Organic Aerosol Sources and Chemistry: Insights from Development and Application of In-Situ Thermal Desorption Gas Chromatograph for Semi-Volatile Organic Compounds (SV-TAG)

by

Yunliang Zhao

A dissertation submitted in partial satisfaction of the requirements for the degree of Doctor of Philosophy in Environmental Science, Policy, and Management in the Graduate Division of the University of California, Berkeley

Committee in charge:

Professor Allen H. Goldstein, Chair
Professor Dennis D. Baldocchi
Professor Ronald C. Cohen

Fall 2012
Organic Aerosol Sources and Chemistry: Insights from Development and Application of In-Situ Thermal Desorption Gas Chromatograph for Semi-Volatile Organic Compounds (SV-TAG)

©2012

by Yunliang Zhao
Abstract

Organic Aerosol Sources and Chemistry: Insights from Development and Application of In-Situ Thermal Desorption Gas Chromatograph for Semi-Volatile Organic Compounds (SV-TAG)

by

Yunliang Zhao

Doctor of Philosophy in Environmental Science, Policy, and Management

University of California, Berkeley

Professor Allen H. Goldstein, Chair

Understanding organic aerosol (OA) sources and secondary OA (SOA) formation is crucial to elucidate their human health and climate change effects, but has been limited by lack of instrumentation capable of in-situ measurements of organic speciation in the atmosphere across the vapor pressure range of semi-volatile organic compounds (SVOCs) and OA. This dissertation describes 1) the development of a novel instrument based on a thermal desorption aerosol gas chromatograph (TAG), called semi-volatile TAG (SV-TAG) which enables quantitative measurements of specific chemical tracers in SVOCs and OA and 2) application of this new instrument to investigate the various source contributions to OA and SOA formation.

The development of the SV-TAG was initiated by employing a denuder difference method to improve the capability of the TAG for quantitative gas/particle separation. Using this technique, hourly time resolution in-situ measurements of organic species were made and then used to investigate the pathways of gas-to-particle partitioning for oxygenated compounds and particle-phase organics were used for source apportionment calculations. The measurements of gas/particle partitioning of phthalic acid, pinonaldehyde and 6, 10, 14-trimethyl-2-pentadecanone were explored to elucidate the pathways of gas-to-particle partitioning whereby SOA was formed. The observations show that multiple pathways of gas-to-particle partitioning contribute to formation of SOA in the atmosphere and the dominance of different pathways are compound-dependent. Absorption into particles is shown to be the dominant pathway for 6, 10, 14-trimethyl-2-pentadecanone to contribute to SOA in Bakersfield, CA. The major pathway to form particle-phase phthalic acid is likely attributed to formation of condensable salts through reactions between phthalic acid and gas-phase ammonia. The observations of pinonaldehyde in particles while inorganic acids in particles were fully neutralized suggest that the occurrence of reactive uptake of pinonaldehyde onto particles does not require the presence of inorganic acids. The relationship between particle-phase pinonaldehyde and RH suggests that aerosol water content plays a significant role in the formation of particle-phase pinonaldehyde. To investigate the contributions of various sources to OA in Bakersfield, CA, positive matrix factorization (PMF) analysis was performed on a subset of the measured particle-phase organic compounds. Six OA source factors were identified, including one representing primary organic aerosol (POA), four different types of secondary organic aerosol (SOA) representing local, regional, and
nighttime production, and one representing a complex mixture of additional OA sources that were not further resolvable. POA accounted for 15% of OA on average with a significant contribution from local vehicles. SOA was the dominant contributor to OA, accounting for on average 72% of OA. The rest of OA was unresolved as a mixture of OA sources. Both local and regional SOA had a significant contribution to OA during the day but regional SOA was the largest contributor to OA during the afternoon. SOA formed from the oxidation of biogenic SOA precursors substantially contributed to OA at night. The absorption of organic compounds into particles is suggested to be the major pathway to form SOA, although other pathways also played significant roles.

To achieve quantitative collection of SVOCs following improved gas/particle separation, a new collection and thermal desorption system was developed with the key component being a passivated metal fiber filter collector. This final configuration of the SV-TAG enabled in-situ quantitative measurements of speciated SVOCs with vapor pressures lower than n-tetradecane (C\textsubscript{14}). The capability for measurements of gas/particle partitioning was demonstrated by measurements of \textit{n}-alkanes in both gas and particle phases. Organic tracers in both gas and particle phases can be quantified. Percentages of speciated organic compounds in total measured organics can be estimated. For example, ~7% and less than 1% of total measured organics in the same retention range of \textit{n}-alkanes (C\textsubscript{14}-C\textsubscript{20}) in the atmosphere in Berkeley, CA were accounted for by the sum of measured \textit{n}-alkanes (C\textsubscript{14}-C\textsubscript{20}) and the sum of \textit{n}-alkylcyclohexanes (C\textsubscript{14}-C\textsubscript{20}).

The SV-TAG has been demonstrated to enable investigation of the pathways of gas-to-particle partitioning and source apportionment of OA with hourly time resolution. The SV-TAG is also capable of quantitative measurements of speciated SVOCs, defining their gas/particle partitioning in-situ for the first time, and providing observational constraints on the abundance of SVOCs with which to investigate their primary emissions, chemical transformation, and fate.
# Table of Contents

Table of Contents.................................................................................................................. i  
List of Figures......................................................................................................................... iii  
List of Tables............................................................................................................................. v  
Acknowledgements.................................................................................................................... vi

**Chapter 1  Introduction** ........................................................................................................ 1
  1.1 Identification of OA sources............................................................................................ 1
  1.2 SOA formation.................................................................................................................. 2
    1.2.1 Gas-to-particle partitioning...................................................................................... 2
    1.2.2 IVOCs and SVOCs................................................................................................ 3
  1.3 Instrumentation................................................................................................................. 4
  1.4 Focus of PhD dissertation............................................................................................... 4
  1.5 References....................................................................................................................... 5

**Chapter 2  Insights into SOA formation mechanisms from measured gas/particle partitioning of specific organic tracer compounds** .................................................................................. 9
  2.1 Abstract............................................................................................................................ 9
  2.2 Introduction...................................................................................................................... 9
  2.3 Methods.......................................................................................................................... 10
    2.3.1 Sampling and analysis.......................................................................................... 11
    2.3.2 Particle-phase fraction calculations.................................................................... 12
  2.4 Results and discussions................................................................................................... 13
    2.4.1 Pinonaldehyde...................................................................................................... 13
    2.4.2 Phthalic acid......................................................................................................... 15
    2.4.3 6,10,14-trimethyl-2-pentadecanone.................................................................. 16
  2.5 Conclusions and implications......................................................................................... 16
  2.6 References....................................................................................................................... 17
  2.7 Tables and Figures.......................................................................................................... 21

**Chapter 3  Sources of organic aerosol investigated using organic compounds as tracers measured during CalNex Bakersfield** ...................................................................................... 26
  3.1 Abstract............................................................................................................................ 26
  3.2 Introduction...................................................................................................................... 26
  3.3 Methods.......................................................................................................................... 28
    3.3.1 Sampling and chemical analysis.......................................................................... 28
    3.3.2 PMF procedures.................................................................................................... 29
  3.4 PMF results...................................................................................................................... 30
    3.4.1 Factor 1: local POA............................................................................................... 30
    3.4.2 Factor 2: A mixture of OA sources...................................................................... 31
    3.4.3 Factor 3: SOA1..................................................................................................... 31
    3.4.4 Factor 4: SOA2..................................................................................................... 31
    3.4.5 Factor 5: SOA3..................................................................................................... 32
    3.4.6 Factor 6: nighttime SOA (SOA4)...................................................................... 32
  3.5 Reconstructed OA............................................................................................................ 32
  3.6 Source contributions to OA mass.................................................................................... 33
  3.7 Formation pathways of SOA.......................................................................................... 34
  3.8 Conclusions and atmospheric implications................................................................... 35
<table>
<thead>
<tr>
<th>Section</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.9 References</td>
<td>35</td>
</tr>
<tr>
<td>3.10 Tables and figures</td>
<td>39</td>
</tr>
<tr>
<td><strong>Chapter 4</strong> Development of an in situ thermal desorption gas chromatography instrument for quantifying atmospheric semi-volatile organic compounds</td>
<td>44</td>
</tr>
<tr>
<td>4.1 Abstract</td>
<td>44</td>
</tr>
<tr>
<td>4.2 Introduction</td>
<td>44</td>
</tr>
<tr>
<td>4.3 Methods</td>
<td>46</td>
</tr>
<tr>
<td>4.3.1 The SV-TAG system</td>
<td>46</td>
</tr>
<tr>
<td>4.3.2 SV-TAG operation</td>
<td>47</td>
</tr>
<tr>
<td>4.3.3 System evaluation</td>
<td>48</td>
</tr>
<tr>
<td>4.3.4 Ambient sampling</td>
<td>50</td>
</tr>
<tr>
<td>4.4 Evaluation results</td>
<td>50</td>
</tr>
<tr>
<td>4.4.1 Thermal desorption and transfer efficiency</td>
<td>50</td>
</tr>
<tr>
<td>4.4.2 SVOC collection efficiency by the F-CTD</td>
<td>51</td>
</tr>
<tr>
<td>4.4.3 Particle collection efficiency by the F-CTD</td>
<td>51</td>
</tr>
<tr>
<td>4.4.4 Denuder vapor collection and particle penetration</td>
<td>52</td>
</tr>
<tr>
<td>4.4.5 Measurements of ambient air</td>
<td>52</td>
</tr>
<tr>
<td>4.5 Summary and atmospheric implications</td>
<td>53</td>
</tr>
<tr>
<td>4.6 References</td>
<td>54</td>
</tr>
<tr>
<td>4.7 Tables and figures</td>
<td>58</td>
</tr>
<tr>
<td><strong>Chapter 5</strong> Conclusions</td>
<td>64</td>
</tr>
<tr>
<td>5.1 Summary</td>
<td>64</td>
</tr>
<tr>
<td>5.1.1 Gas-to-particle partitioning (SOA formation)</td>
<td>64</td>
</tr>
<tr>
<td>5.1.2 Major source contributions to OA</td>
<td>65</td>
</tr>
<tr>
<td>5.1.3 SV-TAG and ambient measurements</td>
<td>65</td>
</tr>
<tr>
<td>5.2 Future work</td>
<td>66</td>
</tr>
<tr>
<td>5.2.1 Measurements of gas/particle partitioning</td>
<td>67</td>
</tr>
<tr>
<td>5.2.2 Measurements of SVOCs</td>
<td>67</td>
</tr>
<tr>
<td>5.2.3 SOA formation and transformation</td>
<td>67</td>
</tr>
<tr>
<td>5.2.4 Instrumentation</td>
<td>67</td>
</tr>
<tr>
<td><strong>Appendix A</strong> Supplemental information for Chapter 2</td>
<td>69</td>
</tr>
<tr>
<td><strong>Appendix B</strong> Supplemental information for Chapter 3</td>
<td>71</td>
</tr>
<tr>
<td><strong>Appendix C</strong> Supplemental information for Chapter 4</td>
<td>75</td>
</tr>
</tbody>
</table>
List of Figures

Chapter 2 - Insights into SOA formation mechanisms from measured gas/particle partitioning of specific organic tracer compounds

Figure 2.1  O/C ratios and subcooled vapor pressures of measured organic species 21
Figure 2.2  Average measured and predicted fractions for oxygenated SVOCs and their corresponding reference compounds 22
Figure 2.3  Measured fraction of pinonaldehyde in particles and the cation-to-anion ratio as a function of RH 23
Figure 2.4  Timelines of the cation-to-anion ratio and the fraction of pinonaldehyde in particles throughout the field campaign 24
Figure 2.5  Correlation between the fraction of phthalic acid in particles and the concentration of gas-phase ammonia 25

Chapter 3 - Sources of organic aerosol investigated using organic compounds as tracers measured during CalNex Bakersfield

Figure 3.1  PMF factor profiles 39
Figure 3.2  Diurnal averages for each PMF factor, the wind direction and the ratio of 1,3,5-TMB to toluene 40
Figure 3.3  Wind rose plots to emphasize the major contributing source directions 41
Figure 3.4  Mean diurnal mass fraction contribution of each factor to total OA 42
Figure 3.5  Diurnal averages for SOA factors from AMS PMF and TAG PMF solutions 43

Chapter 4 - Development of an in situ thermal desorption gas chromatography instrument for quantifying atmospheric semi-volatile organic compounds

Figure 4.1  Schematic of the SV-TAG system 58
Figure 4.2  Recoveries of representative compounds 59
Figure 4.3  Recoveries of n-alkanes ≤ C26 measured with different FTs 60
Figure 4.4  Overall gas collection efficiency of the F-CTD 61
Figure 4.5  Stability of the SV-TAG collection and detection system 62
Figure 4.6  Ambient measurements of gas- and particle-phase organics 63

Appendix A: Supplemental Information for Chapter 2

Figure A1  The stability of gas collection efficiency of the denuder 69
Figure A2  Particle losses inside the denuder 70

Appendix B: Supplemental Information for Chapter 3

Figure B1  (a) The relative difference between the reconstructed and measured OA concentration; (b) the frequency of the different atmospheric OA concentrations occurred at night 72
Figure B2  Factor profiles extracted by PMF analysis from the same group of compounds with or without inclusion of gas-phase contributions to measured SVOCs 73
Figure B3  Correlation of PMF results from the dataset with or without inclusion of
List of Figures

Appendix C: Supplemental Information for Chapter 4

| Figure C1 | Schematic diagrams of the F-CTD and its housing assembly | 78 |
| Figure C2 | Schematic diagram of the FT assembly | 79 |
| Figure C3 | Operating conditions for the evaluation of gas collection efficiency of the F-CTD using thermal desorption method | 80 |
| Figure C4 | Operating conditions for the evaluation of the gas collection efficiency of the F-CTD using evaporation transfer | 81 |
| Figure C5 | Operation conditions of the SV-TAG system during ambient measurements | 82 |
| Figure C6 | Optimization of the recovery of less volatile organic compounds | 83 |
| Figure C7 | Particle collection efficiency of the F-CTD for the mobility diameter from ~30 nm to ~350 nm | 84 |
| Figure C8 | Particle collection efficiency of the F-CTD for the mobility diameter from ~60 nm to ~1000 nm | 85 |
| Figure C9 | Effect of organic loading on the recovery of individual compounds | 86 |
List of Tables

Appendix B: Supplemental Information for Chapter 4

Table B1 - Organic compounds used to evaluate the overall SVOC CE of the F-CTD 77
Acknowledgements

This dissertation is a culmination of research performed over 5 years and could not have been completed without the help and support from my advisor, colleagues, friends and family. I’d like to thank my advisor, Allen Goldstein, who is always supportive and encouraging by sharing his insights, suggestions and excitements for my research. I’m grateful for many challenges, opportunities that I grow to under his guidance. He is a great advisor to work with. I’d like to thank my colleagues in the Goldstein group, Dave, Gabriel, Arthur, Drew, Rachel and Jeong-Hoo, for their help in both laboratory and field measurements. I’d also like to thank other group members for their helpful discussions and making the Goldstein group a supportive environment to do science.

I’d like to thank Nathan Kreisberg and Susanne Hering at Aerosol Dynamic Inc.. Without their help, I could not complete the development of the novel instrument, SV-TAG. I have gone to Nathan numerous times to ask for advice on instrumentation and instrument designs. I’d like to thank Lara Gundel at LNBL as well, from whom I have learned a lot about denuders when I first started the development of the SV-ATG. I also thank University of California Extension Staff and Kern County Staff for logistical support during the Bakersfield CalNex study. I have collected most part of data for my dissertation during the CalNex Campaign. I also thank my other collaborators, including Lynn Russell, Shang Liu and Douglas Day at UC San Diego and Jennifer Murphy, Trevor Vandenboer and Milos Markovic at University of Toronto.

I’d also like to thank my dissertation committee, Professors Allen Goldstein, Dennis Baldocchi and Ronald Cohen, for their thorough and critical readings of my dissertation and for their help to make research and dissertation better.

I am grateful to my parents, Fuying Zhao and Luansheng Wang, for their understanding, support and encouragement all these years. Most of all, I want to thank Ying (Jane) Hong for being such an important part of my life. I own so much of my success to her. Finally, I’d like to thank everyone who helped, supported and encouraged me along the way.
Chapter 1
Introduction

Organic aerosol (OA) constitutes 20–90% of the total fine aerosol mass in most regions (Kanakidou et al., 2005). OA can be categorized into either primary organic aerosol (POA) directly emitted from sources, such as food cooking and vehicle exhausts (e.g. Schauer et al., 1999a, b, 2002) or secondary organic aerosol (SOA) formed in the atmosphere from different precursors through chemical reactions (Odum et al., 1997; Jang et al., 2002; Robinson et al., 2007; Kroll and Seinfeld, 2008). SOA has been found to have different adverse human health effects than POA (Li et al., 2008) and plays a significant role in affecting climate change on both global and regional scales (Hoyle et al., 2009; Goldstein et al., 2009). Understanding OA sources and SOA formation is a critical step toward elucidating their roles in climate change and human health and developing effective control strategies for reducing air pollution.

1.1 Identification of OA sources

Quantification and characterization of OA can be made with a wide range of analytical techniques, such as gas chromatography-mass spectrometry (GC-MS), Fourier transform infrared spectroscopy, Aerodyne Aerosol Mass Spectrometry (AMS), and so on, with chemical resolution spanning from individual organic compounds to function groups to the carbon content and characterization completeness range from less than 20% to 100% (e.g. Turpin et al., 2000; Hallquist et al., 2009). None of these techniques is able to directly determine the amount of POA and SOA or the contributions of various sources to OA, but they provide complementary aerosol chemical composition information which can be analyzed to identify sources and their contributions to OA through statistical analyses and knowledge of source chemical signature and atmospheric transformation pathways.

Measurements of organic speciation made with GC-MS generally resolve less than 20% of the OA mass (Hallquist et al., 2009; Williams et al., 2006), but some of identified organic compounds can serve as molecular markers which are unique to specific OA sources, such as levoglucosan for wood combustion and phthalic acid for SOA (e.g. Schauer et al., 1996). Moreover, SOA tracers can provide information for understanding the chemical formation and transformation mechanisms leading to SOA (e.g. Kleindienst et al., 2007; Zhang et al., 2009; Williams et al., 2010). Chemical mass balance (CMB) and positive matrix factorization (PMF) models are two common methods that use speciated organic molecular marker to estimate source contributions to OA (e.g. Schauer et al., 1996; Schauer and Cass, 2000; Jaekels et al., 2007; Shrivastava et al., 2007; Williams et al., 2010). Both CMB and PMF models assume the measured chemical composition of OA in the atmosphere is the linear sum of contributions of a number of sources which have distinct source profiles (Schauer et al., 1996; Hopke, 2003; Reff et al., 2007; Shrivastava et al., 2007; Zhang et al., 2011).

The CMB model using organic molecular marker compounds directly apportions OA to specific sources, such as diesel vehicles and meat cooking, using their source profiles as model inputs (Schauer et al., 1996). As a result of the requirement of a priori knowledge of the source profile, the CMB model cannot apportion OA mass to sources with unknown or undefined
source profiles, such as SOA. Though the amount of SOA has been estimated by the difference of between apportioned OA and measured OA (Schauer et al., 1996; Zheng et al., 2002; Subramanian et al., 2007), SOA estimated through this approach is poorly constrained.

The PMF model apports organic species into distinct factors according to co-variation between them. The composition of organic species in each factor can be used to identify the presence of either a specific source or atmospheric process (Shrivastava et al., 2007; Williams et al., 2010). Since the PMF model solves for the source profiles, the sources with unknown or undefined source profiles can be determined by inclusion of all organic markers measured in the ambient samples. For example, four types of SOA were determined by performing the PMF analysis on a dataset of organic species measured hourly by a thermal desorption aerosol gas chromatograph (TAG) in a field study site in the Riverside, California (Williams et al., 2010). In the PMF analysis, the contributions of identified sources to OA can be estimated by either directly including OA mass in the PMF dataset as a compound or by using a multivariate fit of the temporal variability of factors onto measured OA (Reff et al., 2007; Shrivastava et al., 2007; Williams et al., 2010).

Bulk analyses of OA provided by optical-thermal elemental carbon (EC) and organic carbon (OC) analyzer or by AMS are particular useful for the split of POA and SOA (e.g. Yu et al., 2007; Zhang et al., 2007), but are not sufficient to conduct source apportionment of POA or SOA to specific sources or processes. Generally, EC tracer-based analysis assumes any organics leading to a higher OC/EC ratio than that in primary emissions must be SOA (Gray et al.1986; Turpin et al., 1991; Strader et al., 1999; Yu et al., 2007). PMF analysis of mass spectra produced by AMS is now routinely used to separate several components of OA, including hydrocarbon-like OA (HOA), low-volatility oxygenated OA (LV-OOA), semi-volatile oxygenated OA, with POA assumed to consist of hydrocarbon-like OA and SOA generally assumed to be represented by the sum of OOA (Ulbrich et al., 2009; Jimenez et al., 2009; Zhang et al., 2011); however, these components of OA cannot be traced to specific sources or processes without further information.

1.2 SOA formation

SOA accounts for the dominant fraction of OA in most regions and its contribution to OA can be over 80% in the afternoon during summer in urban areas (Lim and Turpin, 2002; Zhang et al., 2005; 2007; Williams et al., 2010). However, models, describing SOA formation through gas-phase oxidation of VOCs followed by absorptive partitioning of their low volatility oxidation products into particles, typically predict less SOA than it measured in polluted regions (e.g. Volkamer et al., 2006; Hodzic et al., 2010; Spracklen et al., 2011). These discrepancies between modeled and measured SOA can be in part attributed to uncharacterized gas-to-particle partitioning pathways in the atmosphere (e.g. Kroll et al., 2005; Kroll and Seinfeld 2008) or additional SOA precursors which are likely intermediate-volatility organic compounds (IVOCs) and semi-volatile organic compounds (SVOCs) (e.g. Goldstein and Galbally, 2007; Robinson et al., 2007; Hodzic et al., 2010).

1.2.1 Gas-to-particle partitioning
SOA formation is traditionally modeled by absorption into particles following formation of low-volatility oxidation products, but studies have shown that multiple other pathways of gas-to-particle partitioning are also important in the atmosphere (e.g. Jang et al., 2002; Kroll et al., 2005; Na et al., 2007; Iinuma et al., 2005). In one example, reactive uptake of carbonyls onto acidic particles has been suggested be an important additional pathway of SOA formation (e.g. Jang et al., 2002), even though the compounds involved in this pathway are present in the gas phase according to absorptive partitioning theory (Pankow, 1994). In another example, enhancement of SOA formation resulted from a-pinene ozone reactions has been observed in the presence of gas-phase ammonia and this enhancement was attributed to reactions between gas-phase ammonia and organic acids (Na et al., 2007).

Laboratory studies have shown that reactive uptake of carbonyls on acidic particles depends on specific compounds and experimental conditions. SOA formation from the reactive uptake of small carbonyls was observed to be significantly higher in the presence of an acidic catalyst (e.g. Jang et al., 2002; Iinuma et al., 2005; Kroll et al., 2005), but enhancement in SOA formation varied in different laboratory studies (e.g. Iinuma et al., 2005) and results reported by Kroll et al. (2005) showed that the reactive uptake of some of carbonyls onto particles was insignificant when concentrations of carbonyls used in the laboratory experiments were scaled to atmospheric levels.

Experimental conditions in laboratory studies often differ from the atmospheric conditions in many ways, including NOx level, particle acidity, preexisting aerosol mass, gas-phase organic mass and oxidation state (Kroll and Seinfeld, 2008). As a result, SOA formation pathways occurring in the atmosphere could be different from those characterized in laboratory studies. Therefore, ambient measurements of gas- and particle-phase organic species are necessary to identify the presence of different gas-to-particle partitioning pathways in the atmosphere and investigate the effects of atmospheric conditions, including particle acidity and RH, on gas-to-particle partitioning of different species. Ambient measurements are also necessary to examine the relative importance of different gas-to-particle partitioning pathways.

1.2.2 IVOCs and SVOCs

Recent modeling work has shown that models including estimated I/SVOCs emissions using the volatility basis set (VBS) approach (Robinson et al., 2007; Shrivastava et al., 2008; Grieshop et al., 2009) resulted in better agreement between measured and modeled SOA (Dzepina et al., 2009; Tsimpidi et al., 2010; Hodzic et al., 2010) than those which did not include estimated I/SVOCs (e.g., Volkamer et al., 2006; Hodzic et al., 2010). However, these newer models have not been able to reproduce both the SOA mass and the oxygen to carbon ratio of the bulk aerosol (Hodzic et al., 2010), demonstrating that there remains substantial deficiencies in understanding and modeling of I/SVOCs emissions and their atmospheric transformation into SOA.

These deficiencies of the VBS approach must be rooted in its simplification of the complex composition of I/SVOCs in the atmosphere and the dependence of SOA yields on molecular structure and other parameters. The VBS approach lumps all species with the same effective saturation concentration into a single volatility bin and assumes a constant mass increase for species in a given bin through each oxidation step (Robinson et al., 2007;
Shrivastava et al., 2007). Additionally, the amount of primary I/SVOCs included in these newer models is not measured in the atmosphere but instead is estimated by multiplying concentrations of primary organic aerosol (POA) by a scaling factor derived from dilution measurements of emissions from two primary sources: diesel exhaust and wood burning (Robinson et al., 2007; Grieshop et al., 2009). To better predict SOA production from the oxidation of I/SVOCs, it is critical to apply observational constraints to emissions, speciation, and gas-phase oxidation mechanisms of I/SVOCs.

Since oxygenated compounds in the vapor pressure range of IVOCs are known to partition into particles through non-absorptive gas-to-particle partitioning (e.g. Jang et al., 2002), there is no clear split between IVOCs and SVOCs in the atmosphere according to gas/particle partitioning. We therefore define "SVOCs" to include both IVOCs and SVOCs in this dissertation. There are other operational definitions of SVOCs based on the vapor pressure and air-sampling strategy (Turpin et al., 2000). However, SVOCs are generally in the vapor pressure range approximately equivalent to C_{12}-C_{32} n-alkanes (Turpin et al., 2000; Sihabut et al., 2005; Robinson et al., 2007; de Gouw et al., 2011).

1.3 Instrumentation

To improve the understanding of the primary source contributions to OA and SOA formation in the atmosphere, ambient measurements are needed for both gas- and particle-phase organic marker species which can be used to conduct PMF analyses, provide observational constraints on abundance, speciation and sources of SVOCs, and investigate their gas-to-particle partitioning pathways. Speciated measurements of organic chemicals in aerosols have historically involved integrated field sampling by filtration or impaction with subsequent solvent extraction in the laboratory for GC analysis. Generally the filter sampling has low time resolution (typically 24 hours) and thus obscure the variability in concentrations of organic species which are critical to understanding the dynamic variability of sources, chemistry, gas-to-particle partitioning and atmospheric transport processes. Additionally, these measurements have mainly focused on particle-phase organics and very few of them have been made to measure SVOCs.

The TAG is the first in-situ instrument capable of measuring speciated organic compounds in OA with hourly time resolution and capturing the diurnal trend of gas/particle partitioning in the atmosphere (Williams et al., 2006; Williams et al., 2010). However, the original TAG, equipped with an impactor collection cell, was not designed for collection of SVOCs. Additionally, a filter-based method, utilized by the original TAG to achieve separation of gas- and particle-phase organics, is subject to positive artifacts caused by adsorption of gas-phase organics on the filter as well as negative artifacts due to evaporation of particulate organics. As a result, while the original TAG was able to observe the trends in the concentrations of some speciated SVOCs and their gas/particle partitioning, but it could not do quantitatively or comprehensively.

1.4 Focus of PhD dissertation

The goal of my PhD dissertation is to develop a novel instrument, based on the concept of the TAG, that is capable of quantitative measurements of SVOCs and OA with hourly time
resolution and make ambient measurements with this instrument that would 1) improve our understanding of SOA formation pathways; 2) determine the contributions to OA from a variety of sources and 3) provide observational constraints on abundance, speciation and sources of SVOCs as SOA precursors. These three focuses are addressed in Chapter 2-4, respectively.

This dissertation consists of five chapters:

Chapter 1: Introduction
Chapter 2: Insights into SOA formation mechanisms from measured gas/particle partitioning of specific organic tracer compounds
Chapter 3: Sources of organic aerosol investigated using organic compounds as tracers measured during CalNex Bakersfield
Chapter 4: Development of an in situ thermal desorption gas chromatography instrument for quantifying atmospheric semi-volatile organic compounds
Chapter 5: Conclusions

The development of the SV-TAG was initiated by employing a denuder difference method to improve the separation of gas- and particle-phase organics of the original TAG followed by development of a new system to quantitatively collect and transfer SVOCs into GC/MS. Chapter 2 and 3 focus on the ambient measurements after a denuder was added into the sampling inlet of the original TAG. Ambient measurements were made as part of the CALifornia at the NEXus of Air Quality and Climate Change (CalNex) campaign in Bakersfield. In Chapter 2, the separation of gas- and particle-phase organics by a denuder was evaluated and time-resolved measurements of the gas/particle partitioning of SOA tracers were made to examine the presence of other SOA formation pathways in addition to absorptive partitioning in the ambient atmosphere. In Chapter 3, PMF analysis was performed on particle-phase organic species to investigate the source contributions to OA and provide insights into the importance of different SOA formation pathways in the atmosphere. Chapter 4 presents the development and extensive evaluations of a final configuration of SV-TAG, including the components developed in two development stages. Chapter 4 also reports the first SV-TAG ambient measurements of speciated SVOCs made in Berkeley, CA, which clearly demonstrates the capability to measure abundance, speciation and gas/particle partitioning of SVOCs.

1.5 References


Chapter 2
Insights into SOA formation mechanisms from measured gas/particle partitioning of specific organic tracer compounds

2.1 Abstract
Semi-volatile and intermediate-volatility organic compounds (S/IVOCs) in both gas and particle phases were measured using a Thermal desorption Aerosol Gas chromatograph (TAG) instrument during the CALifornia at the NEXus of Air Quality and Climate Change (CALNEX) campaign in Bakersfield, CA from May 31st to June 27th, 2010. The gas/particle partitioning of phthalic acid, pinonaldehyde and 6, 10, 14-trimethyl-2-pentadecanone is discussed in detail to explore secondary organic aerosol (SOA) formation mechanisms. Measured fractions in the particle phase ($f_{\text{part}}$) of 6, 10, 14-trimethyl-2-pentadecanone were similar to those expected from absorptive gas/particle partitioning theory, suggesting that its partitioning is dominated by absorption processes. However, $f_{\text{part}}$ of phthalic acid and pinonaldehyde were significantly higher than predicted. The formation of low-volatility products from reactions of phthalic acid with ammonia is proposed as one possible mechanism to explain the enhancement of particle-phase phthalic acid. This gas/particle partitioning pathway is expected to lead to high O/C ratios of SOA because it favors the partitioning of gas-phase organic acids into particles by forming condensable ammonium salts. The observations of particle-phase pinonaldehyde when inorganic acids were fully neutralized show that inorganic acids are not required for the occurrence of reactive uptake of pinonaldehyde on particles. The observed relationship between $f_{\text{part}}$ of pinonaldehyde and relative humidity (RH) suggests that the aerosol water content play a significant role in the formation of particle-phase pinonaldehyde. The identification of multiple pathways of oxygenated organics partitioning into particles in the atmosphere demonstrates that multiple pathways of gas/particle partitioning should be included in models to predict SOA and multiple tracers of SOA are needed to reconstruct SOA in source apportionment models.

2.2 Introduction
Secondary organic aerosol (SOA) accounts for the majority of organic aerosol (OA) on a global scale (Kanakidou et al., 2005; Goldstein and Galbally, 2007) and more than 80% in the afternoon during summer in urban areas (Williams et al., 2010a). However, predictions of SOA by traditional models based on laboratory measurements of SOA yields from traditional SOA precursors and absorptive partitioning theory have been shown to substantially underestimate the ambient SOA loadings in polluted regions (Heald et al. 2005; 2010; Volkamer et al., 2006; Spracklen et al., 2011). The discrepancies between measurements and models could in part be attributed to poor understanding of formation pathways of SOA in the ambient atmosphere.

Laboratory studies have shown that SOA formation pathways in addition to absorptive partitioning (Pankow, 1994), such as reactive uptake of gaseous species (Jang et al., 2002; Kroll et al., 2005) and gas-phase non-oxidative reactions (Na et al., 2007), could be important. However, these pathways remain poorly understood. For example, laboratory studies have shown that reactive uptake of oxygenated organic compounds on acidic particles can significantly increase SOA yields, but there is no agreement on the extent of enhancement in SOA yields.
parameterization of SOA formation in the ambient atmosphere and likely lead organic species pathways (gas/particle partitioning) allowing the upper limit of the overestimation to be estimated representing a and evaporation of collected organics from the filter. Additionally, laboratory studies of reactive uptake of oxygenated compounds have focused primarily on small carbonyl compounds and found that not all of them significantly contribute to SOA when their concentrations used in the laboratory studies are scaled to atmospheric levels (e.g., Jang et al., 2002; Kroll et al., 2005). As a result, the contribution of individual compounds to SOA cannot be generalized based solely on their functional groups. Ambient measurements are crucial to examine the importance of laboratory proposed SOA mechanisms.

Ambient measurements with an Aerodyne Quadrupole Aerosol Mass Spectrometer (Q-AMS) have been made to examine the effects of aerosol acidity on SOA formation and the results showed that no significant enhancement in SOA formation was observed during acidic periods identified based on the inorganic ion charge balance (Zhang et al., 2007). However, the importance of acid-catalyzed reactions in SOA formation might not be evident using the inorganic ion charge balance as the indicator of aerosol acidity because organic acids could also provide sufficient acidity for the occurrence of these reactions (Gao et al., 2004). Additionally, the variability in the amount of SOA formed through acid-catalyzed reactions could be obscured by SOA formed through other pathways if there is not an analytical method to distinguish them. In comparison with bulk OA analysis by AMS, time-resolved speciated measurements of gas- and particle-phase organic compounds are particular useful to determine concentrations of organic compounds involved in acid-catalyzed reactions and distinguish SOA products formed through acid-catalyzed reactions from other pathways. Moreover, these time-resolved, speciated measurements provide information to examine factors affecting SOA formation that have previously been investigated in laboratory studies, such as relative humidity (RH) and acidity, and discover new pathways of SOA formation in the atmosphere (Pankow, 1994; Jang et al., 2002; Tillmann et al., 2010).

Williams et al. (2010b) demonstrated that a Thermal desorption Aerosol Gas chromatography (TAG) instrument was able to capture the trend of gas/particle partitioning of individual species, wherein a filter-based sampling method was used to separate gases from particles. In this TAG, the fraction of organic species was overestimated because the its collection cell is designed for particles and is incapable of complete collection of vapors. Additionally, this filter-based method is subjective to sorption of gas-phase organics on the filter and evaporation of collected organics from the filter. Consequently, the extent of overestimation cannot be estimated. In the current study, a denuder-based sampling method was used, representing an improved method to separate gases from particles (Turpin et al., 2000) and allowing the upper limit of the overestimation to be estimated. The investigation of different SOA formation pathways is made by conducting time-resolved, speciated measurements of gas/particle partitioning of oxygenated semi-volatile/intermediate-volatility organic compounds (S/IVOCs) in the ambient atmosphere with this modified TAG. The factors affecting these pathways are investigated using temporal variability of measured gas/particle partitioning of organic species in combination with supporting measurements, such as RH. This study improves the understanding of SOA formation in the ambient atmosphere and likely lead to useful parameterization of SOA formation.

2.3 Methods
2.3.1 Sampling and analysis

A modified TAG instrument was deployed to measure organic species in both gas and particle phases during the CalNex campaign from May 31st to June 27th, 2010 at the Bakersfield California Supersite. The modification was made before this field campaign by adding an active charcoal denuder (30 mm OD, 40 cm length, ~490 channels, Mast carbon, UK) into the sampling inlet as a parallel sampling line to a bypass line made of stainless steel tubing. The denuder was housed inside a home-made aluminum cylindrical tube with a tapered cap in each end connecting to the sampling line upstream and downstream.

Detailed description of operation of TAG can be found elsewhere (Williams et al., 2006; Worton et al., 2011). Only the sampling and operation relevant to this study are described here. During the sampling, ambient air at 10 L/min was sampled from the center of a main flow, drawn from approximately 5 meters above ground at 200 L/min, and sampled through a sharp cut PM2.5 cyclone (10 L/min, BGI Inc., Waltham, MA). Downstream of the cyclone, a flow split was made to discard 10% of air flow. Subsequently, 90% of the ambient flow was sampled through the denuder line (or the bypass line) and delivered into a customized Collection and Thermal Desorption cell (CTD) through a 9 L/min critical orifice for collection of organics. The aerodynamic particle diameter corresponding to 50% collection is ~0.07 μm so that the entire accumulation mode mass falls within the instrument’s collection range (Williams et al., 2006). Gas/particle separation was achieved by alternating ambient air between the denuder line and the bypass line. The samples collected through the denuder (“denuded samples”) were expected to be only particle phase organics while those collected through the bypass line (“undenuded samples”) were the total organics, the sum of the collected gas and particle phase organics. The sampling duration of each sample was 90 minutes from May 31st to June 9th (Sampling period I) and 30 minutes from June 10th to 27th (Sampling period II). The CTD was maintained at 28°C during the ambient sampling and was continuously held at the same temperature for one minute to purge residual air from the CTD with a helium flow of 20 ml/min at the conclusion of ambient sampling. Following the purge, the thermal desorption of collected organics were carried out in a helium flow by heating the CTD from 28°C to 300°C at a rate of ~30°C/min and held at 300°C for nine minutes followed by thermal injection into a gas chromatograph. The chromatographic separation of organic species was achieved by a capillary GC column (Rxi-5Sil MS; 30 m length, 0.25 mm i.d., 0.25 μm film thickness, Restek). The GC oven temperature was held at 45°C for 18 minutes for the sample injection from the CTD to GC followed, in order, by: 1) a ramp to 150°C at 15°C/min; 2) a ramp from 150°C to 330°C at 9°C/min and 3) a hold at 330°C for 4 minutes. Identification and quantification was achieved using a quadrupole mass spectrometer (Agilent, 5973) calibrated based on responses to authentic standards that were manually injected into the CTD at regular time intervals throughout the campaign (Kreisberg et al., 2009).

The gas collection efficiency of the denuder was determined in the beginning, middle and end of the campaign by the difference of the amount of gas-phase organics downstream of the denuder line and the bypass line with a Teflon coated fiber filter placed upstream of the cyclone to remove particles. Particle penetration through the denuder was determined using an optical particle spectrometer (Droplet Measurements, model UHSAS) to measure the number size distributions of ambient particles at both the inlet and outlet of this denuder before this campaign.
A broad suite of complementary measurements were concurrently made at this site, including a full range of meteorological, trace gas and aerosol measurements. The measurements utilized in this study included non-refractory PM$_{1.0}$ inorganic and organic aerosol components, carboxylic acid group, gas-phase ammonia and meteorological data. Non-refractory PM$_{1.0}$ aerosol components were measured by an Aerodyne high-resolution time-of-flight Aerosol Mass Spectrometer (HR-ToF-AMS) using the methods described in Liu et al. (2012). PM$_{1.0}$ was also collected by Teflon filters for measurements of the organic acid group (-COOH) by Fourier transform infrared (FTIR) spectroscopy (Liu et al., 2012). Gas-phase ammonia was measured using an Ambient Ion Monitor/Ion Chromatograph (AIM-IC) (Markovic et al., 2012).

2.3.2 Particle-phase fraction calculations

Because the particle-phase and total organics were not collected simultaneously, measured fraction of a given compound in the particle phase ($f_{\text{part}}$) in sample $n$ is calculated using the particle-phase concentration ($C_{\text{part,n}}$) divided by the average of the previous and subsequent total concentrations ($C_{\text{total,n-1}}$, $C_{\text{total,n+1}}$):

$$f_{\text{part}} = \frac{2C_{\text{part,n}}}{C_{\text{total,n-1}} + C_{\text{total,n+1}}} \quad (2.1)$$

The gas/particle partitioning coefficient ($k_{\text{om}}$) for absorptive uptake into organic aerosol is calculated by the equation described by Pankow (1994):

$$k_{\text{om}} = \frac{RT}{10^6 P_L^0 \delta MW} \quad (2.2)$$

where $R$ is ideal gas constant ($8.2 \times 10^{-5}$ m$^3$ atm mol$^{-1}$ K$^{-1}$), $T$ is temperature (K), $P_L^0$ is the vapor pressure of the pure compound (atm) at the temperature of interest, $\delta$ is the activity coefficient of the compound in the absorbing phase, and $MW$ is the average molecular weight (g mol$^{-1}$) of the absorbing phase. The particle-phase fraction based on partitioning theory ($f_{\text{part,T}}$) was calculated from the partitioning coefficient constant ($k_{\text{om}}$) and the mass concentration of organic aerosols in $\mu$g m$^{-3}$ ($C_{O\text{A}}$):

$$f_{\text{part,T}} = \left(1 + \frac{1}{k_{\text{om}} \times C_{O\text{A}}} \right)^{-1} \quad (2.3)$$

The data collected by other instruments were averaged to match TAG sampling duration of 30 or 90 minutes. The average OA concentration ($C_{O\text{A}}$) from HR-ToF-AMS measurements was $3.7 \pm 1.8$ $\mu$g m$^{-3}$ (0.5 - 11.2 $\mu$g m$^{-3}$). Average temperature was $26 \pm 6$ °C (12 - 40°C). In our study, the theoretical fractions of organic species in the particle phase were calculated using the measured average temperature and average OA concentration ($T = 26^\circ$C, $C_{O\text{A}} = 3.7$ $\mu$g m$^{-3}$) and both molecular weight ($MW = 200$ g mole$^{-1}$) and activity coefficient ($= 0.3$ and 3) from literature (Pankow, 1994; Seinfeld and Pankow, 2003). Subcooled vapor pressures used in this study were from The Estimation Programs Interface (EPI) Suite developed by the US Environmental Protection Agency's Office of Pollution Prevention and Toxics and Syracuse Research Corporation (SRC).
2.4 Results and discussions

More than 150 compounds were measured by TAG, covering a broad vapor pressure range and different functional groups (Figure 2.1). Most identified compounds were present in the vapor pressure range of S/IVOCs defined by Robinson et al., (2007). The gas/particle partitioning of three oxygenated compounds, pinonaldehyde, phthalic acid and 6, 10, 14-trimethyl-2-pentadecanone, are discussed in detail to explore SOA formation in the ambient atmosphere. Pinonaldehyde is a major product of α-pinene ozonolysis with gaseous yields of pinonaldehyde being 0.39-0.69 (Liggio and Li, 2006). Phthalic acid and 6, 10, 14-trimethyl-2-pentadecanone and have been used as SOA tracers in both chemical mass balance and positive matrix factorization model calculations (Zheng et al., 2002; Shrivastava et al., 2007; Williams et al., 2010a).

Although \( f_{\text{part}} \) of the compounds of interest were overestimated because gas-phase organics were only partially collected by the collection cell used in this study which was designed for collecting particle-phase organics, the extent of overestimation can be indicated by \( f_{\text{part}} \) of \( n \)-alkanes. Firstly, the efficient gas/particle separation made by the denuder limits the source of the overestimation to the collection cell. Average collection efficiencies of the denuder for pinonaldehyde, phthalic acid and 6, 10, 14-trimethyl-2-pentadecanone were over 98%. Average losses of the particle number inside the denuder were less than 10% for particle sizes spanning the particle spectrometer’s range (0.05–1 μm) and the performance of this denuder was stable over one month ambient measurements (see Appendix A, Figures A1 and A2). Secondly, the gas/particle partitioning of \( n \)-alkanes can be well described by the gas/particle absorption partitioning theory (Fraser et al., 1997) and \( n \)-alkanes have the lower or same adsorption coefficient constants on the surface of sampling substrates, relative to other compounds with the same vapor pressure (Goss and Schawarzenbach, 1998). As a result, measured particle-phase fractions of \( n \)-alkanes, the sum of absorptive gas/particle partitioning and overestimation caused by incomplete collection of their vapors, are the upper limit of the overestimation of TAG measurements in the vapor pressure range of these \( n \)-alkanes.

The reference compounds selected based on the similar subcooled vapor pressure for pinonaldehyde, phthalic acid and 6, 10, 14-trimethyl-2-pentadecanone are \( n \)-tetradecane, \( n \)-heptadecane and \( n \)-nonadecane, respectively (Figure 2.2). If measured particle-phase fractions of oxygenated organics are far larger than those of their reference compounds, as is the case for phthalic acid and pinonaldehyde, additional SOA formation pathways must occur, beyond absorptive gas/particle partitioning and overestimation due to under collection of gas-phase organics.

2.4.1 Pinonaldehyde

The mean \( f_{\text{part}} \) of pinonaldehyde was 20 ± 20%, much higher than its reference compound of \( n \)-tetradecane (Figure 2.2). The fraction of pinonaldehyde in the particle phase was observed to increase as RH increased (Figure 2.3A), but the fraction contributed by its partitioning into aerosol water is negligible even if a ratio of aerosol water to dry aerosol mass equal to one is assumed and all of the aerosol water is available to take up pinonaldehyde. Moreover, this assumed ratio of the water content to dry mass is inconsistent with the average RH of 34%
during TAG measurements because a ratio of generally less than 0.3 is expected at this average RH (Khlystov et al., 2005; Schuster et al., 2009; Engelhart et al., 2011). The particle-phase pinonaldehyde reported here is with negligible sampling artifacts due to adsorption of gas-phase pinonaldehyde to the collection cell because the denuder efficiently removed organic vapors. Therefore, our observations of particle-phase pinonaldehyde clearly show that gas-phase pinonaldehyde had been converted into forms with the lower vapor pressures prior to the collection.

Low-volatility compounds (e.g. oligomers) formed from monomers with direct involvement of pinonaldehyde have been observed in chamber experiments (Tolocka et al., 2004; Liggio and Li, 2006; Tillman et al., 2010) and ambient samples (Tolocka et al., 2004). In our study, low-volatility compounds were measured as a pinonaldehyde monomer, consistent with previous TAG measurements in a forest area (Worton et al., 2011). The reason may be attributed to the use of the thermal desorption method which could decompose low-volatility compounds into their original monomers (Jang et al., 2002). These low-volatility compounds were not directly measured in our study, but the variability in the concentrations of measured pinonaldehyde can still be useful to investigate factors affecting the formation of low-volatility compounds. In the following discussion, the term of particle-phase pinonaldehyde is taken to include all low-volatility compounds formed with direct involvement of pinonaldehyde. The cation-to-anion ratio, calculated using molar concentrations of ammonium and anions (\(= 2\times[\text{sulfate}] + [\text{nitrates}]\)) measured by HR-ToF-AMS, is used as an indicator of availability of acids in our study. The presence of excess ammonium is indicated when the cation-to-anion ratio is greater than one and the presence of excess acids is indicated when the ratio is less than one.

Particle-phase pinonaldehyde was observed while the cation-to-anion ratio calculated using ammonium, sulfate and nitrate was greater than one, indicating that the presence of inorganic acids were not necessary for the formation of particle-phase pinonaldehyde (Figure 2.4). Since FTIR measures carboxylic acids as an acid group (-COOH) (Russell et al., 2009), excess organic acids, which were not neutralized, were present in particles (Figure 2.4). Our observations of particle-phase pinonaldehyde and availability of acids demonstrate the observations of chamber experiments of \(\alpha\)-pinene ozonolysis which show that oligomers are formed on the neutralized ammonium sulfate particles (Gao et al., 2004; Tolocka et al., 2004) and organic acids produced from gas-phase hydrocarbon oxidation are sufficient to catalyze these heterogeneous reactions (Gao et al., 2004).

Laboratory studies have shown that high aerosol acidity leads to the high yield of oligomers from pinonaldehyde (Liggio and Li, 2006) and the oxidation products of \(\alpha\)-pinene (Gao et al., 2004; Tolock et al., 2004). However, the laboratory observed trend was not displayed by the relationship between \(f_{\text{part}}\) of pinonaldehyde and organic acids measured by FTIR (Figure 2.4). The reason could be that the contribution of organic acids to the aerosol acidity cannot be directly indicated by the cation-to-anion ratio because different organic acids have different dissociation constants and organic acids and their conjugate base can serve as a buffer solution. The cation-to-anion ratio of inorganic ions shows a general trend that the high cation-to-anion ratio was along with the low \(f_{\text{part}}\) of pinonaldehyde (Figure 2.4), but the acidity estimated based on the cation-to-anion ratio would have a large uncertainties when the cation-to-anion ratio is
near one (Xue et al., 2011). Moreover, this trend was not followed when other factors affecting the relationship between aerosol acidity and $f_{\text{part}}$ of pinonaldehyde were considered, such as RH (Figure 2.3B) which can change the composition and mass of SOA (Nguyen et al., 2011) and the aerosol acidity (Liggio and Li, 2006). As shown in Figure 2.3A, $f_{\text{part}}$ of pinonaldehyde exhibited a positive dependence on RH in Sampling Period I, consistent with previous TAG measurements in a forested area (Worton et al., 2011). However, the cation-to-anion ratio calculated from inorganic ions did not consistently decrease as RH increased (Figure 2.3B). As a result, the effect of the aerosol acidity on $f_{\text{part}}$ of pinonaldehyde is not shown by the relationship between the cation-to-anion ratio and $f_{\text{part}}$ of pinonaldehyde.

In comparison with Sampling Period I, $f_{\text{part}}$ of pinonaldehyde was generally lower in Sampling Period II and showed a different dependence on RH (Figures 2.3A and 2.4) while this pattern was not observed for its reference compound, $n$-tetradecane. The difference in relationships between $f_{\text{part}}$ and RH observed during these two sampling periods is also supported by another independent measurement of the cation-to-anion ratio which was different in two sampling periods (Figures 2.3B and 2.4). Therefore, the observed relationship between $f_{\text{part}}$ of pinonaldehyde and RH represents the real relationship between them in the atmosphere.

The enhancement in $f_{\text{part}}$ of pinonaldehyde at the high RH was not observed in Sampling Period II, although the particle-phase concentration of pinonaldehyde was observed to increase as RH increased. The relationships between RH and $f_{\text{part}}$ of pinonaldehyde suggest that RH favors the formation of particle-phase pinonaldehyde in the atmosphere, but it isn't the primary factor affecting the yields of particle-phase pinonaldehyde. Further studies are needed to examine the effect of RH on the yield of particle-phase pinonaldehyde in the atmosphere and laboratory.

2.4.2 Phthalic acid

The mean $f_{\text{part}}$ of phthalic acid was 60 ± 20%, substantially higher than that of its reference compound, $n$-nonadecane (Figure 2.2). To reproduce the mean $f_{\text{part}}$ for phthalic acid using gas/particle partitioning theory, a significantly lower activity coefficient (~5×10⁻³) than the estimated range from 0.3 to 3 for SOA in the atmosphere (Seinfeld and Pankow, 2003) would be needed. The partitioning of phthalic acid into aerosol water cannot explain the mean $f_{\text{part}}$ for phthalic acid based on its Henry's law constant of 2.0×10¹¹ atm m⁻³ mol⁻¹ (USEPA EPI suite) and the assumption that the ratio of aerosol water content to the dry aerosol mass is one and all aerosol water is available to take up phthalic acid. The dissociation of phthalic acid was also considered, but the contribution due to its dissociation to its mean $f_{\text{part}}$ was negligible even when pH was estimated by neutralized inorganic ions without inclusion of other organic acids. Moreover, the aerosol water content is unlikely to be that high at the average RH of 34% in the atmosphere (Khlystov et al., 2005; Schuster et al., 2009; Engelhart et al., 2011). Therefore, there must be an additional partitioning mechanism whereby particle-phase phthalic acid is formed.

We infer that a likely pathway for phthalic acid partitioning to particles is through its reaction with ammonia. This is supported by the presence of excess ammonium in the particle phase indicated by the cation-to-anion ratio of inorganic species measured by HR-ToF-AMS (Figure 2.4). Evidence for reactions between organic acids and gas-phase ammonia is provided by Na et al. (2007), who observed that ammonia could dramatically increase SOA yields by reactions with organic acids in chamber experiments. Furthermore, the positive correlation
between gas-phase ammonia and \( f_{\text{part}} \) of phthalic acid (linear regression \( R^2 = 0.8 \) between average \( f_{\text{part}} \) and the ammonia concentration) supports the hypothesis that phthalic acid partitioning to particles is through reactions with gas-phase ammonia (Figure 2.5).

Reactions with ammonia can convert phthalic acid into ammonium salts with the low vapor pressures and subsequently favor its partitioning into particles. Subcooled vapor pressures of the formed salts can be more than 100 times lower than that of phthalic acid, even if just monoammonium salt was formed (order of magnitude of the subcooled vapor pressure drop is estimated using the vapor pressure drop of organic acids after forming ammonium salt from USEPA EPI suite). Additional support needed for the presence of phthalic acid ammonium salts is that these salts can be measured as phthalic acid by the TAG using a thermal desorption technique to extract collected organics. This support is given in Hajek et al. (1971) wherein the investigation of the thermal decomposition of ammonium salt of isophthalic acid shows that simultaneous release of both ammonia and isophthalic acid molecules from diammonium salts occurs without dehydration or amide formation.

### 2.4.3 6, 10, 14-trimethyl-2-pentadecanone

The mean \( f_{\text{part}} \) of 6, 10, 14-trimethylpentadecanone was 4 ± 2%, similar to that of its reference compound, \( n \)-heptadecane, suggesting that there is no reactive uptake of it on particles during the campaign contrary to the observations of pinonaldehyde and phthalic acid. These results are in agreement with previous studies on the gas/particle partitioning of ketones (Esteve and Noziere, 2005; Kroll et al., 2005). Esteve and Noziere (2005) suggested that aldol condensation was too slow to contribute significantly to SOA under atmospheric conditions. Kroll et al. (2005) showed that ketones did not produce observable volume growth in the presence of acidic seeds even with concentrations of over 500 ppb. Other measured compounds with a ketone functional group in this study, such as benzophenone and 1-hydroxycyclohexy-phenyl methanone, were also present primarily in the gas phase.

### 2.5 Conclusions and atmospheric implications

Measurements of organic compounds in both gas and particle phases improves our understanding of SOA formation mechanisms in the atmosphere. Ketones observed in our study were present predominantly in the gas phase, suggesting that observed reactive uptake to aerosols does not occur and absorption into aerosols is the dominant pathway for them to contribute to SOA in the atmosphere. While absorption of gas-phase phthalic acid into the particles can contribute to observed concentrations of particle-phase phthalic acid, the major pathway to form particle-phase phthalic acid is likely attributed to reactions with gas-phase ammonia. This mechanism is expected to cause high O/C ratios in ammonia enriched areas because reactions with ammonia favor the uptake of carboxylic acids on particles. Therefore, the formation of condensable salts can be a significant area of uncertainty for SOA formation in ammonia enriched areas. Pinonaldehyde contributes to SOA through reactive uptake. Observations of particle-phase pinonaldehyde when inorganic acids were neutralized show that inorganic acids are not required for occurrence of reactive uptake of pinonaldehyde beyond that predicted by the gas/particle partitioning theory. The effect of aerosol acidity on the partitioning of pinonaldehyde into particles observed in laboratory studies is not displayed by our observations using the relationship between the cation-to-anion ratio and the fraction of
pinonaldehyde in particles. The relationship between particle-phase pinonaldehyde and RH suggests that aerosol water content likely plays a significant role in the formation and accumulation of particle-phase pinonaldehyde in the absence of inorganic acids, but it is not the primary one. Our observations highlight that further studies are needed to examine the effects of RH and organic acids on the reactive uptake of pinonaldehyde and other aldehydes on neutral seed particles.

In-situ measurements of both gas- and particle-phase organic compounds in the atmosphere have clearly shown that multiple gas/particle partitioning pathways are present in the atmosphere and other gas/particle partitioning pathways significantly improve the SOA yields, relative to absorptive gas/particle partitioning. Observations of particle-phase pinonaldehyde show that inorganic acids are not required for the occurrence of reactive uptake of pinonaldehyde into particles and subsequently suggest that this pathway is likely widespread. However, the gas/particle partitioning of phthalic acid favored by formation of condensable salts indicates that this pathway could be only significant in ammonia-rich environments. Therefore, further in-situ, time-resolved measurements of gas/particle partitioning covering more oxygenated organic compounds are needed to investigate SOA formation in different areas and provide parameterization for inclusion of these pathways in SOA models. Additionally, since our results show that each of three SOA tracers investigated here has distinct pathways to partition to particles, multiple tracers are needed in source apportionment models to adequately represent SOA formation. However, these tracers are present in both gas and particle phases. As a result, it raises a concern about the accuracy of the source apportionment models using these SOA tracers without correcting gas adsorption on the sampling substrates.

2.6 References


2.7 Tables and figures

**Figure 2.1** O/C ratios of organic compounds measured by TAG as a function of subcooled vapor pressure at 25°C. The compounds of interest are colored in red.
Figure 2.2 Average measured fractions for oxygenated SVOCs (solid markers) and their corresponding reference compounds (empty markers) are shown as markers. Error bars are one standard deviation. The solid and dashed lines are the predicted fractions of organic compounds in the particle phase based on the gas/particle partitioning equation.
Figure 2.3 A) Measured fraction of pinonaldehyde in the particle phase as a function of RH (average 5 or 6 points in each bin in Sampling Period I; 12 points in each bin in Sampling Period II). Error bars are one standard deviation. The X-axis values are the average value of RH in each bin. Fractions larger than 3 standard deviations outside of the mean in each sampling period are considered outliers and excluded in this plot. B) The average cation-to-anion ratio of inorganic species (sulfate, nitrate and ammonium) as a function of RH (average 7 points in each bin in Sampling Period I; 27 points in each bin in Sampling Period II).
Figure 2.4 The temporal change of the cation-to-anion ratio of measured inorganic ions, carboxylic acid group and the fraction of pinonaldehyde in the particle phase.
**Figure 2.5** Fraction of phthalic acid in the particle phase as a function of the concentration of gas-phase ammonia. Each bin of seven bins has 18 data points.
Chapter 3
Sources of organic aerosol investigated using organic compounds as tracers measured during CalNex Bakersfield

3.1 Abstract
To investigate the major sources of organic aerosol (OA) and provide insights into secondary organic aerosol (SOA) formation, positive matrix factorization (PMF) analysis was performed on a large set of organic species measured by in-situ Thermal desorption Aerosol Gas chromatography-mass spectrometry (TAG) during the California at the Nexus of Air Quality and Climate Change (CalNex) campaign in Bakersfield, CA from May 31st through June 27th, 2010. Six OA source factors were identified, including one representing primary organic aerosol (POA), four different types of secondary organic aerosol (SOA) representing local, regional, and nighttime production, and one representing a complex mixture of additional OA sources that were not further resolvable. The average POA contribution to total OA was 15% throughout the campaign with a diurnal range from 10% during the day to 19% at night and motor vehicles are suggested to be a significant contributor. The contribution of the four distinct types of SOA to total OA averaged 72% throughout the campaign and varied diurnally from 78% during the day to 66% at night. The complex mixture of additional OA sources in the final factor contributed an average of 14% to total OA. Both regional and local SOA were significant OA sources, but regional SOA had a larger contribution to total OA than local SOA during the day. Biogenic VOC oxidation products contributed a significant fraction of total OA at night. We conclude that SOA was formed through multiple pathways that peak at different times of day, with the largest SOA contribution occurring through condensation of gas-phase oxidation products onto particles. Effective control measures to reduce OA in Bakersfield should focus on reducing sources of VOCs that serve as precursors to SOA during the day, and we conclude that the major source of SOA is regional.

3.2 Introduction
Organic compounds constitute a major mass fraction (20-90%) of atmospheric fine particulate matter in most environments (Kanakidou et al., 2005). These organic compounds have been categorized into either primary organic aerosol (POA), directly emitted from various primary sources such as food cooking and vehicle exhausts (e.g. Schauer et al. 1999a, b, 2002a), or secondary organic aerosol (SOA), formed in the atmosphere through chemical reactions (Odum et al., 1997; Jang et al., 2002; Robinson et al., 2007; Kroll and Seinfeld, 2008). The majority of organic aerosol (OA) in rural and urban areas is secondary (Zhang et al., 2007; Shrivastava et al., 2007; Williams et al., 2010; Jimenez et al., 2009). SOA and POA have been shown to have different adverse human health effects (Li et al., 2009), so quantifying source contributions to OA has important implications for air quality regulation.

Source apportionment between POA and SOA has generally been done in one of three ways; (i) bulk organic spectra measured by the Aerodyne Aerosol Mass Spectrometer (e.g. Jimenez et al., 2009; Zhang et al., 2011), (ii) thermal-optical elemental carbon/organic carbon analyzer (e.g. Turpin et al., 1991; Strader et al., 1999) and (iii) molecular speciation measured by
gas chromatograph/mass spectrometer (GC/MS) (e.g. Shrivastava et al., 2007; Williams et al., 2010). In comparison with bulk organic analysis, the molecular speciation resolves a smaller fraction of OA, but the identified organic compounds can serve as tracers for specific source types (Schauer et al., 1996; Schauer and Cass, 2000) and provide information for understanding SOA formation and transformation in the atmosphere (e.g. Kleindienst et al., 2007; Williams et al., 2010). Two common source apportionment methods using organic compounds are the chemical mass balance (CMB) and positive matrix factorization (PMF) models (e.g. Schauer et al., 1996; Shrivastava et al., 2007; Williams et al., 2010).

The CMB model uses organic compounds to estimate OA contributions from a variety of sources, such as biomass burning, meat cooking, or diesel vehicle emissions, but requires a priori knowledge of their source profiles as inputs to solve the model (Schauer et al., 1996; Schauer and Cass, 2000; Zheng et al., 2002). As a result, CMB analysis is sensitive to the provided source profiles (Robinson et al., 2006; Subramanian et al., 2007). The products and SOA yields measured for a few known volatile organic compounds (VOCs) oxidized in chamber experiments have been considered as source profiles of SOA (Kleindienst et al., 2007; Stone et al., 2009). However, the relevance of these source profiles is limited to these known compounds and the specific experimental conditions used in the chamber to derive the profiles. Source profiles for SOA are difficult to establish due to the complexity of oxidation processes and the wide range of precursors and atmospheric conditions, so CMB models have typically been unable to provide adequate constraints on the SOA mass or composition. Though SOA mass can be estimated by CMB models as the difference between apportioned OA and measured OA, they do not provide detailed information about SOA sources (Schauer et al., 1996; Zheng et al., 2002; Subramanian et al., 2007).

The PMF model uses covariation between organic compounds to categorize them into unique groups or factors that could represent emissions from contemporaneous source or formation processes (Shrivastava et al., 2007; Zhang et al., 2009; Williams et al., 2010). Since a priori knowledge of source profiles is not required as inputs, factors likely representing poorly constrained or unknown sources, such as SOA, can be determined. Characterization of these factors based on their compositions and temporal variations therefore provide an effective tool for differentiating complex sources (Shrivastava et al., 2007; Zhang et al., 2009; Williams et al., 2010). Past work has demonstrated the ability of this method to separate multiple SOA types with distinct diurnal patterns in the absence of known source profiles (Williams et al., 2010). Despite its utility for source apportionment, PMF analysis is not widely performed on organic compounds in the particle phase because it typically requires a larger number of samples which poses significant challenges when measurements of speciated OA are mainly made by filter sampling with 24-hour collection periods in the field followed by solvent extraction in the laboratory for analysis (Jaeckels et al., 2007; Shrivastava et al., 2007). Recently, speciated measurements of OA have been facilitated by the development of an in-situ Thermal desorption Aerosol Gas chromatography instrument (TAG) which provides hourly time resolved chemical speciation of OA (Williams et al., 2006). As a result of the hourly time resolution, TAG provides as many data points in a few weeks of measurements as solvent extracted filters are able to in a year, thus enabling multivariate statistical analyses. TAG data has been analyzed by PMF in previous work to resolve nine different types of OA in an urban environment, including various SOA and POA sources (Williams et al., 2010).
In this work, we focus on data from a field site in Bakersfield, CA, one of two supersites of the CALifornia at the NEXus between Air Quality and Climate Change (CalNex) campaign in June 2010. Before the CalNex campaign. Previous studies of source contributions to OA in the Bakersfield area were based on the EC-tracer method and CMB and focused on OA in winter (Magliano et al., 1999; Strader et al., 1999; Schauer et al., 2000). Neither the EC-tracer method nor CMB was able to provide insights into different SOA types. Recently, new results of studies of major OA components and SOA formation in Bakersfield areas during the CalNex campaign have been reported. Liu et al. (2012) has shown that SOA was the dominant component of OA in the Bakersfield site, accounting for 80~90% of OA, and the majority of SOA was likely formed through gas-to-particle condensation of daytime oxidation products of volatile organic compounds (VOCs). Rollins et al. (2012) reported that organic nitrate aerosol accounted for 1/3 of nighttime increase of OA and suggested that reductions in NOx emissions can reduce OA concentrations. In this study, we take a different approach to investigate source contributions to OA and atmospheric processes to form SOA using organic tracers, instead of bulk organic mass spectra (Liu et al. 2012) and organic nitrates (Rollins et al. 2012). These organic tracers bear specific information for OA sources and SOA formation pathways. For example, Zhao et al. (2012) has shown that multiple gas-to-particle partitioning pathways contributed to SOA formation in Bakersfield areas by examining the measured gas/particle partitioning of known SOA tracers. By performing PMF analysis on these organic tracers, including primary and secondary tracers measured by the TAG in Bakersfield, CA, distinct factors can be determined, which are associated with specific source types and atmospheric processes in the context of an urban area and the agricultural, natural, and industrial sources present in the region. The effective control strategy to reduce OA concentrations are suggested based on our investigation of source contributions to OA and formation pathways of SOA.

3.3 Methods
3.3.1 Sampling and chemical analysis

Speciated measurements of particle-phase organics were made with the TAG instrument during the CalNex field campaign from May 31st to June 27th 2010 at the Bakersfield supersite, CA. A description of the TAG is provided by Williams et al. (2006), and details on updates and operation of the TAG used in this work are provided in Chapter 2. Briefly, the TAG was updated by addition of a denuder into the sampling inlet so that there were two parallel sampling lines, one denuder line and one bypass (no denuder) line made of stainless steel tubing. During sampling, ambient air was pulled at 10 L/min through a PM2.5 cyclone from the center of a main sampling flow of 200 L/min drawn from ~5 meters above the ground through a 6-inch (i.d.) rigid duct and gas- and particle-phase organics were collected by an impactor. The sampling flow was alternated between the denuder line to collect only particle-phase organics, and the bypass line to collect total organics, the sum of gas- and particle-phase organics. Following sampling, the collected organics were thermally desorbed and injected into a gas chromatograph-mass spectrometer (GC/MS) for analysis. Identification and quantification were achieved using a quadrupole mass spectrometer (Agilent, 5973) calibrated based on responses to authentic standards that were manually injected into the impactor cell at regular intervals throughout the campaign (Kreisberg et al., 2009). During this campaign, TAG measurements were made in two
periods wherein the sampling durations were different. The duration of each sample was 90 minutes from May 31st to June 9th (Sampling period I) and 30 minutes from June 10th to 27th (Sampling period II). Over the 27-days of measurements, 244 samples of speciated OA were acquired and over 100 particle-phase organic compounds were identified and quantified.

Supporting measurements conducted concurrently that are relevant to data analysis and discussion in this study included detailed characterization of meteorological conditions, OA, and volatile organic compounds (VOCs). Wind speed and direction were monitored by a propeller wind monitor (R.M. Young, 5130). Details on measurements of submicron OA made with an Aerodyne High-Resolution Time-of-Flight Aerosol Mass Spectrometer (HR-ToF-AMS) are provided in Liu et al., (2012). Gas-phase VOCs were measured by an automated in-situ GC-FID/MS system (Gentner et al., 2012). In addition, ozone was measured using a UV photometric ozone analyzer (Dasibi Inc., model 1008 RS).

3.3.2 PMF procedures

The positive matrix factorization (PMF) model describes observed concentrations of organic species as the linear combination of the contributions from a number of sources with constant profiles (Paatero and Tapper, 1993; Hopke, 2003; Reff et al., 2007; Ulbrich et al., 2009):

\[ x_{ij} = \sum_p g_{ip} f_{pj} + e_{ij} \]  

(3.1)

where \( x_{ij} \) is the concentration of species \( j \) measured in sample \( i \), \( g_{ip} \) is the contribution of factor \( p \) to sample \( i \), \( f_{pj} \) is the concentration of species \( j \) in factor \( p \) and \( e_{ij} \) is the residual not fit by the model. PMF solves the equations for \( g_{ip} \) and \( f_{pj} \) to best reproduce \( x_{ij} \) without \textit{a priori} knowledge of them based on the selected number of factors (\( p \)). The solution to PMF minimizes the object function, \( Q \), defined as:

\[ Q = \sum_{i=1}^{n} \sum_{j=1}^{m} \frac{(e_{ij})^2}{s_{ij}} \]  

(3.2)

where \( s_{ij} \) is the estimated uncertainty of species \( j \) measured in sample \( i \).

As the PMF model allows each data point to be individually weighted, the uncertainties associated with organic compounds at different concentration levels are estimated separately. By estimating the uncertainty using this approach, undetected data points can be weighted, so compounds or days with many missing points do not need to be removed. This approach provides an advantage over many other methods by achieving longer timelines of observations (Pollissar et al., 1998). In our study, data points, which were not detected because of the low concentration, were assigned a concentration of one third of the compounds detection limit, and uncertainty was assigned to be four times its concentration, similar to the estimation made in Pollissar et al. (1998). The uncertainty for the measured data point was estimated using the same equations described in Williams et al. (2010). In contrast to that work, however, the detection limit of each compound was used in the current study to better account for the instrument noise, instead of instrument precision derived from the standard deviation of the difference between consecutive data points used in Williams et al., (2010).

To investigate the major OA sources in this study, only the particle-phase organics were included in the PMF analysis. In Appendix B, we demonstrated that the inclusion of gas-phase
contributions to semi-volatile organic compounds (SVOCs) significantly bias the results of source apportionment. The criteria we used for inclusion of compounds were that the timeline of observations covered the entire period of TAG measurements, the compounds were in the particle phase, and the percentage of above detection limit data points was greater than 50% over the full timeline of observations. As a result, 30 compounds (listed in Figure 3.1) in 244 samples were included in the PMF analysis. PMF was applied using the Igor based PMF Evaluation Panel v2.04 (Ulbrich et al., 2009).

Detailed discussions of determination of the optimum number of PMF factors can be found in the literature of Reff et al. (2007), Ulbrich et al. (2009) and Williams et al. (2010), and interpretability of these factors is an important criterion. In our study, the linkage between factors and specific OA source types or atmospheric processes was initially made based on the existing knowledge of specific sources of individual compounds in each factor and was further corroborated by their diurnal profiles and supporting measurements, such as VOCs and meteorological conditions.

Although the sum of concentrations of all species included in PMF analysis accounts for only a small fraction of AMS measured OA, the mass contribution of each factor to measured OA can be calculated by a multivariate fit of the temporal contributions of PMF factors onto total AMS measured OA mass concentrations (Reff et al., 2007; Williams et al., 2010). The sum of these calculated mass contributions are defined as reconstructed OA. The regression coefficients for factors provide an additional constraint on the number of factors to be included, since negative regression coefficients are a good indication that the number of factors exceeds the optimum (Reff et al., 2007).

3.4 PMF results

The variability of the TAG data is best explained by six factors with a value of $Q/Q_{expected}$ equal to 2.5 with $F_{peak}$ set to 0. Since the interpretability is an important criterion in determining the optimal number of factors, 5-factor and 7-factor solutions were also explored. In comparison with the 6-factor solution, the 5-factor solution resulted in a convolution of factors 1 and 5. The 7-factor solution split factor 6, which is a mixture of anthropogenic and biogenic SOA factors, without providing additional meaningful information. We therefore chose to examine the 6-factor solution in detail to characterize the major components of total OA. To facilitate discussion, we define 8:00-20:00 PST as the day and 20:00-8:00 PST as the night.

3.4.1 Factor 1: Local POA

We define factor 1 as local POA. Factor 1 has the highest contribution from hydrocarbons among the six factors (Figure 3.1) and has a diurnal profile with higher concentrations at night, consistent with the buildup of local emissions into a shallow nighttime boundary layer (Figure 3.2). Highest concentrations of this factor occurred during the night, at low wind speeds, and with wind coming from all directions (Figures 3.2 and 3.3) indicating that it originated mainly from local sources because Bakersfield is located near the southern end of the San Joaquin Valley. One of the two main oxygenated compounds present in this factor was 9-fluorenone, which has been identified as a primary emission from motor vehicles (Schauer et al., 1999). The other prominent oxygenated compound present in this factor was phthalic acid.
Phthalic acid can be produced from oxidation of naphthalene and has been used as a tracer for anthropogenic SOA (Schauer et al., 2002b; Fine et al., 2004; Shrivastava et al., 2007), but its average concentration in this factor accounts for less than 1% of its total measured concentration so we consider this a very minor contribution of SOA adding to this POA dominated factor.

The major known POA sources in Bakersfield are wood burning, meat cooking and motor vehicles (Schauer et al., 2000) and the average contribution of unknown POA sources to OA was small (~10%) (Schauer et al., 2000; Strader et al., 1999). Our POA factor correlated with several measured hopanes (r > 0.54), which were not included in the PMF analysis because over 50% of their data points were below detection limit. The good correlations with hopanes suggest that this factor is related to motor vehicles that have been identified as the dominant source of hopanes in urban environments (Schauer et al., 1996; 2002; Schauer and Cass, 2000). Tracers for meat cooking, such as palmitoleic acid and cholesterol, were not detected and the detected tracer for wood burning, retene, was only observed in a few particle-phase samples.

3.4.2 Factor 2: A mixture of OA sources

We define factor 2 as a mixture of OA sources with contributions from both POA and SOA. While hydrocarbons accounted for a significant fraction of this factor, relative to local POA, there was a smaller fraction of each PAH and a larger fraction of each phthalate (Figure 3.1). Additionally, an SOA tracer, 6, 10, 14-methyl-2-pentadecanone (Shrivastava et al., 2007), apparently contributed to this factor, relative to the POA factor wherein this tracer was not present. The composition of this factor suggests that it was affected by a variety of different sources and atmospheric chemical processes. The large difference in the concentration range in alternating intervals of its 2-hr diurnal profile is caused by very different concentrations observed during two sampling periods. The concentrations in Sampling Period I were higher than those in Sampling Period II and were plotted in every other sampling interval. The diurnal profile for the concentration of this factor was examined separately for each sampling period, but neither period had a clear diurnal cycle, suggesting again that this factor is mostly likely a mixture of OA sources.

3.4.3 Factors 3: SOA1

We define factor 3 as SOA1. The contribution of organic species to this factor was dominated by oxygenated compounds (Figure 3.1). Methyl phthalic acid has been used as a tracer for anthropogenic SOA (Fine et al., 2004; Williams et al., 2010) and 9,10-anthraquinone has been observed as oxidation products of PAHs (Helmig et al., 1992; Helmig and Harger, 1994). The elevated concentration of this factor during 20:00-24:00 PST (Figure 3.2) is attributed to the low wind speed and low temperature, which favors the accumulation of gas-phase oxygenated compounds in the afternoon and subsequent condensation of them onto particles to contribute to SOA. These condensable oxygenated compounds can be formed locally, and probably include some organic nitrates (Rollins et al., 2012), and can also be produced upwind and transported to the site during the afternoon along with SOA3 (discussed below).

3.4.4 Factor 4: SOA2

We define factor 4 as SOA2. The dominant contribution of organic species to this factor was from two SOA tracers, phthalic acid and methyl-phthalic acid, though other compounds, both hydrocarbons and oxygenated compounds, were present but with much smaller
contributions. Evidence that this factor was associated with photochemically produced SOA is apparent when comparing the diurnal profile of this factor with that of the ratio of 1,3,5-trimethylbenzene (TMB) to toluene. Because both 1,3,5-TMB and toluene have similar source types in Bakersfield (Gentner et al., 2012) and 1,3,5-TMB reacts faster with OH radicals than toluene does (Atkinson and Arey, 2003), a smaller ratio of 1,3,5-TMB to toluene indicates that the air mass is more oxidized. An opposite trend between the average diurnal profile of this factor and that of the ratio of 1,3,5-TMB to toluene clearly shows that higher concentrations of this factor occurred when gas-phase organics were more oxidized (Figure 3.2). The highest concentration of this factor occurred in the morning (8:00-9:00 PST) when the wind speed was low (Figure 3.2), indicating that this SOA was formed locally.

3.4.5 Factor 5: SOA

We define factor 5 as SOA3. This factor had its highest concentrations in the afternoon (Figure 3.2) and correlated with ozone (r=0.62). The dominant contribution of organic species to this factor was from oxygenated compounds (Figure 3.1). However, the diurnal patterns of the ratio of 1,3,5-TMB to toluene did not correlate well with this factor, especially in the afternoon (Figure 3.2). The afternoon ratio of 1,3,5-TMB to toluene was close to the nighttime ratio, indicating that observed VOCs were fresh and emitted locally without having undergone significant oxidation. Therefore, we infer that SOA in this factor was not dominantly formed through local VOC oxidation by OH, but instead was composed primarily of regionally formed, transported SOA. This inference is supported by relatively high wind speeds (from the northeast) during the observations of high concentrations of this factor (Figure 3.3) would carry regional source contributions to the observation site, but would dilute local source contributions.

3.4.6 Factor 6: Nighttime SOA (SOA4)

We define factor 6 as nighttime SOA (SOA4). This factor had the highest concentrations at night with the dominant contribution from SOA tracers phthalic acid, methyl phthalic acid, 6, 10, 14-methyl-2-pentadecanone and pinonaldehyde. Pinonaldehyde is a volatile oxidation product of a biogenic VOC, α-pinene, and can contribute to SOA by forming low vapor pressure products, such as dimers (Tolocka et al., 2004; Liggio and Li, 2006). The factors controlling gas/particle partitioning of pinonaldehyde at this site has been analyzed in Chapter 2. Deconvolution of biogenic and anthropogenic contributions is difficult due to the co-variation of chemical transformations leading to SOA, but it is evident that SOA from biogenic VOC oxidation contributed significantly to SOA4 as the highest concentrations of this factor were most frequently observed when the wind blew from the vegetated areas located to the east of the field site (Figure 3.3).

3.5 Reconstructed OA

Reconstructed OA based on TAG-derived factors was compared with measured OA to evaluate the capability of the TAG-derived factors to capture the variability of OA in the atmosphere. Good agreement between the reconstructed and measured OA was demonstrated by differences between them, which were less than 20% for over 70% of observations (Figure B1a). The reconstructed OA had largest uncertainties at the lowest measured OA concentrations and was generally lower than the measured OA at high measured OA concentrations. This could be attributed to larger uncertainties related to organic species when the atmospheric OA
concentration was low and lack of inclusion of additional organic tracers in the current dataset when the atmospheric OA concentration was high (because they were below detection limit in more than 50% of observations or cannot be measured by TAG). As shown by Figure B1b, the negative relative residuals mainly occurred at night. One reason for these negative residuals could be that the TAG did not measure a tracer for organic nitrates which have been shown to significantly contribute to nighttime OA in this region (Rollins et al., 2012).

### 3.6 Source contributions to OA mass

The local POA source accounted for an average 15% of total OA and motor vehicles were shown to have a significant contribution to local POA. The average contribution of POA to total OA could be up to 28% if we assumed that all of the mass of the mixture of OA sources is POA, but this is surely an overestimate. The contributions from other specific primary sources cannot be determined individually in this study because they are small and PMF analysis cannot reliably separate very small factors among the noise and variation of the larger factors. The emissions in the Bakersfield region are highly complex with cars, trucks, oil and gas extraction and refining operations, agricultural activities and other sources all potentially contributing some to these POA factors.

The average contribution of the sum of SOA (SOA1–4) was 72% of total OA and the contribution of SOA could be up to 85% if the mass in the mixture of OA sources is only considered as SOA. The dominance of SOA in total OA derived from organic species in this study is consistent with the results reported by Liu et al. (2012) based on PMF analysis of AMS observations from the same site which showed that 80% to 90% of total OA is accounted for by SOA. We conclude that efforts to control atmospheric OA in this region must focus on understanding and then controlling the sources of SOA precursors or factors leading their transformation into SOA.

We calculated the diurnal cycle of relative contribution from each identified component to OA (Figure 3.4) to highlight the dominant components of OA in different periods of the day and night. The factors contributing the largest fraction of total OA was different during the day and night, but both of them were SOA. The largest daytime contribution to total OA was from SOA3, and this factor accounted for ~50% of total OA throughout the day. SOA4 was dominant at night when it accounted for 39% of total OA, but it provided only a very small contribution to total OA during the day. SOA2 also contributed substantially to total OA, accounting for 21% of the total OA when it reached its highest concentration in the morning (8:00 - 12:00 PST). SOA1 always had a small contribution to the total OA, accounting for < 10% of total OA over all periods of the day. Both SOA2 and SOA3 substantially contributed to total OA (12–21% and 40–56%, respectively) during the day. Since SOA2 was more local and SOA3 was more regional, we suggest control of SOA precursor emissions on both local and regional scales are needed to reduce the daytime OA concentration. It will the most effective to control the regional precursors in the afternoon. SOA4, comprising both anthropogenic and biogenic SOA, was the largest OA source at night, but it's unclear if control of anthropogenic SOA precursors is able to reduce OA concentrations because the contribution of SOA formed by oxidation of anthropogenic precursors is less constrained.

33
3.7 Formation pathways of SOA

Controlling emissions of pollutants involved in the formation pathways of SOA has the potential to play a significant role in reducing regional OA concentrations (e.g. Liggio and Li, 2006; Na et al., 2007). In areas where biogenic emissions are oxidized in the presence of anthropogenic pollutants such as SO$_2$, NO$_x$ and black carbon, it has become increasingly apparent that SOA formation from biogenic VOC is substantially enhanced (Goldstein et al., 2009; Surratt et al., 2010; Carlton et al., 2010; Spracklen et al., 2011). Rollins et al. (2012) has shown that in the Bakersfield region reductions in NO$_x$ can reduce the OA concentration because of the involvement of NO$_x$ in formation of organic nitrates. Through the following discussion, we expect to not only suggest the main formation pathway of SOA formation in the Bakersfield site, but also examine if control of other species involved in SOA formation in addition to NOx and organic precursors can reduce OA concentrations. TAG-derived factors can be indicative of the dominant formation pathway for each SOA type identified. The pathways of oxygenated compounds contributing to SOA in each factor can be used to draw a distinction between direct gas-to-particle condensation wherein oxidation reactions produce low volatility products that condense, secondary gas-to-particle condensation wherein non-oxidation reactions produce lower volatility products that condense, such as reactions between carboxylic acids and ammonia, and reactive uptake wherein compounds expected to be in the gas-phase enter the particle phase through acid-catalyzed reactions. Evidence for these formation pathways in Bakersfield during CalNex based on individual compounds – specifically phthalic acid, pinonaldehyde and 6, 10, 14-trimethyl-2-pentadecanone has been explored and reported in Chapter 2. Ketones enter the particle phase primarily through direct gas-to-particle condensation, while phthalic acid was shown to enter the particle phase mainly through reaction with ammonia in the gas phase. It is likely that SOA1 and SOA3 were formed at least partially by direct gas-to-particle condensation because the dominant contribution to the factor profiles of these two factors are from compounds with ketone functional groups. The factor profile of SOA2 is primarily determined by particle-phase phthalic acid formed through reactions with gas-phase ammonia, indicating the reactions of carboxylic acids and ammonia play a significant role in the formation of SOA2. This formation pathway is also supported by the good correlation (r>0.5) between the concentration of SOA2 and the concentration of excess ammonium defined as the amount of ammonium which was not neutralized by sulfate and nitrate measured HR-ToF-AMS. This formation pathway could be site-specific because it needs excess ammonia to allow these reactions with carboxylic acids to occur. For SOA4 (i.e., the nighttime SOA) the factor profile contains pinonaldehyde, as well as phthalic acid and 6, 10, 14-2-pentadecanone. The presence of pinonaldehyde in the particles indicates the occurrence of acid-catalyzed reactions in the particles. However, the relative contribution to the nighttime OA from these particle-phase reactions cannot be distinguished from the direct gas-to-particle condensation. Chapter 2 has shown that secondary gas-to-particle partitioning and reactive uptake can significantly increase SOA yields of phthalic acid and pinonaldehyde relative to direct gas-to-particle partitioning. Therefore, control of species involved in these pathways of gas-to-particle partitioning such as ammonia and organic acids could lead to reductions in the OA concentration, in addition to control of organic precursors. However, the efficiency of reduction of SOA concentrations by control ammonia needs further investigation. For example, the reduction of ammonia would increase the aerosol acidity which improve the SOA formation from particle-phase reactions (Jang et al., 2002; Liggio and Li, 2006).
To substantiate that the formation pathways of SOA can be indicated by individual compounds, daytime SOA types derived from PMF analysis of organic species were compared to those from PMF analysis of bulk organic mass spectra measured by AMS (Figure 3.5). The contribution of the daytime SOA (the sum of SOA2 and SOA3) derived from organic species to total OA in this study displays a similar diurnal profile to SOA of the sum of high O/C alkane (O/C=0.68), high O/C aromatic (O/C=0.63) and petroleum (O/C=0.20) SOA derived from AMS mass spectra reported in Liu et al. (2012) from the same measurement campaign, which were selected because their mass fraction exhibited a daytime enhancement. The consistency of daytime SOA between these two studies supports that oxygenated organic compounds observed by TAG are able to reproduce the trend of SOA formation during the day.

3.8 Conclusions and atmospheric implications

PMF analysis was performed on 244 particle-phase speciated organic samples, acquired over the span of a month, to investigate OA sources. The variability of this dataset was best explained by six types of OA sources using PMF analysis. The concentration of reconstructed OA based on these six factors was in good agreement with the concentration of measured OA. Local POA accounted for 15% of the total OA on average and is suggested to be mostly contributed by motor vehicles. SOA was the dominant component of total OA throughout the day and night. SOA (the sum of SOA1-4) accounted for an average 72% of total OA with an average diurnal variation in the range from 66% at night to 78% during the day. SOA3 (regional SOA, 56%) was dominant during the afternoon and SOA4 (nighttime SOA, 39%) was dominant during the night. SOA2 (Local SOA) contributed significantly in the morning, accounting for 21% of total OA from 8:00 am to 12:00 pm PST.

Our results suggest that the formation of SOA during this study occurred dominantly through direct gas-to-particle condensation. Particle-phase reactions also contributed to formation of nighttime SOA, but their contributions are not well-constrained in this study. The formation of SOA in this study occurred through multiple pathways, some of which may be regionally-specific due to the unique mixture of emissions in this airshed. Our investigation of gas-to-particle partitioning suggested that the amount of SOA can be reproduced by the traditional SOA model using absorptive gas-to-particle partitioning.

The best control strategy for each type of SOA (SOA1-4) to enable effective reductions in the regional OA concentration may be different. The control of SOA precursor emissions on both local (SOA1 and 2) and regional (SOA3) scales are needed during the day, but control of regional SOA precursor emissions is likely to be most effective in the afternoon. Control of ammonia would also reduce the concentration of local SOA (SOA2). However, as discussed above, control of ammonia emissions could also lead to more nighttime SOA. At night, The contributions of both biogenic and anthropogenic precursors to the formation of nighttime SOA (SOA4) were evident, but neither of their contributions can be determined in this study. Consequently, it leaves it unclear if the concentration of nighttime SOA is effectively reduced by control of anthropogenic organic precursors.

3.9 References


Goldstein, A. H., C. D. Koven, C. L. Heald, and I. Y. Fung (2009), Biogenic carbon and anthropogenic pollutants combine to form a cooling haze over the southeastern United States, P. Natl. Acad. Sci. USA, 106(22), 8835-8840.


3.10 Tables and figures

Figure 3.1 PMF factor profiles. Compounds are generally grouped into hydrocarbons and oxygenated organic compounds. The y-axis is the fraction of each compound contributing to the total mass of all compounds in that factor.
Figure 3.2 Diurnal profiles of the mass concentration of each factor and the average diurnal profile of wind direction and the ratio of 1,3,5-TMB to toluene. In the box plots, the center line of each box is the median of the data, the top and bottom of the box are 75th and 25th percentiles and top and bottom whiskers are 90th and 10th percentiles. The solid square marker is the mean concentration. In average diurnal profiles, the mean value is plotted with ± one standard deviation.
Figure 3.3 Wind rose plots for six PMF factors using only concentrations larger than the mean concentration in each factor to emphasize the major contributing source directions. The wind direction is shown in degrees with 0 (360) as north, 90 as east, 180 as south and 270 as west. The relative concentration level is indicated by different colors: yellow > blue > green >red. The frequency of observations is represented by the length of each wedge.
Figure 3.4 Mean diurnal mass fraction contribution of each factor to total OA during six different times of day.
Figure 3.5 Average diurnal profiles of the mass fraction of OA identified as SOA by AMS PMF (Liu et al., 2012) and by TAG PMF (this study). The TAG data are shifted to the right for clarity.
Chapter 4
Development of an in situ thermal desorption gas chromatography instrument for quantifying atmospheric semi-volatile organic compounds

4.1 Abstract

Semi-volatile organic compounds (SVOCs) play a significant role in the formation of secondary organic aerosol, but their atmospheric abundance and chemical composition are poorly understood. We have developed a new system for the Thermal desorption Aerosol Gas chromatograph (TAG) that extends its capability to quantitatively speciate SVOCs (SV-TAG) with the vapor pressure lower than tetradecane (C_{14}) in hourly time resolution. The key component is a passivated stainless steel fiber filter that quantitatively collects both gas and particle phase organic compounds. A separation between gas and particle phase collection is determined through a difference method by periodically sampling ambient air through a multi-channel charcoal denuder that efficiently removes gas phase compounds. Measurements made with this new instrument will provide constraints on the abundance, chemical composition, and gas/particle partitioning of atmospheric SVOCs, and provide opportunities to improve our understanding of secondary organic aerosol formation in the atmosphere.

4.2 Introduction

Directly emitted, intermediate volatility organic compounds (IVOCs, defined as having an effective saturation concentration C* of $10^3$ to $10^6$ µg m$^{-3}$) and semi-volatile organic compounds (SVOCs, C* of $10^{-1}$ to $10^3$ µg m$^{-3}$) have been proposed as a substantial, yet unaccounted for source of secondary organic aerosol (SOA) precursors (Robinson et al., 2007; Weitkamp et al., 2007). Estimated atmospheric abundances of directly emitted I/SVOCs can be an order of magnitude larger than primary organic aerosol (Robinson et al., 2007). Chamber experiments show that oxidation of compounds in the IVOC range produces SOA efficiently (Lim and Ziemann, 2009; Chan et al., 2009; Presto et al., 2010; de Gouw et al., 2011). The importance of I/SVOCs to SOA formation is further supported by recent modeling work showing that the inclusion of estimated I/SVOCs emissions by means of the volatility basis-set approach (Robinson et al., 2007; Shrivastava et al., 2008; Grieshop et al., 2009), resulted in better agreement between measured and modeled SOA (Dzepina et al., 2009; Tsimpidi et al., 2010; Hodzic et al., 2010) than models that did not include I/SVOCs (e.g., Volkamer et al., 2006; Hodzic et al., 2010). However, these newer models cannot reproduce both the SOA mass and the oxygen to carbon ratio of the bulk aerosol (Hodzic et al., 2010), which highlights substantial deficiencies in the understanding of I/SVOCs emissions and oxidation mechanisms.

The volatility basis-set approach used in these newer models estimate primary I/SVOCs by multiplying concentrations of primary organic aerosol (POA) by a scaling factor derived from dilution measurements of emissions from two primary sources: diesel exhaust and wood burning (Robinson et al., 2007; Grieshop et al., 2009). The corresponding uncertainty in estimated IVOC abundance could be as much as three times the magnitude of POA emissions (Robinson et al., 2007). Gas-phase oxidation of I/SVOCs proposed by Robinson et al. (2007) and updated by Grieshop et al. (2009) lumps all species with the same effective saturation concentration into a single volatility bin and assumes a constant mass increase for species in a given bin through each
oxidation step. However, it is likely that a large number of isomers are present for each carbon number in the vapor pressure range of I/SVOCs (Goldstein and Galbally, 2007; Isaacman et al., 2012) and laboratory experiments have shown that SOA yields from gas phase OH radical initiated oxidation of alkane isomers differ according to their molecular structure in the order cyclic > linear > branched (Lim and Ziemann, 2009). In addition to absorptive gas-to-particle partitioning, previous studies have shown that other formation pathways, such as reactive uptake of small carbonyls on acidic particles (e.g. Jang et al., 2002; Tolocka et al., 2004) and reactions between carboxylic acids and ammonia (Na et al., 2007), are important for SOA formation. Presto et al. (2012) recently introduces a chromatography-based approach to determine the volatility distribution of POA emissions without speciation analysis based on relationship between the retention time of known compounds and saturation concentration of them. The observational constraints on the volatility distribution of S/IVOCs can be provided by using this approach, but S/IVOCs need to be quantitatively collected. To better predict SOA production from the oxidation of I/SVOCs, it is critical to apply observational constraints to emissions, speciation, and gas-phase oxidation mechanisms and pathways of gas-to-particle partitioning of I/SVOCs.

The Thermal desorption Aerosol Gas chromatograph (TAG, Williams et al., 2006) has shown the capability to capture hourly trends in the gas/particle partitioning of speciated organics and the variability of low-volatility compound vapors (Williams et al., 2010). However, the original TAG, with its impactor-based collector, was not designed for capture of vapor phase organics. As a result, this instrument could not quantitatively measure speciated I/SVOCs. As mentioned above that multiple SOA formation pathways are present in the atmosphere, oxygenated compounds in the vapor pressure range of IVOCs can therefore partition into particles through non-absorptive gas-to-particle partitioning (e.g. Jang et al., 2002). As a result, there is no clear split between IVOCs and SVOCs in the atmosphere according to gas/particle partitioning. We will take "SVOCs" in this paper to include both IVOCs and SVOCs described above. There are other operational definitions of SVOCs based on the vapor pressure and air-sampling strategy (Turpin et al., 2000). However, SVOCs are generally in vapor pressure range approximately equivalent to C_{12}~C_{32} n-alkanes (Turpin et al., 2000; Sihabut et al., 2005; Robinson et al., 2007; de Gouw et al., 2011). In this paper, we present the development and evaluation of a new configuration for TAG, called semi-volatile TAG (SV-TAG), that enables the quantitative measurement of speciated SVOCs with hourly resolution.

In selecting the sampling methodology for SV-TAG, consideration was given to denuder-based methods, wherein a sorbent tubular system is used to remove vapor phase organics prior to particle collection, and to filter-based methods that consist of a filter and sorbent trap in series. Filter-sorbent trap methods have been used extensively (Fraser et al., 1997; Simcik et al., 1998; Schauer et al., 1999; 2002; Subramanian et al., 2004), but are subject to positive artifacts caused by adsorption of gas-phase organics on a filter as well as negative artifacts due to evaporation of particulate organics. While sorption onto the filter media may be assessed experimentally (Turpin et al. 2000), it is not possible to determine the magnitude of volatilization from the filter, nor the absorption onto already-deposited organic material. The latter could be an even more important pathway contributing to underestimation of gas-phase organics (Storey et al., 1995). Denuder-based methods, with laboratory extraction and analysis of the substrates, have been used to measure SVOCs in the atmosphere (e.g. Gundel et al., 1995; Cui et al., 1998; Possanzini
et al., 2004) and from primary emission sources (e.g. Schauer et al., 1999; 2002). Additionally, thermally extractable, multi-capillary denuders have been developed using short sections of gas chromatography capillary columns (Krieger and Hites, 1992; Tobias et al., 2007) and stainless steel honeycombs coated by cross-linked and bonded polydimethylsiloxane (Rowe and Perlinger, 2010). If the denuder efficiently removes and retains vapor compounds of interest, the artifacts are smaller because gas-phase organics are collected by the denuder surface while particles pass through and are subsequently captured by the filter or the adsorbent bed downstream (e.g. Gundel et al., 1995; Eatough et al., 1993; Cui et al., 1998; Turpin et al., 2000). Concerns have been raised concerning the disequilibrium of gas/particle partitioning following the removal of gas-phase organics, but this is minimized if the residence time inside the denuder is short (Kamens and Coe, 1997). Moreover, vapor collection efficiencies and particle losses within the denuder can be determined experimentally (Eatough et al., 1993; Gundel et al., 1995). For these reasons we selected a denuder-based method for SV-TAG.

Our SV-TAG instrument combines denuder-difference sampling with in-situ thermal desorption and gas chromatography/mass spectrometry (GC/MS) analysis. The collection cell is designed to quantitatively collect both organic vapors and particles. The gas/particle partitioning is determined by sampling alternately with, and without the denuder in-line upstream. Although thermally extractable denuders have been discussed in the literature (Tobias et al., 2007; Rowe and Perlinger, 2010), we opted for the difference approach because it is a less complex system that allows us to combine the gas and particle collection and thermal desorption parts into a single component. This paper describes the evaluation of the SV-TAG denuder and collector components and procedures, and presents example ambient data.

4.3 Methods
4.3.1 The SV-TAG system
A schematic of the SV-TAG is shown in Figure 4.1. Components of SV-TAG that differ from the original TAG are (1) an activated carbon denuder (2) a metal fiber filter collection and thermal desorption cell (F-CTD), and (3) a valve-less injection interface (VLI) with a secondary focusing trap (FT). The denuder is switched in and out of the sample line, upstream of the F-CTD, and removes organic vapors while allowing particles to penetrate. The F-CTD enables efficient collection of SVOCs from the gas phase while simultaneously enabling efficient collection of all particles, all on a filter surface that can be directly thermally desorbed and then reused for the next sample. The FT facilitates the use of a much higher desorption flow rate (up to 200 ml min⁻¹) than is possible with the direct gas chromatography column transfer where injection flow rates are typically limited to ~1-2 ml min⁻¹.

The SV-TAG sample inlet uses a sharp cut PM₂.₅ cyclone (BGI Inc., Waltham, MA), followed by two parallel sampling lines (a denuder line and a bypass line) and a 3-way ball valve (V1, Figure 4.1). The denuder is a multi-channel, activated carbon cylindrical monolith made from pyrolized phenolic extrusions (3 cm OD, 20 cm length, ~ 490 channels, Mast carbon, UK). It is housed inside an aluminum tube and sealed by Viton o-rings with tapered caps at each end connecting to ~1.0 cm (3/8 inch) OD sampling lines upstream and downstream. The bypass sampling line is made of ~1.0 cm OD stainless steel tubing.
The F-CTD uses a 37 mm in diameter Bekipor® stainless steel (SS) fiber filter (3AL3, Bekaert Fiber Technology, Belgium). This filter has a total fiber surface area of ~160 cm$^2$ per cm$^2$ of filter surface based on manufacturer specifications. This is higher than the total fiber surface area of ~130 cm$^2$ per cm$^2$ of filter surface for quartz fiber filters (Mader and Pankow, 2001). The goal of this study is to develop an in-situ instrument capable of collecting SVOC samples. We hypothesized that a passivated SS fiber filter should be quantitatively capable of collecting SVOCs if it had large enough surface area, and it would be suitable for repeated thermal desorption of the sample for analysis. This hypothesis was based on previous observations that breakthrough of SVOCs on quartz filters is small in a 4 or 6-hour sampling duration (Subramanian et al., 2004) and the demonstration in our original TAG that passivated SS is a suitable material for thermal desorption of organics (Williams et al., 2006; Kreisberg et al., 2009).

The F-CTD is fabricated by permanently sealing this filter inside a 316 SS, brazed capsule with integrated sampling inlet and outlet tubes and a small bore tube for transferring desorbed organics to the GC (see Appendix C, Figure C1). Ridges located inside the capsule clamp the filter in place and ensure rapid heat transfer to the entire filter surface during thermal desorption. Following fabrication, the F-CTD is passivated with an Inertium® coating (Advanced Materials Components Express, Lemont, PA). The F-CTD is housed between two aluminum disks outfitted with four cartridge heaters and two temperature sensing thermocouples. Cooling fins are machined into the edges of the aluminum disks. A time programmed blower delivers room air to cool the F-CTD to room temperature after completion of thermal desorption.

The VLI and FT allow the collected sample to be transferred from the F-CTD, focused, and injected into the GC for analysis (Figure 4.1B) without having to pass through a valve. As described by Kreisberg et al (2012), the VLI provides more reproducible sample transfer than did the 6-port valve of the original TAG.

The VLI used in this study consists of two 50 cm long stainless steel capillaries with inner diameters of 125 µm connected by a pair of VICI tees (Valco Instruments Co. Inc.) housed in a square aluminum heating block (Figure 4.1) held at 300 ºC. Similar to the F-CTD, the capillaries and VICI tees are passivated with an Inertium® treatment.

The FT for the final configuration is made of a 50-cm section of a SS GC capillary column (MXT-5, 0.5 m x 0.53 mm ID, 5 µm film thickness, Restek) and integrated into the vent line of the VLI as shown in Figure 4.1. The FT has its own aluminum heating and cooling block with heat sink fins on both vertical edges (Figure C2). A thin layer of insulation between the VLI and the FT housing minimizes heat transfer. A time programmed box fan is used to maintain the FT at a low enough temperature to re-focus organics desorbed from the F-CTD. This FT does not require liquid nitrogen or other consumables.

4.3.2 SV-TAG operation

SV-TAG alternates between two modes of operation: (1) GC sample analysis with simultaneous collection of the next sample and (2) thermal desorption of a sample into the GC. After collection, the deposited organics are thermally desorbed from the F-CTD and refocused onto the FT ("sample trapping") followed by thermal transfer from the FT to the GC ("sample
injection"). Following sample injection, the F-CTD is cooled in preparation for collection of the next sample. Sample collection alternates between the bypass (undenuded) and denuder inlet lines. Gas/particle partitioning is determined by difference, with the undenuded sample providing "total organics" and the denuded sample providing "particle-phase organics".

For sampling, ambient air at 10 L/min is pulled from the center of a larger transport flow air stream, through the cyclone, through either the denuder or bypass line, and then delivered to the F-CTD which is maintained at 28 °C. After completion of ambient sampling, the 3-way valve (V1) and sample valve (V2) are closed and helium flow is set to 200 ml/min for one minute with the cell still at 28°C to flush any residual air from the F-CTD. The flow is then reduced to 10 ml/min and organics are thermally desorbed by heating the F-CTD from 28 °C to 305 °C at a rate of ~35 °C/min, then to 315 °C at a rate of 5 °C/min and holding at 315 °C for 4 minutes. Midway through the desorption, the helium flow is increased from 10 to around 150 ml/min to facilitate the transfer of the less volatile material. The majority (~80%) of the helium desorption flow enters the F-CTD through its outlet and the remainder through the inlet such that the filter is predominantly back-flushed. Desorbed organics are refocused onto the FT which is held at 40 °C.

After sample injection, the system is switched back to the GC analysis mode for organic chemical analysis (see Appendix C2). The F-CTD and FT heaters are turned off and fans are turned on to cool the F-CTD and FT. Typically, sample collection time is 30 min (or 90 min for 2-hr cycle time), desorption and transfer step is about 20 min, followed by an 8-min cooling step. The GC analysis is typically 35 min and is initiated at the conclusion of the desorption, and continues during the collection of the next sample.

4.3.3 System evaluation

The combined thermal desorption and transfer efficiency of organics from the F-CTD to the head of the GC column were evaluated using a C₈-C₄₀ n-alkane liquid standard, which covers the SVOC vapor pressure range. The liquid standard was introduced into the F-CTD via a micro-liter syringe through a customized injection port and thermally desorbed and transferred to the GC/MS for analysis. The thermal desorption flow was varied for these tests. Initially transfer efficiencies were evaluated without a focusing trap, which restricted the thermal desorption flow to the GC column flow of 1-2 ml/min. Subsequent experiments, reported here, employed a FT which decouples the thermal desorption flow from the chromatographic flow and facilitated evaluation of the transfer efficiencies at higher carrier gas flows, as much as 200 ml/min.

After the addition of a FT into the system, the combined thermal desorption and transfer efficiency from the F-CTD to the head of the column, defined as the recovery, depends on the thermal desorption efficiency of particle-phase organics from the F-CTD, the efficiency of the FT to trap the most volatile compounds in the vapor pressure range of SVOCs, and the transfer
from the FT to the column. Initially, the thermal desorption from the F-CTD was optimized by examining the recovery of the particle-phase alkanes, followed by optimizing the recovery of SVOCs from the FT. Our evaluation compared the relative recovery of individual alkanes from a standard comprised of equal alkane masses at each carbon number. The relative recovery of each n-alkane was expressed as the ratio of abundance of this n-alkane to the maximum abundance of n-alkanes in the C_{14}-C_{40} range for each run.

The selection of the FT materials is based on the goal to minimize any extra losses caused by the addition of the FT. Since all of the components exposed to organics after thermal desorption were made of SS and the GC column, three different media were evaluated: unpassivated SS tubing and two SS MXT-5 columns with 3µm and 5µm film thickness, respectively. The FT made of the same length of each material was initially evaluated to determine which one would be optimized for the recovery of SVOCs by increasing the length of the FT and reducing the thermal desorption flow rate. For all experiments, the temperature of the FT was held at 40°C during sample trapping because of its proximity to the heated VLI.

The collection efficiency (CE) and volatilization losses of the F-CTD were evaluated using authentic standards (Table B1) under simulated ambient sampling conditions. The authentic standards covered the vapor pressure range of SVOCs and spanned several different functional groups, typically observed in the atmosphere. These standards were introduced using an impactor collection and thermal desorption cell (I-CTD) placed immediately upstream of the F-CTD to act as a vaporizing source. The I-CTD was used in the original TAG, and is a compact collection cell equipped with an injection port for liquid standards (Kreisberg et al, 2009).

To evaluate volatilization losses, we utilized a "thermal desorption transfer" method whereby liquid standards introduced into the I-CTD with a micro-syringe were transferred to the F-CTD by vaporizing them at 300 °C in a 150 ml/min helium flow. This low flow rate ensured efficient capture by the F-CTD. Then, particle and hydrocarbon free air from a zero air generator (model 737, AADCO, Cleves, OH) was drawn at 10 L/min though the F-CTD for either 30 or 90 minutes, to simulate normal ambient sampling (either 30 or 90 minutes). Profiles of the flows, F-CTD and FT temperatures are shown in Figure C3.

To evaluate the overall CE of the F-CTD, including both CE and volatilization losses, we utilized an "evaporation transfer" method whereby organic standards deposited in the I-CTD were transferred to the F-CTD over either 30 min or 90 min period in a 10 L/min zero air flow at room temperature. This simulated organic vapor collection during real ambient sampling. The material on the F-CTD was analyzed at the end of the room temperature transfer. Then the I-CTD was heated in a 150 ml/min He flow to transfer the remaining standard and the F-CTD was analyzed a second time. These data provide the total retention by the F-CTD by compounds. This retention is the sum of collected material. Details are shown in Figure C4. The detailed description of the thermal desorption method and evaporation transfer method are also provided in Appendix C3.

The gas CE of the denuder was determined by examining the difference between the undenuded and denuded samples in Berkeley, CA when a Teflon filter was placed upstream of the cyclone to remove particles. The size-dependent penetration of particles through the denuder...
and the particle CE by the F-CTD were evaluated through measurements of upstream and downstream particle size distributions as presented in Appendix C4.

4.3.4 Ambient sampling

The SV-TAG was initially tested in Berkeley, CA in an hourly automated in-situ mode of operation for two periods. In the first period of June 13 - 20, 2011, both denuded and undenuded samples were collected to investigate gas/particle partitioning and overall denuder performance. In the second period of Oct. 1- 4, 2011, only undenuded samples were collected to investigate the variability of SVOCs in the atmosphere. Additionally, in the second period deuterated internal standards were co-injected into the F-CTD with every ambient sample to evaluate the reproducibility of the thermal transfer efficiency. For all ambient runs SV-TAG was operated with the 50 cm, 5 µm film FT with the two step purge. The exact temperature and flow settings are presented in Figure C5. The denuder performance was also evaluated as part of the California Research at the Nexus of Air Quality and Climate Change campaign in Bakersfield during June-July 2010.

4.4 Evaluation results

4.4.1 Thermal desorption and transfer efficiency

Tests of the thermal desorption from the F-CTD were first conducted using a 15 cm long section of unpassivated SS tube as the FT. Figure 4.2 shows the recovery of n-pentadecane (C₁₅H₃₂, a gas-phase alkane) and n-tritriacontane (C₃₃H₆₈, a particle-phase alkane) as a function of the thermal desorption flow rate (recoveries of the full range of n-alkanes are shown in Figure C6). The recovery of n-tritriacontane was significantly enhanced by increasing the thermal desorption flow rate up to 150 ml/min. The recovery of n-pentadecane decreased with increasing the desorption flow rate, which was attributed to its breakthrough from the unpassivated SS FT.

The dependence of the recovery on the desorption flow rate was also evaluated with two other types of FTs made of 15 cm long GC columns (MXT-5 0.53mm OD, 3 and 5 µm film thickness). While the recovery of \( > C_{20}\) n-alkanes was observed to be similar for all three types of FTs under the same thermal desorption method; the recovery of n-alkanes with carbon number \( \leq 20\) from the FTs made of the SS GC columns was significantly different from the FT made of the unpassivated SS tube (Figure 4.3a). The relationship between the recovery and carbon number from the FTs made of GC columns exhibited a steeper drop, relative to the unpassivated SS tube in the lower end of carbon numbers. The steeper drop reduces the number of compounds whose recoveries would be estimated by interpolating the recovery curve of n-alkanes. Additionally, the recovery of untested compounds from the FT made of a section of a GC column can be derived using their retention time determined through the GC analysis and the recovery curve constructed using the recovery and retention time of n-alkanes. The thicker film (5 µm) FT improved the recovery of more volatile n-alkanes, relative to the FT made of the GC column with 3 µm film thickness, due to the increased capacity of the film to trap organics.

The recovery of \( < C_{18}\) n-alkanes from the FT was optimized using the FT made of a section of the GC column with the 5 µm film thickness. The optimal recovery for SVOCs was achieved when the length of the FT was increased to 50 cm and the total volume of desorption flow was reduced by shortening the total desorption time from 18 to 14 minutes and by using a
two step purge with a flow of 10 ml/min for 6 minutes followed by a higher flow of 150 ml/min for 8 minutes. With the two step purge and 50 cm long FT, the recovery of n-tetradecane (C_{14}H_{30}) from the FT was about 40%, while the recovery of C_{15}-C_{26} alkanes was above 95% (Figure 4.3b) and C_{27}-C_{32} was 50-90% (Figure C6). Assuming a Gaussian elution profile, we estimate that recovering close to 100% of n-tetradecane would require doubling the length of the FT, improvements for more volatile components would best be achieved through redesign of the FT housing to facilitate active cooling.

4.4.2 SVOC collection by the F-CTD

The upper limit for losses of collected organics due to volatilization was evaluated using the thermal desorption transfer method. The average recovery (n = 5) of most compounds was around 100% after a 30 min zero air purge (Figure 4.4a). The recovery of each compound with a 90 min purge was approximately the same as with a 30 min purge and the average recovery of all compounds after 30 min and 90 min purges was 0.99 ± 0.02 and 0.99 ± 0.05, respectively. Except for the concentrations listed in Table B1, volatilization of collected n-alkanes from the F-CTD was evaluated at concentrations of both 16.7 ng/m$^3$ and 66.7 ng/m$^3$ for 30 min ambient sampling. This range of concentrations was chosen to mimic the ambient concentration range of n-pentadecane (0 to 55 ng/m$^3$) reported by Fraser et al. (1997). The fraction of each compound observed to be retained on the F-CTD was in the range of 0.9 to 1.05. Since organic compounds with lower vapor pressures show stronger adsorption constants on the surface of sampling substrates (Goss and Schwarzenbach, 1998), we conclude that losses due to volatilization from the surface of the F-CTD are negligible for organics with vapor pressures lower than n-tetradecane under these sampling conditions.

The overall CE of the F-CTD for gas-phase organics was directly evaluated using the evaporation transfer method. All compounds had CE in the range of 0.8 to 1.0 for 30 min simulated ambient sampling, with the exceptions of 1-tridecanal and 2-pentadecanone (Figure 4.4b). A similar overall CE was also observed for 90 min simulated ambient sampling and the average ratio of overall CE for all compounds between 30 min and 90 min simulation was 1.00 ± 0.05 (Figure 4.4b). The similar overall CE for all other compounds observed in both 30 min and 90 min simulation demonstrates that the F-CTD is capable of quantitative measurements of SVOCs with vapor pressures lower than n-tetradecane for 90 min sampling at 10 L/min.

It's unclear what caused the lower measured overall CE for 1-tridecanal and 2-pentadecanone. The diffusivity of 1-tridecanal and 2-pentadecanone is larger than most other compounds, such as n-heptadecane, yet the recovery of n-heptadecane is close to 100% in the same evaluation. Additionally, we have demonstrated that the losses of collected organics due to volatilization are negligible. However, the measured overall CE was reproducible allowing for an empirical correction.

4.4.3 Particle collection efficiency by the F-CTD

Particle collection efficiencies for the F-CTD are presented in Appendix C (Figures C7 and C8). Mobility based measurements with test aerosols data show collection of 94% - 100%, with a minimum at 100 nm diameter for sampling at 10 L/min (Figure C7). Optical counter data with ambient particles show a similar profile, but with a minimum CE of 90% at 100 nm (Figure
C8). On a mass basis, the average mass CE for PM\textsubscript{1.0} ambient particles was 98% as estimated from the upstream and downstream particle volume distribution. In contrast to the particle CE of the original I-CTD which had a 50% cut-point at 0.1 µm, the F-CTD improved the CE of ultrafine particles to over 90%.

### 4.4.4 Denuder vapor collection and particle penetration

For atmospheric measurements, the concentration of gas-phase organics measured downstream of a Teflon filter followed by the denuder was less than 1% of that measured downstream of the Teflon filter alone, indicating a CE of over 99%. Moreover, these measurements were observed to be stable for more than a month of continuous ambient sampling during the California Research at the Nexus of Air Quality and Climate Change campaign in Bakersfield during June-July 2010 (See Appendix A, Figure A1). Average losses of particles inside the denuder were less than 10% by number for particle sizes spanning the range 0.05~1.00 µm (See Appendix A, Figure A2).

### 4.4.5 Measurements of ambient air

The stability of SV-TAG for ambient run of 3 days was evaluated by examining the consistency of internal standards (ISs) injected with each sample. As shown in Figure 4.5, the variability of an IS around its mean value was within 10% for the more volatile IS (hexadecane-d\textsubscript{34}, eicosane-d\textsubscript{42}, and phenanthrene-d\textsubscript{8}). The range of variability for later eluting internal standards, such as n-tetracosane-d\textsubscript{50}, was greater and was found to be strongly related to changes in the total organics collected by the F-CTD (organic loading) (Figure C9). The effect of the organic loading on the recovery of individual compounds has been reported in measurements of ambient organic aerosol with a previous version of the TAG (Lambe et al., 2010) and an independent study of thermal desorption of organics from charcoal (Senf and Frank, 1990). The observed relationship between the variability of each IS and the organic loading establishes the need to use ISs introduced on the top of ambient samples to correct for the effect of varying organic loadings. It should be noted that the effect of the organic loading could be overestimated in the current system because ISs were introduced into the F-CTD from its downstream side and thus increasing the surface area to which internal standards were exposed, relative to the ambient samples collected on the upstream surface. The injection of internal standards will be changed in future designs to better mimic ambient sampling.

A pair of undenuded and denuded samples are shown in Figure 4.6a. Over one hundred compounds were identified in the undenuded sample using authentic standards and according to the best matches available with the NIST2008 mass spectral database. The identified compounds span a broad vapor pressure range and many different functional groups, such as alkanes, acids and aromatics. For example, the observed n-alkanes demonstrate the wide vapor pressure range from SVOCs to particle-phase organics that are possible to detect and quantify with this SV-TAG instrument and they also highlight the capability to quantify gas to particle partitioning of organic compounds (Figure 4.6b). Though the undenuded and denuded samples were not collected simultaneously, the previous version of the TAG has demonstrated that this sampling strategy is able to capture the variability of gas/particle partitioning in a long timeline of observations (Zhao et al., 2012).
Some of the identified compounds, which are predominantly present in the gas phase, can be used as organic markers to provide observational constraints on the abundance of primary emissions of gas-phase SVOCs. Pristane and phytane are potential tracers for the emissions of SVOCs from diesel and gasoline powered vehicles (Fraser et al., 1997; Schauer et al., 1999; 2002). Other identified organic compounds, such as phthalic acid, 6, 10, 14-trimethyl-2-pentadecanone, alkyl cyclohexanes and dibenzothiophene that are frequently used as source markers for organic aerosol in source apportionment models are traditionally considered to be only present in particles (e.g. Shrivastava et al., 2007). However, our observations of gas/particle partitioning of these compounds clearly show that they are present in both gas and particle phases as SVOCs. With the SV-TAG, we can clearly separate and quantify both the gas phase and particle phase abundance of these compounds, and thus investigate their partitioning and use them with less uncertainty for source apportionment with hourly resolution.

The measured abundance of gas phase organics was about an order of magnitude larger than the particle-phase organics in Berkeley CA over the SVOC range observed by the SV-TAG. The gas-phase organics, determined by the difference between the total (undenuded) and particle-phase (denuded) organics, were dominant early in the chromatogram and mostly formed an unresolved complex mixture (Figure 4.6a). The sum of measured n-alkanes (C_{14}-C_{20}), which were dominantly present in the gas phase, accounted for ~7% of total measured organics in the same retention time range as n-alkanes (C_{14}-C_{20}), defined as high-volatility SVOCs which were dominantly present in the gas phase and calculated by integrating the total ion in the retention time range of n-alkanes (C_{14}-C_{20}). The sum of measured of n-alkylcyclohexanes (C_{14}-C_{20}) accounted for less than 1% of high-volatility SVOCs. The upper limit of the contribution from straight-chain and branched alkanes to high-volatility SVOCs was estimated to be ~32% by assuming that m/z 57 (C_4H_9^+) was just contributed by straight-chain and branched alkanes. This restricts the upper limit for branched alkanes to be ~ 25% of measured high-volatility SVOCs and indicates that branched hydrocarbons likely made up a substantial fraction of organics in the ambient air in Berkeley, CA.

### 4.5 Summary and atmospheric implications

In this chapter, we demonstrated that the SV-TAG is capable of hourly automated quantitation of speciated gas- and particle-phase organics with their vapor pressure lower than tetradecane (C_{14}). The recovery of n-alkanes relative to eicosane (C_{20}) is 40% for C_{14}, 95-100% for C_{15}-C_{26} and 50-90% for C_{27}-C_{32}. The recovery of individual compounds during the ambient measurements can be tracked and corrected using ISs introduced into each sample. The F-CTD cell efficiently collects the entire particle size range of PM_{2.5} with collection efficiency near 100%, except for a slight dip to ~90% in a narrow range around 0.1 μm. The SV-TAG facilities quantitative measurements of the gas to particle partitioning of SVOC compounds through a denuder difference method. We also showed that traditional molecular markers for organic aerosol can be routinely measured by the SV-TAG with minimal sampling artifacts. We observed that many traditional organic aerosol markers are actually SVOCs with a significant fraction in the gas phase, which has important implications for the use of these organic tracers in source apportionment models. The SV-TAG is now ready for field deployments and will provide unique datasets for the study of emissions, transformation, and partitioning of SVOCs and for
studies of SOA formation and transformation in the atmosphere over hourly to seasonal timescales.

4.6 References


4.7 Tables and figures

(a) Schematic of the SV-TAG system with the major components labeled. (b) Schematic of the F-CTD, FT and valveless injector showing the flow direction, valve states and temperature states in each mode of operation.

Figure 4.1 a) Schematic of the SV-TAG system with the major components labeled. (b) Schematic of the F-CTD, FT and valveless injector showing the flow direction, valve states and temperature states in each mode of operation.
Figure 4.2 Recovery of representative compounds as a function of flow rate through the F-CTD and an unpassivated SS FT.
Figure 4.3 Recoveries of n-alkanes as a function of carbon number (≤ 26) measured with different FTs. a) Selection of FT materials; b) Optimization of FT length and purge method. For the one-step purge, thermal desorption flow was set to 150 ml/min for 18 min. For the two-step purge, thermal desorption flow rate was set to 10 ml/min for 6 min then 150 ml/min for 8 min.
Figure 4.4 Overall gas collection efficiency of the F-CTD measured by (a) thermal desorption transfer and (b) evaporation transfer. The compounds are generally grouped by alkanes, aromatic hydrocarbons and oxygenated compounds and their chemical formulae are listed in Table B1.
Figure 4.5 Stability of the SV-TAG collection and detection system over a 3 day period of continuous ambient sampling as indicated by the response of 6 per-deuterated internal standards that were co-injected with each ambient sample.
Figure 4.6 (a) Total ion chromatograms of undenuded (black) and denuded (green) ambient samples collected in Berkeley, CA. The sampling volumes were 0.5 m$^3$ each, collected at 10 L/min for 50 minutes, and the denuded sample was collected 40 minutes after the undenuded sample; two large peaks in the chromatogram of the denuder sample were peaks of column bleed. (b) Measured concentrations of n-alkanes (left axis) and particle phase fraction (right axis) calculated by dividing the particle-phase concentration (denuded sample) by the total concentration (undenuded sample).
Chapter 5
Conclusions

5.1 Summary

In this thesis, a novel instrument (SV-TAG) has been successfully developed to quantitatively measure SVOCs and OA in the atmosphere. The design was based on the principles of thermal desorption followed by GC/MS analysis (TAG). Ambient measurements of speciated OA and SVOCs were made with this SV-TAG at its two development stages: 1) improved separation of gas and particle phase organics and 2) quantitative collection and transfer of SVOCs to the GC/MS. Insights into the various source contributions to OA and SOA formation were gained through ambient measurements made in Bakersfield, CA during CalNex campaign following the first development stage. The capability of the SV-TAG to quantitatively measure speciated SVOCs was demonstrated by laboratory evaluation and ambient measurements in Berkeley, CA following the second development stage.

5.1.1 Gas-to-particle partitioning (SOA formation)

In the first stage of development of the SV-TAG, a denuder was added into the sampling inlet to improve separation of gas- and particle-phase organics. However, the original TAG impactor-based cell, designed for collection of particles but had reduced collection of organic vapors, was used. Consequently, the overall fraction of organic compounds in the particles was overestimated. After the improvement of addition of a denuder into its sampling inlet, the upper limit of the overestimation of fractions of organic compounds in particles was able to be estimated using observations of \( n \)-alkanes and consequently investigation of different pathways of gas-to-particle partitioning of oxygenated compounds was allowed to be done.

Measurements of gas/particle partitioning of phthalic acid, pinonaldehyde and 6, 10, 14-trimethyl-2-pentadecanone were selected to explore the pathways of gas-to-particle partitioning whereby SOA was formed. The results indicate that absorption into particles is the dominant pathway for 6, 10, 14-trimethyl-2-pentadecanone to contribute to SOA in the atmosphere. Absorption of gas-phase phthalic acid into particles can contribute to observed particle-phase phthalic acid concentrations, but the major pathway forming particle-phase phthalic acid is likely through reactions with gas-phase ammonia to form condensable salts. The observations of pinonaldehyde in the particle phase when inorganic acids were neutralized indicate that inorganic acids are not required for occurrence of reactive uptake of pinonaldehyde into particles. The laboratory observed effect of aerosol acidity on reactive uptake of pinonaldehyde observed in ambient particles. The relationship between particle-phase pinonaldehyde and RH suggests that aerosol water content likely plays a significant role in the formation of particle-phase pinonaldehyde. These results highlight the needs to investigate the effects of RH and organic acids on the reactive uptake of pinonaldehyde and other aldehydes onto neutral seed particles in chamber experiments.

Though some of the major pathways of gas-to-particle partitioning were successfully investigated without quantified gas-phase organics, an instrument capable of quantitative
measurements of gas/particle partitioning is desirable to allow direct comparisons of partitioning coefficients between ambient and laboratory measurements, to provide accurate parameterizations of gas/particle partitioning in SOA models, and to provide quantitative data on SVOC concentrations in the atmosphere.

5.1.2 Major source contributions to OA

In Chapter 2, many compounds, including traditionally SOA tracers, have been shown to be present in both gas and particle phases. With the use of a denuder in the sampling inlet (described in Chapter 2), temporal variability of these tracers purely in the particles can be directly quantified. In Chapter 3, the measured particle phase concentrations were used in PMF analysis to determine POA and SOA sources contributing the OA in Bakersfield, CA during 2010 CalNex.

Six OA sources were identified, including a POA source, a mixture of OA sources, and four types of SOA sources. Local vehicles were suggested to significantly contribute to POA. Four types of SOA (1-4) displayed distinct diurnal profiles. Each of SOA1-3 displayed an enhancement in its contributions to OA at different time during the day. SOA1 and 2 were mainly local while SOA3 was more regional, transported. SOA4 mainly occurred during the night and included contributions from both anthropogenic and biogenic SOA contributions.

SOA was the dominant component of OA, with four types of SOA accounting for an combined 72% of OA, varying diurnally from 66% at night to 78% during the day. Regional SOA (SOA3, 56%) was the largest contributor to OA during the afternoon and SOA4 (nighttime SOA, 39%) was the largest one during the night. A clear split between local and regional SOA during the day and anthropogenic and biogenic SOA during night cannot be made in this study. However, the results indicate that regional SOA had a larger contribution to OA than local SOA during the day and biogenic SOA contributed a significant fraction of OA during the night.

The best control strategy for each type of SOA (SOA1-4) to enable effective reductions in the regional OA concentration may be different. The control of SOA precursor emissions on both local and regional scales are needed during the day, but control of regional SOA precursor emissions is likely to be most effective in the afternoon. At night, our observations suggest that biogenic SOA could contribute a significant fraction of total OA, yet its contribution is not well constrained. Consequently, it leaves it unclear if control of anthropogenic SOA precursor emissions can efficiently reduce OA concentrations. In addition to organic precursors of SOA, other species involved in SOA formation can be controlled to reduce OA concentrations. The major pathway is indicated to be absorptive gas/particle partitioning, but other formation pathways, including reactions between organic acids and ammonia to form condensable salts and reactive uptake of carbonyls onto particles, also played a significant role in SOA formation. Therefore, reductions in emissions of the species involved in the SOA formation, such as gas-phase ammonia, should also reduce the SOA concentration in the atmosphere in this region. However, the efficiency to reduce OA by control of ammonia needs further investigation because reductions in ammonia emissions could lead to high aerosol acidity and subsequently increase nighttime SOA formation.

5.1.3 SV-TAG and ambient measurements
As described in Chapter 2, the original TAG with improved gas/particle separation is able to investigate gas/particle partitioning, but it is not capable of quantification of speciated SVOCs and the gas/particle partitioning coefficient constants which are critical to providing observational constraints on abundance and speciation of SVOCs and parameterize the gas/particle partitioning in models. In response to these needs, a new collection and thermal desorption system was developed, enabling the SV-TAG to quantitatively collect and transfer speciated SVOCs to the GC/MS.

The development and evaluation of the SV-TAG was presented in Chapter 4. The major components differing from the original TAG include: (1) an activated carbon denuder, (2) a metal-fiber filter collection and thermal desorption cell (F-CTD), and (3) a valve-less injection interface (VLI) with a secondary focusing trap (FT). These new components enabled the quantitative measurements of speciated SVOCs with vapor pressures lower than n-tetradecane (C_{14}) within a 90-min sampling period.

Ambient measurements in Berkeley CA showed that the abundance of gas-phase organics was an order of magnitude larger than the particle-phase organics over the SVOC vapor pressure range. Of these measured organics, the sum of measured n-alkanes (C_{14}-C_{20}) accounted for ~7% of measured high volatility SVOCs and the sum of measured n-alkylcyclohexanes (C_{14}-C_{20}) accounted for less than 1% of measured high volatility SVOCs. The upper limit of abundance for branched alkanes was estimated to be ~ 25% of measured high volatility SVOCs, indicating that branched hydrocarbons accounted for a substantial fraction of organics in the ambient air.

Tracers in the vapor pressure range of high volatility of SVOCs, such as phytane and pristane, were quantified in the ambient samples. Other identified organic compounds traditionally considered to be only present in particles were observed to be present in both gas and particle phases. With the SV-TAG, these tracers can be quantitatively measured and subsequently used in source apportionment models for both gas and particle phase organics.

In summary, this thesis has improved our understanding of SOA formation and source contributions to OA by developing a novel instrument capable of time-resolved measurements of gas- and particle-phase organic species and applying it to ambient measurements. This thesis work has provided direct evidence that multiple pathways of gas-to-particle partitioning of organic compounds occur in the atmosphere, are compound-dependent and are different from the standard absorptive/partitioning theory. This thesis also suggests effective control strategy to reduce OA concentrations. The development of SV-TAG enables not only investigation of the pathways of gas-to-particle partitioning with hourly time resolution, but also quantitatively measurements of speciated SVOCs. These measurement capabilities open a avenue to quantify the gas/particle partitioning in the atmosphere and provide observational constraints on the abundance of SVOCs, and enable investigation the primary emissions of SVOCs.

5.2 Future work

With the capability of the SV-TAG to quantitatively measure both gas- and particle-phase organics in an automated, continuous mode of operation, further work can be done, including 1) measurements of gas/particle partitioning; 2) source apportionment of SVOCs; 3) examination of
SOA formation and transformation using PMF in combination with SOA tracers, although further improvements in the instrumentation are still needed.

5.2.1 Measurements of gas/particle partitioning

SOA formation, including oxidation of gas-phase organics and the pathways of gas-to-particle partitioning of their oxidation products, has been mainly investigated in chamber experiments wherein the thermodynamics of gas/particle partitioning may be different from that in the atmosphere. Therefore, the unique capability of SV-TAG in measurements of gas/particle partitioning in the atmosphere is a powerful tool to examine our understanding of SOA formation in the atmosphere. Chapter 2 has shown that ambient measurements of gas/particle partitioning enables identification of other pathways of gas-to-partitioning and the investigation of parameters affecting gas/particle partitioning, but no quantitative constraints can be provided from these measurements due to incomplete collection of gas-phase organics.

With the configuration described in Chapter 4, quantitative measurements of gas/particle partitioning are able to be made to: 1) provide observational constraints on the mean molecular weight of atmospheric OA and activity coefficients of organic compounds in atmospheric OA whose gas-to-particle partitioning is through absorption; 2) identify the presence of other pathways of gas-phase organics entering particles other than absorption into particles and 3) subsequently provide observational constraints on the contributions of individual organic compounds to SOA and parameterization of factors affecting these pathways. Additionally, using the SV-TAG, the oxidation of SOA precursors can now be better mapped through measurements of gas/particle partitioning of their oxidation products and consequently provide constraints on oxidation mechanisms used in SOA models.

5.2.2 Measurements of SVOCs

The need for measurements of atmospheric SVOCs to provide observational constraints on speciation, abundance and partitioning of SVOCs was highlighted in the introduction of Chapter 4. The chemical composition and abundance of SVOCs vary spatially owing to difference in the composition of primary sources in different regions. Therefore, measurements covering more regions are desirable. In addition to measurements of SVOCs in the atmosphere, direct measurements of primary emission sources are needed for source apportionment of SVOCs.

5.2.3 SOA formation and transformation

In combination with investigation of gas/particle partitioning in Chapter 2, Chapter 3 has demonstrated that SOA formation and transport can be explored using PMF analysis of time-resolved organic species. In this thesis, a novel approach to investigate SOA formation in the atmosphere was presented providing insights into the discrepancies between modeled and measured SOA. This approach needs to be applied to more areas to improve our understanding of SOA formation and offer implications for improving air quality regulation.

5.2.4 Instrumentation

Gas/particle partitioning is currently determined by a denuder difference method wherein the denuder is alternated into the sampling line every other run due to a single collection cell employed in the configuration. Though the current configuration makes the collection and
thermal desorption system simple and is able to provide useful insights into SOA formation and gas/particle partitioning, this sampling strategy could be improved to provide simultaneous observations of gas and particle phases to examine in more detail how partitioning varies with meteorological conditions, atmospheric chemistry and OA matrix. One future alternative which can be tried is a dual-cell setup, in which one cell is used to collect particle-phase organics and another is used to simultaneously collect total organics (gas-phase and particle-phase organics), with sequential desorption and analysis through the GC/MS.

Chapter 4 has shown that current operating temperature of the FT is a limiting factor which could be reduced to improve its recovery for small-molecular-weight compounds. Two main factors affecting the temperature of the FT when it is trapping the organics desorbed from the F-CTD are the temperature of the cooling air (room temperature) and the heat transfer from the VL injector. The large thermal mass of the current heating block of the VL injector limits the time required to ramp the temperature of the VL injector during the SV-TAG operation. If the current configuration of the VL injector and FT is used, active cooling should be applied to the FT to increase the higher volatility range of measureable species. Future work also should include reducing the thermal masses of heating blocks of the VL injector and the contact surface between the heating blocks for the VL and the FT.

Many potentially important SOA tracers, such as dicarboxylic acids, are typically too polar to completely elute from the GC column. This severely limits the capability of SV-TAG to quantify potentially dominant SOA tracers. The transmission efficiency of these polar compounds through the GC can be increased by derivatization, such as methylation and silylation. Future work should therefore include development of an on-line derivatization method to improve the capability of SV-TAG to quantify more polar SOA tracers.
Figure A1 The stability of gas collection efficiency of the denuder throughout the 1-month field campaign, displayed by its collection efficiency for n-pentadecane.
Figure A2 Particle losses inside the denuder.
Appendix B: Supplemental information for Chapter 3

Appendix B1 Effect of inclusion of gas-phase contributions to particle-phase SVOCs on the PMF solution

Many compounds present in the particle phase, especially SOA tracers, are SVOCs. As a result of the utility of SVOCs in PMF analysis, inclusion of gas-phase contributions can bias the temporal variability in particle-phase concentrations of these SVOCs and subsequently lead to changes in correlations between different compounds which PMF analysis relies on to extract different factors and explain the OA observations. In our study, the effect of gas-phase contributions on the PMF solution, caused by gas-phase organic adsorption to the collection cell when sampling in bypass mode (no denuder), is investigated by comparing the same number of factors extracted from the same group of compounds with or without inclusion of gas-phase contributions. The factor profiles from measurements of organics with and without inclusion of gas-phase contributions are significantly different (Figure B2). To visualize the difference caused by inclusion of gas-phase contributions, the correlation between two factor profiles and the correlation between their time series were plotted against each other. As shown in Figure B3, the difference in the temporal variability and chemical composition of the factors are apparent for factor 1, 4, 5. Though the correlation coefficients (r) for both temporal variability and chemical composition of the factor 2, 3, 6 are good, they are still less than 0.8. This comparison establishes that the exclusion of gas-phase contributions to SVOCs used in PMF analysis is crucial to achieving correct source apportionment.
Figure B1 (a) The box plot of the relative residues determined using the difference between reconstructed and measured OA divided by measured OA. The center line of each box is the median of the data, the top and bottom of the box are 75th and 25th percentiles and top and bottom whiskers are 90th and 10th percentiles. The data points are evenly distributed to each interval. (b) the frequency of the atmospheric OA concentrations in each interval observed at night.
**Figure B2** Factor profiles extracted by PMF analysis from the same group of compounds with and without inclusion of gas-phase contributions to the measured organic species.
**Figure B3** Correlation of the timeline of observations and correlation of factor profile of the corresponding factor extracted by PMF analysis from the dataset with or without inclusion of gas-phase contributions. The labeled number refers to the factor number in Figure B2.
Appendix C: Supplemental information for Chapter 4

Appendix C1. Valve-less injection interface (VLI) operation

The VLI controls the helium gas flow direction in one of its two capillaries that serves to connect the sampling and analysis sides of the system. This flow switch is controlled through the vent and purge valves and by adjusting the pressure at the tee joining the two capillaries with the GC column. Pressure balance is achieved using an electronic pressure controller (EPC) module of the GC system (model 6890, Agilent, CA) and a fixed external pressure regulator on the purge supply line. Normally, the helium carrier gas supplied by the EPC is split between the GC column for sample analysis and the cell or the vent. During sample injection, the purge valve is opened and the vent valve closed, resulting in transfer of the sample from the FT to the GC.

Appendix C2. Organic chemical analysis

The chromatographic separation of desorbed organics is achieved using a GC (Agilent 6890) with a capillary column (Rxi-5Sil MS; 30 m length x 0.25 mm ID, 0.25 µm film thickness, Restek). The GC oven temperature is increased at 15 °C/min from 45 °C to 150 °C then at 9 °C/min from 150 °C to 330 °C, and held at 330 °C for 4 minutes. Identification and quantification are achieved using a quadrupole mass selective detector (Agilent 5973 MSD) calibrated with authentic standards (Kreisberg et al., 2009) that are injected into the F-CTD. The transfer efficiency and detector drift during ambient sampling are monitored by co-injection of internal standards with each ambient run (Worton et al., 2012). The internal standards are introduced into the downstream side of the F-CTD using an automatic liquid injection system described elsewhere (Isaacman et al., 2011).

Appendix C3. Operating conditions for evaluations of overall CE

The operating conditions for thermal desorption transfer method are shown in Figure C3 and described as follows: 1) liquid standards were injected into the I-CTD; 2) liquid standards were vaporized at 300 °C and transferred to the F-CTD by helium carrier gas at 150 ml/min for three minutes; 3) following completion of the transfer and after the I-CTD was cooled down to room temperature, the F-CTD was purged by zero air at 10 L/min for 30 or 90 minutes; 4) after the zero air purge, organics remaining in the F-CTD ("measured amount") were thermally desorbed and transferred into the GC/MS for analysis. The amount of organics determined using the same approach but without any zero air purging was defined as the reference amount. The CE of the F-CTD for each compound was calculated by dividing the measured amount for each component by its corresponding reference amount.

The operating conditions for the evaporation transfer method are shown in Figure C4 and described as follows: 1) liquid standards were injected into the I-CTD; 2) standards evaporating from the I-CTD at room temperature were transferred in 10 L/min zero air flow into the F-CTD; 3) after 30 minor 90 min collection, organics collected by the F-CTD were thermally desorbed and introduced into the GC/MS while the I-CTD was maintained at room temperature; 4) after the F-CTD was cooled back down to 28 °C, organics remaining on the I-CTD were thermally desorbed and transferred into the F-CTD; 5) these less volatile organics collected by the F-CTD were thermally transferred to the GC/MS. The CE for each compound was calculated as the sum
of the measured amount in both steps 3 and 5 divided by the reference amount for each compound.

**Appendix C4. Particle losses inside the denuder and particle collection efficiency of the F-CTD**

Particle losses within the denuder and the particle collection efficiency (CE) of the F-CTD were determined by measuring the difference between upstream and downstream ambient particle number size distributions, measured by an optical particle spectrometer (Droplet Measurements, model UHSAS) in the size range of ~0.05 to 1.00 μm optical equivalent diameter at 10 L/min sampling flow. The collection efficiency of the F-CTD for particles less than 0.05 μm was measured by a home built scanning mobility particle sizer using TSI model 3081 differential mobility analyzer using laboratory generated particles in the range of ~30 to 340 nm. The particles were generated by nebulizing dioctyl sebacate.

**Appendix C5. Reference:**


Appendix C6 Tables and figures

Table C1. Chemical formulae, retention times, and concentrations of organic compounds used to evaluate the overall SVOC CE of the F-CTD.

<table>
<thead>
<tr>
<th>Compounds</th>
<th>Chemical Formula</th>
<th>Retention Time</th>
<th>Simulated ambient concentrations</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>30 min sampling</td>
</tr>
<tr>
<td>n-tetradecane</td>
<td>C_{14}H_{30}</td>
<td>15.02</td>
<td>33.33</td>
</tr>
<tr>
<td>n-pentadecane</td>
<td>C_{15}H_{32}</td>
<td>16.07</td>
<td>33.33</td>
</tr>
<tr>
<td>n-hexadecane</td>
<td>C_{16}H_{34}</td>
<td>17.15</td>
<td>33.33</td>
</tr>
<tr>
<td>n-heptadecane</td>
<td>C_{17}H_{36}</td>
<td>18.24</td>
<td>33.33</td>
</tr>
<tr>
<td>n-octadecane</td>
<td>C_{18}H_{38}</td>
<td>19.34</td>
<td>33.33</td>
</tr>
<tr>
<td>n-nonadecane</td>
<td>C_{19}H_{40}</td>
<td>20.42</td>
<td>33.33</td>
</tr>
<tr>
<td>n-eicosane</td>
<td>C_{20}H_{42}</td>
<td>21.47</td>
<td>33.33</td>
</tr>
<tr>
<td>phytane</td>
<td>C_{20}H_{42}</td>
<td>19.41</td>
<td>33.33</td>
</tr>
<tr>
<td>dodecylbenzene</td>
<td>C_{16}H_{30}</td>
<td>20.29</td>
<td>83.33</td>
</tr>
<tr>
<td>1,5-dimethylnaphthalene</td>
<td>C_{12}H_{12}</td>
<td>15.56</td>
<td>16.67</td>
</tr>
<tr>
<td>1,2-dimethylnaphthalene</td>
<td>C_{12}H_{12}</td>
<td>15.69</td>
<td>16.67</td>
</tr>
<tr>
<td>phenanthrene</td>
<td>C_{14}H_{10}</td>
<td>19.28</td>
<td>16.67</td>
</tr>
<tr>
<td>chrysene</td>
<td>C_{12}H_{12}</td>
<td>26.04</td>
<td>16.67</td>
</tr>
<tr>
<td>9-fluorenone</td>
<td>C_{13}H_{8}O</td>
<td>18.79</td>
<td>41.67</td>
</tr>
<tr>
<td>xanthone</td>
<td>C_{13}H_{8}O_{2}</td>
<td>20.11</td>
<td>41.67</td>
</tr>
<tr>
<td>anthraquinone</td>
<td>C_{14}H_{8}O_{2}</td>
<td>21.29</td>
<td>41.67</td>
</tr>
<tr>
<td>4,4'-dimethoxybenzophenone</td>
<td>C_{15}H_{14}O_{3}</td>
<td>23.96</td>
<td>41.67</td>
</tr>
<tr>
<td>5-octyldihydro-2(3H)-furanone</td>
<td>C_{12}H_{22}O_{2}</td>
<td>18.01</td>
<td>100</td>
</tr>
<tr>
<td>2-pentadecanone</td>
<td>C_{15}H_{30}O</td>
<td>18.21</td>
<td>83.33</td>
</tr>
<tr>
<td>2-octadecanone</td>
<td>C_{18}H_{36}O_{2}</td>
<td>21.49</td>
<td>83.33</td>
</tr>
<tr>
<td>1-tridecanal</td>
<td>C_{13}H_{26}O_{2}</td>
<td>16.19</td>
<td>33.33</td>
</tr>
</tbody>
</table>
The transfer line
The metal fiber filter

**Figure C1** Schematic diagrams of the F-CTD and F-CTD housing assembly. The inlet (~ 0.5 cm ID) and outlet (~ 0.2 cm ID) of the F-CTD consist of 2.5 cm long ~0.6 cm (1/4 inch) and ~0.3 cm (1/8 inch) OD tube extensions for compression fitting seals. The transfer line is made of a 10 cm long SS capillary with ~1.6 mm (1/16 inch) OD and ~0.5 mm (1/50 inch) ID, which is brazed into the capsule on the upstream side of the filter to allow back-flushing during thermal desorption. The F-CTD can be heated to 300°C in approximately 4 minutes and cooled back down to 30°C in approximately 8 minutes when room temperature is ~ 20°C. The temperature in the center of the filter, the furthest point from the heater measured by a thermocouple in contact with the filter surface, reached its setpoint at approximately the same time as the heating block and the temperature difference between the center of the filter and the heating block was <10 °C when the temperature of the heating block was held isothermally at its setpoint.
Figure C2 Schematic diagram of the focusing trap assembly.
**Figure C3** Operating conditions for the evaluation of gas CE of the F-CTD using the thermal desorption method: (1) Injection of organics into the I-CTD; (2) Transfer organics from the I-CTD to F-CTD by thermal desorption; (3) Zero air purge for 30 min (shown) or 90 min; (4) Thermal desorption of the collected organics, including sample trapping (4a) and sample injection (4b).
Figure C4 Operating conditions for the evaluation of the gas CE of the F-CTD using evaporation transfer: (1) Injection of liquid standards; (2) Zero air purge for 30 min (shown) or 90 min; (3) Thermal desorption of the collected organics; (4) Transfer from the I-CTD to F-CTD; (5) Thermal desorption of the collected organics. Thermal desorption consists of sample trapping and sample injection (see Figure C3).
Figure C5 Operating conditions of the SV-TAG system during ambient measurements. 1) the previous ambient sampling; 2) thermal desorption of the previous samples and initiate the GC analysis at the conclusion of thermal desorption from the F-CTD; 3) cool down the F-CTD and FT in preparation for next ambient sampling; 4) the ambient sampling while the GC is analyzing the previous sample.
Figure C6 Optimization of the recovery of less volatile organic compounds: recovery of n-alkanes as a function of carbon number at different purge flow rates.
Figure C7 Particle collection efficiency of the F-CTD as a function of the mobility diameter down to ~ 30 nm. The measurements were made with a home built scanning mobility particle sizer using TSI model 3081 differential mobility analyzer. The particles were generated by nebulizing dioctyl sebacate (DOS).
Figure C8 Particle collection efficiency of the F-CTD (left axis) inferred from the number concentration of ambient particles measured upstream and downstream of the F-CTD. The volume size distribution is shown on the right axis.
Figure C9 Effect of organic loading on the recovery of individual compounds. The abundance of the total ion is normalized to its mean value during the sampling period.