemical generation from snowpacks (Chen et al. 2019) and photolysis of nitrate aerosol (Ye et al. 2017), although the latter remains controversial (Romer et al. 2018). Given the variety of potential HONO sources and associated uncertainties, future measurements are needed to ascertain dominant formation pathways.

Dissolution of trace gases and subsequent aqueous-phase reactions within fog droplets and deliquesced aerosol can also lead to secondary aerosol formation, including sulfate and high molecular weight organic compounds (Ervens et al. 2011). Previous SJV wintertime studies have shown aqueous-phase formation of high molecular weight organic compounds
(Qin and Prather 2006), as well as hydroxymethanesulfoante (Whiteaker and Prather 2003). This compound has been suggested as a significant winter sulfate source in highly polluted regions such as China (Song et al. 2019). Aqueous phase sulfur oxidation driven by NO$_2$ at high aerosol pH has recently been proposed to explain rapid sulfate formation in China (Wang et al. 2011). Fog pH has been increasing in recent years within the SJV, where ammonium nitrate is a major component in the north and sulfate is prevalent in the south (Herckes et al. 2015), providing a potential test of this and other known and proposed aqueous sulfur oxidation mechanisms (Alexander et al. 2009; Calvert et al. 1985; Green et al. 2019). Studies of the multiphase processes remain too sparse to accurately parameterize their contribution to winter air quality.

O$_3$ and NO$_2$ are often collectively termed odd oxygen (O$_x$) and tracked together to determine the growth of O$_3$ beyond its rapid interconversion with NO$_2$ (Wood et al. 2009). Odd oxygen is frequently defined more broadly to include other nighttime reservoirs such as NO$_3$, N$_2$O$_5$, and gas phase HNO$_3$ (S. G. Brown et al. 2006; Liu 1977). Recent work has expanded the definition of O$_x$ further (referred to as O$_{x,\text{total}}$) to include additional reactive nitrogen compounds and particulate nitrate (Womack et al. 2019). Tracking O$_{x,\text{total}}$ allows for the investigation of the role that oxidation plays in the buildup of pollutants in boundary layers. During the Utah Winter Fine Particulate Study (UWFPS) (Baasandorj et al. 2018), O$_{x,\text{total}}$ increased dramatically above the 45 ppbv O$_3$ background during pollution episodes and rapidly became dominated by NO$_2$ and particulate nitrate under conditions of depleted O$_3$. Box modeling demonstrated that the high ratio of NO$_2$ to VOCs in the SLV altered the traditional O$_3$ photochemical cycle (Kleinman 1994; Lin et al. 1988) to primarily form particulate nitrate (Womack et al. 2019). This finding contrasts with the adjacent Uintah Basin, UT, where high concentrations of VOCs from oil and gas extraction with relatively lower NO$_x$ levels have been shown to force the photochemical cycle to produce O$_{x,\text{total}}$ in the form of wintertime O$_3$ (Ahmadov et al. 2015; Edwards et al. 2013; Edwards et al. 2014). These two analyses of air pollution in adjacent basins, SLV and Uintah, demonstrated the important role of photochemistry during winter and the utility of O$_{x,\text{total}}$ as a parameter for evaluating its role in the generation of both O$_3$ and particulate nitrate pollution. The contrast between the two basins shows that particulate nitrate and O$_3$ are closely coupled and can be regarded as chemically identical (Meng et al. 1997). Modeling techniques that have long been used for evaluating mitigation strategies for ozone can also be used for particulate nitrate (Nguyen and Dabdub 2002; Pun et al. 2009). In the wintertime SLV, a NO$_x$–VOC sensitivity isopleth demonstrated that the formation of O$_{x,\text{total}}$ is most sensitive to reductions in VOCs, and is NO$_x$-saturated, meaning that reductions in NO$_x$ emissions would initially increase aerosol pollution (Womack et al. 2019), a result that is counterintuitive since NO$_x$ is also the precursor to particulate nitrate, the dominant component of PM$_{2.5}$ in SLV. However, the NO$_x$–VOC relationship for particulate nitrate must be understood in detail in the context of the competing processes that govern particulate nitrate formation and chemical and meteorologically driven loss, which remain incompletely understood.

**Processes governing particulate matter formation and loss**

The formation and loss of atmospheric aerosols is tightly linked to the partitioning of the chemical constituents between the particle and gas phases, and the multiphase reactions described in the previous section. Observations demonstrate that particulate nitrate and OA dominate wintertime composition in a variety of urban areas within valleys (e.g., Fresno, CA; Salt Lake City, UT; Beijing, China; Po Valley, Italy) (Bressi et al. 2016; Franchin et al. 2018; Lu et al. 2019; Young et al. 2016). Analysis of the OA composition indicates oxygenated OA, likely synonymous with SOA, often comprises a large fraction of the OA, along with OA derived from major primary sources, including biomass combustion, vehicles, and cooking (Bressi et al. 2016; Crippa et al. 2013; Lu et al. 2019; Young et al. 2016; Zhang et al. 2007;
Paglione et al. 2020). Yet, substantial challenges remain regarding the ability of models to quantitatively predict ambient aerosol concentrations in these wintertime environments, especially of the secondary components (Fountoukis et al. 2016; Kleeman 2019; Schroder et al. 2018). These challenges arise from the incomplete understanding of the physical, chemical, and thermodynamic processes that govern aerosol formation and partitioning, and how these are impacted by wintertime meteorological conditions. In addition, differences in the chemical environment and processes occurring within the nocturnal residual layer versus the surface layers are of particular importance in wintertime owing to shallower nocturnal boundary layers coupled with less available sunlight compared to other seasons (Baasandorj et al. 2017; Prabhakar et al. 2017; Pusede et al. 2016; Wang et al. 2018).

Dry deposition may influence ozone and particle loss rates, but particle deposition velocities in the key submicron particle range are generally small and thus dry deposition will likely only play a considerable role when the nocturnal boundary layer is particularly shallow (Emerson et al. 2020) and the aerodynamic resistance is low (i.e., vigorous turbulent mixing.) Dry deposition of gases such as HNO$_3$ or oxygenated VOCs can drive evaporation of particulate nitrate and organic aerosol and thus indirectly contribute to loss of PM$_{2.5}$ (Pusede et al. 2016; Knote et al. 2015). For HNO$_3$, such indirect loss processes are most important in warmer, ammonia-deficient conditions, when HNO$_3$ comprises a large fraction of the total nitrate, and when the boundary layer is particularly shallow (Prabhakar et al. 2017) and loss of OA will be similarly enhanced in warmer conditions. Direct measurements of dry deposition of such gases can help constrain understanding of the importance of these loss processes (e.g., Nguyen et al. 2015), and further examined through the use of process-level models (e.g., Prabhakar et al. 2017). Ozone dry deposition is highly variable based on the surface characteristics and, if not measured directly, it can presumably be parameterized. However, particle loss via scavenging by fog may be important in valley regions when the fog penetrates to the surface (Gilardoni et al. 2014); chemical processing within fog may also play a role in transforming the ambient aerosol composition (Gilardoni et al. 2016). Improved treatment of scavenging by wet deposition that accounts for variability in in-cloud water has been shown to improve model skill at predicting nitric acid and particle phase nitrate under wintertime conditions (Luo et al. 2019). Strong, multiday pollution events are often ended by a change in synoptic conditions that brings increased ventilation and, potentially, precipitation (Largeron and Staquet 2016).

One consideration is cold conditions with low absolute humidity but variable relative humidity characteristic of many wintertime valley environments. The limited availability of thermodynamic data for electrolyte solutions—especially mixtures containing organics—at low temperatures means that predictions of gas–particle partitioning and aerosol pH using existing thermodynamic models are extrapolations and may not be accurate. Direct measurements of the gas–particle distribution of inorganic and some organic species (e.g., organic acids) can serve to evaluate model predictions of equilibrium phase partitioning and aerosol pH (Murphy et al. 2017; Guo et al. 2016; Nah et al. 2018). However, the low temperatures may give rise to long equilibration time scales, making assumptions of thermodynamic equilibrium on typical model time scales questionable. Comparison between sufficiently detailed observations and model predictions will allow for investigation of sensitivities to temperature and RH (Evanoski-Cole et al. 2017). The viscosity of particles varies over many orders of magnitude, dependent on the particle composition, ambient temperature, and relative humidity (Renbaum-Wolff et al. 2013). Consequently, particles exhibit material properties, e.g., diffusivities, ranging from liquid-like to solid (Koop et al. 2011), which affects gas–particle partitioning and equilibration time scales (Shiraiwa and Seinfeld 2012; Shiraiwa et al. 2013a), heterogeneous reactions (Gaston et al. 2014; Li et al. 2018; Liu et al. 2018), and particle growth dynamics (Shiraiwa et al. 2013b; Zaveri et al. 2014; Zaveri et al. 2018). Additionally,
components within particles can undergo phase-phase separation into an organic-rich and inorganic/aqueous-rich phase, leading to unexpected internal morphologies within individual particles (Song et al. 2012; You et al. 2012). It is unclear how such phenomena impact rates of mass transfer between particles and the rates of aerosol-forming chemical processes, such as N₂O₅ reactive uptake, that occur on and within particles in the cold wintertime conditions.

Observations of N₂O₅ reactive uptake coefficients exhibit complex dependencies on particle OA mass fraction (Bertram et al. 2009), along with the OA chemical composition (e.g., the O:C atomic ratio) and potential for liquid–liquid phase separation (Gaston et al. 2014). The N₂O₅ reactive uptake coefficient may increase with decreasing temperature, perhaps owing to increased solubility of N₂O₅ (Abbatt et al. 2012; Hallquist et al. 2003; Schweitzer et al. 1998; Wagner et al. 2013), but requires further characterization for particle compositions reflective of wintertime valleys.

Particle composition and abundance vary with size, with especially large differences between submicron and supermicron particles. Compositional differences consequently engender size-varying differences in aerosol water content—critical to N₂O₅ uptake and other heterogeneous processes. The importance of considering and characterizing refractory and supermicron particle variations varies between regions. For example, supermicron particle contributions in the wintertime SJV are often small (Parworth et al. 2017), while in the colder SLV it can be substantial as a result of dust, road salt and lake salt (Hrdina et al. 2021). Compared to submicron particles, field observations of supermicron particle concentration, size, and composition are often limited, challenging model–measurement comparisons. Particle composition affects both N₂O₅ uptake and resulting products (HNO₃ versus ClNO₂), and often there are differences between laboratory parameterizations and field observations of ClNO₂ yields (McDuffie et al. 2018). McNamara et al. (2020) recently showed that ClNO₂ production, driven by the surface reaction of N₂O₅ (Gaston and Thornton 2016), is explained by surface area-weighted single-particle composition and developed a new parameterization that requires testing in other environments. Notably, the SLV has two unique chloride sources—playa dust and road salt—that likely contribute to ClNO₂ (Mitroo et al. 2019; McNamara et al. 2020). Measurements completely characterizing the time and size-varying particle composition in wintertime valley environments are needed to accurately simulate nighttime production of HNO₃ via N₂O₅ uptake.

Gas–particle partitioning of HNO₃ and other acids depends on size-varying particle and gas composition. In submicron particles, acid–anion phase partitioning is strongly controlled by NH₃ availability, alongside temperature, RH, and aerosol pH, the latter of which depends on the identity and abundance of other aerosol constituents (e.g., particulate sulfate or organic aerosol) (Murphy et al. 2017). In contrast, HNO₃ can react effectively irreversibly with supermicron dust or sea salt particles, displacing CO₂ (from CaCO₃) (Usher et al. 2003) or HCl (from NaCl) (Gard et al. 1998). As dry deposition is much faster for supermicron particles than accumulation mode, uptake to supermicron particles can thus suppress submicron nitrate formation. An observationally constrained understanding of the dynamic links between HNO₃, NH₃, and submicron versus supermicron reactive uptake and partitioning is needed.

Formation of SOA in valley environments during winter has received relatively little consideration compared to summer, although SOA can comprise a substantial fraction of aerosol dependent upon the conditions. Some observations indicate that the [SOA]/[NO₃⁻] ratio, or the [OA]/[NO₃⁻] ratio when the former is not available, decreases as the total aerosol loading increases, although the absolute SOA concentration tends to increase with aerosol loading (Bressi et al. 2016; Franchin et al. 2018; Lu et al. 2019; Young et al. 2016). This suggests some link between SOA and nitrate formation, potentially related to the dependence of aerosol liquid water content on particulate nitrate; higher aerosol water content can lead to enhanced uptake and reaction of water-soluble organic compounds—including at night.
Aqueous processing, especially of VOCs from residential wood combustion, lead to production of high molecular weight (Qin and Prather 2006) and absorbing organic aerosol (i.e., brown carbon, BrC) (Laskin et al. 2015; Gilardoni et al. 2016), which can influence the local and regional radiative budget (Mohr et al. 2013). Understanding of SOA and BrC formation in wintertime environments is lacking, both within and outside of valleys, with large model–measurement differences in some environments and a strong sensitivity to the particular SOA parameterization used (Chrit et al. 2018; Fountoukis et al. 2016; Meroni et al. 2017; Schroder et al. 2018; Shah et al. 2019). It may be that current parameterizations, developed almost entirely using room-temperature experimental data, are not properly capturing T-dependent changes in both gas- and particle-phase chemical pathways and in compound volatility (Bianchi et al. 2019; Stolzenburg et al. 2018; Ye et al. 2019). Winter SOA formation via reaction of NO$_3$ radicals with VOCs, especially from sources such as VCPs and residential wood combustion in poorly mixed nighttime boundary layer structures, has received relatively little attention (Mohr et al. 2013; Yuan et al. 2016), even though NO$_3$ chemistry has been proposed as a dominant SOA source during summer in the SJV (Rollins et al. 2012).

Overall, the identity and contribution of major submicron aerosol constituents in wintertime valleys is well known, with less known about the supermicron aerosol. Still, gaps in process-level understanding of the formation pathways and diurnally varying partitioning confront the development of clear control strategies and major improvements in wintertime air quality in such regions. Further, particulate matter loss processes, including wet and dry deposition, are poorly constrained by observations, as are the loss processes of PM precursors. Targeted, comprehensive measurements that can be compared with models are necessary to close these gaps.

**Greenhouse gas emissions**

The proposed field study will yield unique insights into wintertime GHG emissions and how they relate to air pollutant and associated precursor emissions in western U.S. basins. Wintertime emissions likely differ from other seasons owing to the presence of heating needs (Gurney et al. 2012; Mitchell et al. 2018a,b). Specifically, this field campaign will link GHG to emission sectors to understand how shifts in these emissions are associated with changes in short-lived pollutants and use GHG emissions as transport tracers to elucidate meteorological processes.

Three GHG emissions sectors will be carefully considered: transportation, agriculture, and oil/gas and energy distribution. These are most relevant in the western United States. The on-road transportation sector is one of the largest emitters in urban settings, often accounting for nearly 50% of CO$_2$ and other pollutants, but is generally the least well-constrained sector within carbon emission inventories, with large uncertainties (Gately and Hutrya 2017; Mendoza et al. 2013). Agriculture is an important source of GHG and air pollutants in the western United States, in particular methane (CH$_4$) and nitrous oxide (N$_2$O), respectively comprising nearly 40% and 80% of total emissions (U.S. Environmental Protection Agency 2019). Several studies have now shown that inventories underestimate methane emissions from oil and gas production (Alvarez et al. 2018; Robertson et al. 2017; Lyon et al. 2015). To constrain the contribution of all of these (and other) source types to degradation in air quality, will require monitoring a variety of trace gases.

Changes in vehicle fleets and improvements in technology have reduced mobile-sector emissions of both CO$_2$ (Gately et al. 2015) and air pollutants (McDonald et al. 2018). For example, NOx and carbon monoxide (CO) emissions in cities have fallen (Parrish et al. 2011; Hassler et al. 2016), although the decline has slowed recently (Jiang et al. 2018). Also, changes in the U.S. fuel mix (U.S. Energy Information Administration 2019), may affect emissions.
of both GHGs and pollutants. These trends point to changes in the urban atmospheric composition that might manifest themselves in observed concentrations. Work in Salt Lake City (Lin et al. 2018), with the longest running urban CO$_2$ network, indicates that CO$_2$ enhancements have remained flat in the urban core. This is despite population growth, implying energy efficiency improvements, and increased CO$_2$ enhancements in the suburbs due to new anthropogenic emissions occurring where previously there were none (Mitchell et al. 2018a). These long-running datasets in Salt Lake City are complemented by a novel platform on the public transit light-rail trains that measure GHGs and air pollutants as the trains traverse the city (Mitchell et al. 2018b; Mendoza et al. 2019). Concentration ratios from this platform have been used to fingerprint NO$_x$ emission sources (Mitchell et al. 2018b) and characterize how fine-scale meteorological phenomena, such as canyon flows and interbasin transport, affect air pollution distributions (Baasandorj et al. 2018). The proposed field campaign will utilize GHG and pollutant datasets from aircraft, mobile laboratories and ground sites.

The chemical inertness of GHGs such as CO$_2$ and CH$_4$ allows for their use as passive tracers of atmospheric transport (Pataki et al. 2005). This is particularly true in the wintertime, when photosynthetic uptake of CO$_2$ is minimal (Pataki et al. 2003; Strong et al. 2011). Measurements of CO$_2$ in the SLV have revealed strong relationships between levels of CO$_2$ and atmospheric stability, with the presence of PCAPs associated with significant enhancements in CO$_2$ concentrations (Pataki et al. 2005; Bares et al. 2018; Lin et al. 2018). Therefore, if GHG emissions are constrained, levels of GHG can potentially be used to quantify the effects of atmospheric transport processes and separate the contributions of such transport effects from atmospheric chemistry.

**Campaign design**

The recent 2017 Utah Winter Fine Particulate Study (UWFPS) in SLV was an aircraft and ground-based campaign utilizing the NOAA Twin Otter, a medium sized research aircraft (Baasandorj et al. 2018). The study demonstrated the capability to conduct in situ aircraft measurements in shallow winter boundary layers within complex terrain during periodic episodes of low visibility. A clear lesson learned from the UWFPS study was that the complexity of the science governing wintertime air quality is better addressed through a broader suite of instrumentation and focal area than could be accommodated by the payload and range of a Twin Otter. A larger aircraft (e.g., NSF C-130, NOAA P-3) would allow for a more extensive payload providing detailed in situ and remote sensing instrumentation, as well as the ability to sample multiple basins within a domain that could encompass most or all of the western United States (see Fig. 1).

The workshop considered the optimal combination of both ground-based and airborne measurements and platforms required to address the science questions identified. A recurring theme was the importance of combining collocated, comprehensive meteorological and chemical measurements. The workshop attendees stressed the need for large aircraft to carry the payload required to characterize both the organic and inorganic gases and particle composition and understand their relationships. Participants also expressed the need for airborne and surface measurements of coupled meteorological and chemical processes to bring insight to not only the formation and dissipation of the PCAPs that allow for the buildup of these extreme air quality events, but especially the vertical and horizontal transport and mixing processes modulating the concentrations of pollutants and precursors and constraining chemical processes. Based on the climatology of PCAPs in these regions (Whiteman et al. 2014), the optimum period for an airborne field campaign is from approximately in mid-December early through mid-February. Measurements may not be continuous during this period and would depend on logistical considerations, such as total aircraft hour allocation to the project.
Key measurements that will be needed on board the aircraft includes standard meteorological parameters along with fast-response measurements of VOCs, most likely using chemical ionization mass spectrometry, the various NO\textsubscript{x} and NO\textsubscript{y} species (e.g., NO, NO\textsubscript{2}, HNO\textsubscript{3}, N\textsubscript{2}O\textsubscript{5}, HONO, NO\textsubscript{3}) and other important trace gases (e.g., O\textsubscript{3}, SO\textsubscript{2}, HCHO, CO) and greenhouse gases (CO\textsubscript{2}, CH\textsubscript{4}, N\textsubscript{2}O), and submicron and supermicron aerosol composition and size distributions. A suite of ground-based measurements (in situ and remote sensing) will complement aircraft data to characterize the coupled meteorological–chemical system in three dimensions during the field campaign. These include wind and aerosol lidars, radio acoustic sounders, rawinsonde, ceilometers, and long-path Differential Optical Absorption Spectroscopy (DOAS) to provide continuous observations of vertical profiles (e.g., temperature, wind speed, and trace gases such as NO\textsubscript{2}, SO\textsubscript{2}, O\textsubscript{3}, HONO, HCHO, and NO\textsubscript{3}). Specifically, a western U.S. winter air quality study should include comprehensive measurements of major radical sources, including CH\textsubscript{2}O and other aldehydes, HONO, CINO\textsubscript{3}, and O\textsubscript{3}, together with measurements of actinic flux to define photolysis rates and radical sources. Measurements of major radical species, including OH, HO\textsubscript{2} and preferably RO\textsubscript{2}, together with nighttime measurements of NO\textsubscript{3} radicals, must also be available to constrain diel radical cycles and concentrations and to compare with the process-level predictions. Speciated measurements of these radicals and their precursors are most easily carried out from ground sites. However, measuring a subset of these compounds from a research aircraft allows for extension of the conclusions regionally and as a function of altitude. Comparison of continuous measurements at a single location to vertical and horizontal distributions of radicals and/or radical precursors provides a more complete view of the radical budget.

Complete characterization requires fully instrumented aircraft measuring O\textsubscript{x,total} and its partitioning between nitrogen and oxygen species (O\textsubscript{3}, NO\textsubscript{2}, NO\textsubscript{3}, HNO\textsubscript{3}, N\textsubscript{2}O\textsubscript{5}, CINO\textsubscript{3}, etc.) within and above the boundary layer during daytime and nighttime. In general, total budgets have not been fully quantified in valleys in the western United States. However, it is clear that odd-oxygen species play a substantial role in the buildup of wintertime pollutants. Owing to the significance and uncertainty in HONO sources, nitrous acid measurement via multiple techniques and from ground and aircraft platforms will be an essential component of a future western U.S. winter air quality study.

High temporal and spatial resolution measurements of NO\textsubscript{x}, VOCs from multiple sources (transportation, VCPS, cooking, wood combustion, biogenic, industrial, agriculture), NH\textsubscript{3} and amines are required. Multiple tracers will enable quantitative attribution of GHG emissions to different sectors through their enhancement ratios. These include enhancement ratios of CO\textsubscript{2} and CH\textsubscript{4} relative to tracers such as CO and NO\textsubscript{2}. For instance, using the CH\textsubscript{4}:CO ratio measured from aircraft, a recent study revealed that CH\textsubscript{4} emissions from urban areas in the East Coast corridor were drastically underestimated by inventories (Plant et al. 2019). Similarly, observed VOC:GHG correlations would allow attribution to oil/gas emissions (Pétron et al. 2012). In addition, novel tracers like atmospheric O\textsubscript{2} will be considered for source attribution as part of the planned field campaign.

Summary and conclusions
The interactions between complex wintertime chemistry and numerous meteorological processes regulating the occurrence and rates of chemical processes in PCAPs has not yet been observed in a detail needed to provide satisfactory understanding of the evolution of these complex pollution episodes. This paper provides an outline of the comprehensive chemical and meteorological datasets needed from airborne platforms with extensive range, coupled to a variety of surface-based measurements with high vertical resolution at numerous representative sites to provide enhance understanding of these pollution episodes. The workshop has outlined the design and requirements for a field campaign.
investigating wintertime basins to provide the breadth of observations necessary to characterize the meteorological–chemical linkages to validate complex processes within coupled atmosphere–chemistry models.

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