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ALDEHYDE AND OTHER VOLATILE ORGANIC CHEMICAL EMISSIONS IN FOUR FEMA TEMPORARY HOUSING UNITS - FINAL REPORT

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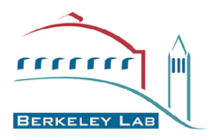
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Randy Maddalena, Marion Russell, Douglas P. Sullivan, and Michael G. Apte

Environmental Energy Technologies Division

November 2008



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Aldehyde and other Volatile Organic Chemical Emissions in Four FEMA Temporary Housing Units – Final Report

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November 2008

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ABSTRACT

As part of an ongoing effort with the U.S. Centers for Disease Control and Prevention (CDC), Lawrence Berkeley National Laboratory (LBNL) entered into an interagency agreement with CDC to help identify mitigation strategies for reducing indoor emissions of volatile organic compounds (VOCs) including formaldehyde in Federal Emergency Management Administration (FEMA) temporary housing units (THUs)¹. Four unoccupied FEMA THUs were studied to assess their indoor emissions of VOC including formaldehyde. Indoor measurement of whole-THU VOC and aldehyde emission factors ($\mu g h^{-1}$ per m² of floor area) for each of the four THUs were made at FEMA's Purvis Mississippi staging yard using a mass balance approach. Measurements were made in the morning, and again in the afternoon in each THU. Steady-state indoor formaldehyde concentrations ranged from 378 μ g m⁻³ (0.31ppm) to 632 μ g m⁻³ (0.52 ppm) in the morning, and from 433 μ g m⁻³ (0.35 ppm) to 926 μ g m⁻³ (0.78 ppm) in the afternoon. THU air exchange rates ranged from 0.15 h^{-1} to 0.39 h^{-1} . A total of 45 small (approximately 0.025 m^2) samples of surface material, 16 types, were collected directly from the four THUs and shipped to Lawrence Berkeley Laboratory. The material samples were analyzed for VOC and aldehvde emissions in small stainless steel chambers using a standard, accurate mass balance method. Quantification methods for the VOCs included high performance liquid chromatography for formaldehyde and acetaldehyde, ion chromatography for the acetic acid, and gas chromatography-mass spectrometry for the remaining VOCs. Material specific emission factors (μ g h⁻¹ per m² of material) were quantified. Approximately 80 unique VOCs were initially identified in the THU field samples, of which forty-five were quantified either because of their toxicological significance or because their concentrations were high. Whole-trailer and material specific emission factors were calculated for 33 compounds. The THU emission factors and those from their component materials were compared against those measured in other types of housing and the materials used in their construction. Whole THU emission factors for most VOCs were similar to those from comparative housing. The three exceptions were large emissions of formaldehyde, acetic acid, TMPD-DIB (a common plasticizer in vinyl products), and somewhat elevated emission of phenol. Of these compounds, formaldehyde was the only one with toxicological significance at the observed concentrations. Whole THU formaldehyde emissions ranged from 173 to 266 μ g m⁻² h⁻¹ in the morning and 257 to 347 μ g m⁻² h⁻¹ in the afternoon. Median formaldehyde emissions in previously studied site-built and manufactured homes were 31 and 45 μ g m⁻² h⁻¹, respectively. Only one of the composite wood materials that was tested appeared to exceed the US Department of Housing and Urban Development (HUD) formaldehyde emission standard for new material but several of the materials exceeded if the decline in emission with aging is considered. The high loading factor (material surface area divided by THU volume) of composite wood products in the THUs and the low fresh air exchange relative to the material surface area may be responsible for the excessive concentrations observed for some of the VOCs and formaldehvde.

¹ This is a final project report which supersedes the previously submitted interim report by the same authors titled INTERIM REPORT: VOC AND ALDEHYDE EMISSIONS IN FOUR FEMA TEMPORARY HOUSING UNITS dated 4 May, 2008.

TABLE OF CONTENTS

ABSTRACT	i
TABLE OF CONTENTS	ii
LIST OF FIGURES	iii
LIST OF TABLES	iv
LIST OF ABBREVIATIONS	vi
EXECUTIVE SUMMARY	vii
RECOMMENDATIONS FOR FUTURE WORK	
INTRODUCTION	
EMISSION FACTORS	1
FORMALDEHYDE EMISSIONS FROM BUILDING MATERIALS – BACKGROUND INFORMATION Formaldehyde Emission Behavior VOLATILE ORGANIC COMPOUND (VOC) EMISSIONS FROM BUILDING MATERIALS - BACKGROUND INFORMAT	3
METHODS	
OVERVIEW OF EXPERIMENTAL APPROACH	4
DESCRIPTION OF STUDY UNITS	4
AIR SAMPLING AND ANALYSIS	
Volatile Organic Chemicals	
Low Molecular Weight Aldehydes	
Acetic Acid QUALITY ASSURANCE	
QUALITY ASSURANCE	
MEASUREMENT OF WHOLE TRAILER CONCENTRATIONS	
Collection and Characterization of Indoor Materials	
MEASUREMENT OF MATERIAL SPECIFIC EMISSION FACTORS	
DATA ANALYSIS	
RESULTS	
MATERIAL SPECIFIC LOADING RATIOS	9
WHOLE TRAILER VENTILATION AND VOC MEASUREMENTS	
MATERIAL SPECIFIC VOC MEASUREMENTS	
PERCENT CONTRIBUTION OF MATERIAL SPECIFIC EMISSIONS TO WHOLE TRAILER MEASUREMENTS	
DISCUSSION	11
VOC EMISSIONS FROM BUILDING MATERIALS	12
ALDEHYDE EMISSIONS FROM BUILDING MATERIALS	
CONCLUSIONS	15
RECOMMENDATIONS FOR FUTURE WORK	16
REFERENCES:	17
TABLES:	19
FIGURES:	42

LIST OF FIGURES

Figure 1. Preparation for indoor sampling in a THU. Half inch holes were drilled into the THU door for insertion of ¼" stainless steel sampling tubes. A sampling tube and sample pump are seen in the foreground
Figure 2. Collection of indoor sample through the THU door
Figure 3. Example of tracer gas experiment determining ventilation rate in trailer showing initial stabilization period followed by the linear decay region. The ventilation rate is determined from the slope of the decay curve in the linear region as described in the text. The response shown here for Trailer 1 is typical of all the units tested
Figure 4. Comparison of measured indoor air concentration (ppm) data for new site-built and manufactured homes (Hodgson et. al., 2000), German residences (Hippelein, 2004) and the four THUs. The data are reported as geometric mean (GM) with error bars representing one geometric standard deviation (GSD). The GSD for the Hippelein (2004) data were calculated from the arithmetic mean and standard deviation
Figure 5. Comparison of GM (GSD error bars) measured whole building VOC emission factors (emissions per floor area) for seven new site built houses, four new manufactured houses (Hodgson et. al. 2000), and the four THUs studied in this project. Note that this chart is plotted on a logarithmic scale

LIST OF TABLES

Table 1. Specifications and Ventilation Characteristics of the Temporary Housing Units 1	19
Table 2. Projected Surface Area of Indoor Materials (m ²) 1	19
Table 3. Description of Surface Materials Harvested from Trailers and Tested for Emissions2	20
Table 4. Surface Coverings and Finishes on Tested Materials	21
Table 5. Surface Loading Ratios and Area-Specific Clean Air Flow Rates	22
Table 6. Environmental Conditions 2	23
Table 7 Target VOCs Identified in Temporary Housing Units 2	24
Table 8. Measured Steady-state VOC Concentrations ($\mu g/m^3$) in Field Samples	25
Table 9. Whole Trailer Emission Rates Normalized to Floor Area ($\mu g m^{-2} h^{-1}$)	26
Table 10. Material Specific Emission Factors ($\mu g m^{-2} h^{-1}$) for the Dutchmen trailer	28
Table 11. Material Specific Emission Factors ($\mu g m^{-2} h^{-1}$) for the Pilgrim trailer	29
Table 12. Material Specific Emission Factors ($\mu g m^{-2} h^{-1}$) for the Coachman trailer	30
Table 13. Material Specific Emission Factors ($\mu g m^{-2} h^{-1}$) for the Cavalier trailer	31
Table 14. Material Emission Factors Normalized to Whole Trailer Floor Area (μ g m ⁻² h ⁻¹) for the Dutchmen trailer	
Table 15. Material Emission Factors Normalized to Whole Trailer Floor Area (μ g m ⁻² h ⁻¹) for the Pilgrim trailer	9 33
Table 16. Material Emission Factors Normalized to Whole Trailer Floor Area (μ g m ⁻² h ⁻¹) for the Coachmen trailer	9 34
Table 17. Material Emission Factors Normalized to Whole Trailer Floor Area (μ g m ⁻² h ⁻¹) for the Cavalier trailer	
Table 18. Total ($\mu g \text{ m}^{-2} \text{ h}^{-1}$) and Percent Contribution of Each Material to Area Normalized Whole Trailer Emission Rates for the Duchmen	36
Table 19. Total (µg m ⁻² h ⁻¹) and Percent Contribution of Each Material to Area Normalized Whole Trailer Emission Rates for the Pilgrim	37
Table 20. Total (µg m ⁻² h ⁻¹) and Percent Contribution of Each Material to Area Normalized Whole Trailer Emission Rates for Coachmen	37

Table 21. Total (µg m ⁻² h ⁻¹) and Percent Contribution of Each Material to Area Normalized Whole Trailer Emission Rates for the Cavalier
Table 22. Comparison of Sum Material Specific Emission* (µg m ⁻² h ⁻¹) with Measured Whole Trailer Emission
Table 23. Material specific aldehyde emissions from cabinetry, passage door, and subfloor used to fabricate a new manufactured house
Table 24. Material specific emission factors of terpene hydrocarbons from indoor sources used to fabricate a new manufactured house
Table 25. Reported Formaldehyde Emission Factors from CARB's Battelle (1996) study ¹ 41

LIST OF ABBREVIATIONS

ACH	Air Exchanges per Hour
ASTM	American Society for Testing and Materials
CARB	California Air Resources Board
CDC	Centers for Disease Control and Prevention
CPSC	US Consumer Product Safety Commission
CV	Coefficient of Variation
FEMA	Federal Emergency Management Administration
GM	Geometric Mean
GSD	Geometric Standard Deviation
HUD	US Department of Housing and Urban Development
HWPW	Hardwood Plywood
LBNL	Lawrence Berkeley National Laboratory
NCEH	National Center for Environmental Health
NIOSH	National Institute for Occupational Safety and Health
NIST	National Institute of Standards and Technology
ORNL	Oak Ridge National Laboratory
ppm	parts per million
ppb	parts per billion
PVC	polyvinylchloride
REL	Recommended Exposure Limit
TAC	Toxic Air Contaminant
THU	Temporary Housing Units
TMPD-DIB	2,2,4-Trimethyl-1,3-pentanediol diisobutyrate
TMPD-MIB	2,2,4-Trimethyl-1,3-pentanediol monoisobutyrate
UFFI	Urea Formaldehyde Foam Insulation or expandable foam insulation
VOC	Volatile Organic Chemical

EXECUTIVE SUMMARY

This document superseded the previously submitted interim report titled "INTERIM REPORT: VOC AND ALDEHYDE EMISSIONS IN FOUR FEMA TEMPORARY HOUSING UNITS" by the same authors dated 4 May, 2008. This final report includes updated data on acetic acid, a more detailed comparison of the results to regulatory product standards and published emission factors and updated figures and tables.

The objectives of this study have been to 1) directly measure indoor concentrations and whole trailer emission factors in four unoccupied temporary housing units (THUs) for a range of volatile organic compounds (VOCs) and aldehydes including formaldehyde, and 2) determine materials specific emission factors of these compounds from individual surface materials collected directly from the THUs providing initial information into the magnitude and potential sources of indoor pollutant loadings in the tested THUs. The results of this study will be used by the U.S. Centers for Disease Control and Prevention to help identify mitigation strategies for reducing indoor emissions of VOCs including formaldehyde.

The Federal Emergency Management Administration (FEMA) has supplied over 100,000 emergency THUs to families that lost their homes in Louisiana and Mississippi during the Hurricane Katrina and Rita disasters. FEMA owns approximately 160,000 of these THUs. Some are deployed to other parts of the U.S., some are used to house emergency workers, and many are in storage. Concerns about the indoor environmental quality in the THUs have arisen based on occupant health complaints and concerns. These health concerns have been identified by physicians treating THU occupants, and through risk analyses of indoor air quality measurements made in both occupied and unoccupied units. These measurements were reported by the Sierra Club and by the Centers for Disease Control and Prevention (CDC). Formaldehyde concentrations observed in both occupied and unoccupied THUs have exceeded the National Institute for Occupational Safety and Health (NIOSH) recommended exposure limit (REL) of 0.016 ppm, often by a factor of 10 or greater. The NIOSH REL was based on the analytical limit of detection and not on health effects data.

Measured emission factors for individual building materials can be used to assess the contribution of specific materials to the overall indoor pollutant load using mass balance modeling. Emission factors describe the mass of a particular compound released by a specific material per hour, per unit area of material. Measured emission factors provide a means to directly compare emission characteristics from one material to another. Emission factors from materials are dependent on a range of environmental parameters such as temperature, relative humidity and boundary layer diffusion characteristics, which are influenced by air flow across the surface. These parameters need to be considered when emission factors are compared.

When describing emissions from a single material, i.e., fiberboard or flooring, emission factors are typically expressed in terms of the projected surface area of the material itself. However, when describing the emissions from a composite assembly of materials, such as a house or travel trailer that is constructed from a variety of component pieces, it is difficult to isolate a single emission source. In this case, it is customary to present emissions of a particular compound as a net mass emitted per unit area of floor, per hour. Both of these emission factor metrics have the same units: $\mu g m^{-2} h^{-1}$. It is important to understand the distinction between emissions on a permaterial area versus a net per-floor area basis when studying material emission characteristics.

Sources contributing to elevated formaldehyde indoors are primarily related to building products and furnishings. Formaldehyde is only one compound of concern that is emitted from these materials. A range of VOCs typically present when formaldehyde is observed, are also emitted from materials. Like formaldehyde, which is a toxic air contaminant, many of the VOCs are known to have low odor thresholds, high potency as respiratory irritants, and in some cases carcinogenicity.

This report presents results from experiments designed to assess their indoor emissions of VOCs, including formaldehyde, in four unoccupied FEMA THUs. Whole-THU emissions were measured, and then selected material samples were collected directly from the four units and their material specific emission rates were measured in small chambers. Standard analytical methods employing rigorous quality control were used to quantify a range of VOCs and aldehydes in the air in both the whole-THU and chamber emissions experiments and these measurements were used along with characteristics of the THUs to estimate emission rates.

The THUs selected for study were all of the travel trailer design including a Thor Industries Dutchmen manufactured September 2005, a Pilgrim International manufactured October 2005, a Coachmen's Spirit of America manufactured October 2006 and a Gulfstream Coach Cavalier manufactured March 2006. The units were in excellent condition. The approximate floor areas ranged from 221 - 240 square feet. The Dutchman trailer was equipped with an additional pullout section approximately 14 feet long by 3 feet (~42 ft²) that was not opened up during sampling.

Measurement of whole-THU VOC and aldehyde emission factors (μ g h⁻¹ per m² of floor area) for each of the four THUs were made at FEMA's Purvis Mississippi staging yard. Measurements were made in the morning, and again in the afternoon in each THU. Steady-state indoor formaldehyde concentrations ranged from 378 μ g m⁻³ (0.31ppm) to 632 μ g m⁻³ (0.52 ppm) in the morning, and from 433 μ g m⁻³ (0.35 ppm) to 926 μ g m⁻³ (0.78 ppm) in the afternoon. THU air exchange rates ranged from 0.15 h⁻¹ to 0.39 h⁻¹.

A total of 45 small (approximately 0.025 m^2) samples of surface material, 16 types, were collected directly from the four THUs and shipped to Lawrence Berkeley Laboratory (LBNL). Material specific emission factors were determined using small chambers generally following the ASTM Standard Guide D-5116-97. The material samples were analyzed for VOC emissions in small stainless steel chambers using a standard, accurate mass balance method. Quantification of VOCs was done via gas chromatography – mass spectrometry; low molecular weight aldehydes via high performance liquid chromatography; and acetic acid via an ion chromatography method. Material specific emission factors (μ g h⁻¹ per m² of material) were quantified. Approximately 80 unique VOCs were initially identified in the THU field samples, of which forty-five were quantified either because of their toxicological significance or because their concentrations were high.

All THUs had a significant fraction of the internal surface area (walls, ceiling, cabinet walls) constructed of 1/8-inch plywood with a vinyl or PVC skin or simulated wood finish. All units had sheet vinyl flooring while the Dutchmen and Pilgrim also had carpeted areas. All countertops were particleboard surfaced with high-pressure laminate. A variety of wood products were used for the sub-floor and for the bench and bed platforms.

Whole-trailer and material specific emission factors were calculated for 33 compounds. The THU emission factors and those from their component materials were compared against those

measured from other types of housing and the materials used in their construction. Whole THU emission factors for most VOCs were similar to those from comparative housing measured in the U.S (all approximately 6 months old or less). Four exceptions were large emissions of formaldehyde, acetic acid and TMPD-DIB (2,2,4-Trimethyl-1,3-pentanediol diisobutyrate - a common plasticizer in vinyl products), and somewhat elevated emissions for phenol. A detailed assessment of the toxicology and risk associated with the identified indoor pollutants was not performed but of the compounds that were identified, the observed concentration of formaldehyde is thought to be of toxicological significance. Additionally, several VOCs (dodecane, tridecane, p-xylene, alpha-pinene, beta-pinene and hexanal) were measured in the four THUs at concentrations above those reported in a study of 39 German homes. Acetic acid was also measured at levels that were well above reported odor thresholds for VOCs.

Whole THU formaldehyde emissions ranged from 173 to 266 μ g m⁻² h⁻¹ in the morning and 257 to 347 μ g m⁻² h⁻¹ in the afternoon. Median formaldehyde emissions in previously studied sitebuilt and manufactured homes (approximately 6 months old or less) were 31 and 45 μ g m⁻² h⁻¹, respectively.

The material specific formaldehyde emission factor measurements showed that the highest material emissions were from the cabinet walls, sub flooring, and the bench materials (the fabric and foam materials also showed elevated emissions, but these may be due to the re-emission of formaldehyde that had sorbed to the material from the indoor air, rather than as primary emitters). Only one material, the Cavalier plywood cabinet wall (490 μ g m⁻² h⁻¹) exhibited emissions in excess of the HUD standard of 130 μ g m⁻² h⁻¹ for new material. However, the material in the THUs had aged and as a result the emission rates are expected to be lower than initial emissions from new material. If this aging is accounted for then several of the materials in the THUs may have emission that exceeded the HUD standard when the materials were new. Knowledge on the rate of decline in emission rates with aging is limited so it is difficult to conclude that the aged materials would have been below the HUD standard or within previously reported ranges for other structures. Nevertheless, even with the aged materials, the emission factors for phenol, TMPD-DIB, acetic acid and formaldehyde remained higher in the THUs than the new homes.

We conclude that whole trailer formaldehyde emission factors are high, but the materials emission factors may be within those commonly found in the building industry. This indicates a difference in the construction/design that may lead to elevated concentrations and whole trailer emission rates. Three features of material application in the THUs differ from most other dwellings: 1) the extensive use of lightweight composite wood products, 2) very high surface loading of composite wood products and 3) low fresh air per unit surface area of composite wood products in the THUs.

Much of the projected surface area in the THUs (wall, ceiling, and cabinetry) use 1/8" hardwood plywood (HWPW). The wood product loading factor of the THU is far higher than in houses that use gypsum board for walls and ceilings. These high loading factors in combination with observed ventilation rates may be the primary reason for the unusually high rates at which formaldehyde mass is emitted into the THU. Considering this in terms of the area-specific clean air flow rates, the high material loading ratio in the units combined with relatively low fresh air ventilation rates results in area-specific air flow rates that are quite low relative to other housing types. With all other factors being equal, the steady-state concentrations indoors are inversely proportional to the air exchange rates. The THUs in this study are not outfitted for increased

ventilation and may be under ventilated for housing with such small volume. Although low ventilation does not directly affect the measured formaldehyde emission rates presented in this report, it can influence the concentrations experienced by the THU occupants.

The results of this study are not statistically representative of the entire fleet of FEMA THUs because the study was based on a convenience sample of four THUs. Nonetheless, the measured material-specific emission factors for volatile organic compounds, including formaldehyde, were similar to values reported in the literature for materials. However, it is important to consider that the materials in this study were both aged and allowed to interact with emissions from other materials. Formaldehyde and some of the other VOCs measured in the unoccupied THUs and the associated whole trailer emission factors were found to be higher, sometimes much higher, than what is typically found in residential environments. The difference between these THUs and other housing appears to be the very high composite wood surface areas relative to room volume used in the travel trailer design and the low ventilation rates in terms of low area-specific fresh air flow rates in the THUs.

Recommendations for future work

This report provides a preliminary assessment into the effect of THU design and material choices on indoor VOC and aldehyde concentrations. It is by no means definitive because we studied only four THU models produced by four manufactures and the focus of this study was limited to the travel trailers, while a significant portion of THUs are park trailer models, and mobile homes. A systematic assessment across a wider range of THU makes and models including a better characterization of fresh air ventilation rates under occupied conditions could provide a better understanding of the time integrated exposure concentrations in occupied units.

The results in this report do not yet address temperature and humidity effects on material emissions within the studied units. It is hypothesized that at the higher temperature and relative humidity conditions found in the summertime in the southeastern portions of the US, emissions of formaldehyde from the urea-formaldehyde composite materials will increase. Chamber experiments and a seasonal study designed to investigate the potential effects of temperature and humidity should be completed.

INTRODUCTION

This document superseded the previously submitted interim report titled "INTERIM REPORT: VOC AND ALDEHYDE EMISSIONS IN FOUR FEMA TEMPORARY HOUSING UNITS" by the same authors dated 4 May, 2008. This final report includes updated data on acetic acid, a more detailed comparison of the results to regulatory product standards and published emission factors and updated figures and tables.

FEMA has supplied over 100,000 emergency THUs to families that lost their homes in Louisiana and Mississippi during the Hurricane Katrina and Rita disasters. FEMA owns approximately 160,000 of these THUs. Some are deployed to other parts of the U.S., some are used to house emergency workers, and many are in storage. Concerns about the indoor environmental quality in the THUs have arisen based on occupant health complaints and concerns. These concerns have been identified by physicians treating THU occupants, and through risk analyses of indoor air quality measurements made in both occupied and unoccupied units. These measurements were reported by the Sierra Club and by the Centers for Disease Control. Formaldehyde concentrations observed in both occupied and unoccupied THUs have exceeded the National Institute for Occupational Safety and Health (NIOSH) recommended exposure limit (REL) of 0.016 ppm, often by a factor of 10 or greater. The NIOSH REL was based on the analytical limit of detection and not on health effects data.

Although formaldehyde levels in the THUs was highlighted by the Sierra Club survey, and by media focus, a concern has existed that other irritating, odorous, or potentially toxic volatile organic compounds (VOC) may be emitted from the THU construction materials and furnishings, and that the design of the THUs, including extensive use of plywood, particle board and laminated material in combinations with low ventilation rates may lead to elevated exposure concentrations. A careful study of indoor VOC emissions in the whole trailers and from the individual THU materials was needed to identify sources of indoor pollutants and begin to assess exposure concentrations that result from these emissions.

Emission Factors

Measured emission factors for individual building materials can be used to assess the contribution of specific materials to the overall indoor pollutant load using mass balance modeling (Hodgson et. al., 2004). Emission factors describe the mass of a particular chemical released by a material per hour and per unit surface area. Measured emission factors provide a means to directly compare emission characteristics from one material to another. Emission factors from materials can be influenced by a range of environmental parameters such as temperature, relative humidity and boundary layer diffusion characteristics, which are influenced by air flow across the surface. It is important that these parameters are consistent when emission factors are compared. Measurement method standardization helps to ensure this.

When describing emissions from a single material, i.e., fiberboard or flooring, emission factors are typically expressed in terms of the projected surface area of the material itself. However, when describing the emissions in a complete structure, such as a house or travel trailer that is composed of a variety of different materials that can release and absorb air pollutants, it is difficult to isolate a single emission source. In this case, it is customary to present emissions of a particular compound as a net mass emitted per unit area of floor, per hour. Both of these emission factor metrics have the same units: $\mu g m^{-2} h^{-1}$. It is important to recognize the

distinction between emissions on a per-material area versus a net per-floor area basis when studying material emission characteristics. The convention followed in this work is to report whole-trailer emission factors on a floor area basis while the individual materials are reported on a projected surface area basis.

Formaldehyde Emissions From Building Materials – Background Information

Sources contributing to elevated formaldehyde indoors are primarily related to building products and furnishings. Formaldehyde is only one compound of concern that is emitted from these materials. A range of VOCs typically present when formaldehyde is observed, are also emitted from materials (Hodgson, 1999). Like formaldehyde, which is a toxic air contaminant, many of the VOCs are known to have low odor thresholds, high potency as respiratory irritants, and in some cases carcinogenicity. However, given the toxicity and prevalence of formaldehyde in residences it has been the focus of a number of studies over the last several decades.

The problem of excessive formaldehyde emissions from building materials reached national awareness starting in the early 1980s with the increase in commercial and industrial use of urea formaldehyde as a bonding agent and as an expanded foam insulation (UFFI). The US Consumer Product Safety Commission (CPSC) had reported health complaints caused by UFFI since 1972. In 1980 the National Academy of Science advised maintaining the lowest practical formaldehyde concentrations in order to minimize possible adverse effects on public health, based upon emerging results from an ongoing carcinogenicity study (NAS 1980). A heightened concern began with the emergence of health effects in occupants of mobile homes (Hileman, 1982). In 1982 the Consumer Product Safety Commission placed a ban on UFFI (CPSC 1982). This ban was subsequently lifted a year later by court order (CPSC 1983). However, the use of UFFI as a building material was curtailed by the industry.

In 1984 the U.S. Department of Housing and Urban Development (HUD) established formaldehyde product standards for all plywood and particleboard materials using bonding, coating, or surface finishing systems containing formaldehyde when installed in manufactured homes (Turner et al. 1996). The standard is embodied in the HUD Standard 24 CFR Chapter XX Part 3280, *Manufactured Home Construction and Safety Standards* (HUD 2001). The standard is based upon the ASTM emission testing method E-1333 that continues to be used (ASTM 2002). The standard was intended to cap the mass of formaldehyde that emanated from fresh wood composite materials in terms of concentration in a test chamber using standardized surface loading ratios and area specific air flows or air exchange rates. The standard was developed for testing newly manufactured wood products prior to their use in construction².

The wood products industry adopted the HUD standard in the U.S. during the 1980s. Subsequent surveys indicated that because the reduction of the mass emission rate of formaldehyde from wood products and the discontinuation of the use of UFFI in residential

² The HUD safety standards for certified plywood and particleboard used in manufactured home construction require that formaldehyde concentrations not exceed 200 ppb (0.246 mg/m³) from plywood and 300 ppb (0.369 mg/m³) from particleboard, as measured under the conditions specified in ASTM Method E1333. Engineered wood products are tested with specified loading ratios for particleboard and plywood of 0.43 m² of material per m³ of test chamber volume (0.13 ft²/ft³), and 0.95 m²/m³ (0.29 ft²/ft³), respectively. Using the operating conditions and formaldehyde emissions rate equation specified in the standard, the initial formaldehyde emissions rates from the newly manufactured materials are 430 µg/m²/h (8.81 x 10⁻⁸ lb/ft²/h) for particleboard and 130 µg/m² • h (2.66 x 10⁻⁸ lb/ft² • h) for plywood.

construction, formaldehyde levels in residences dropped substantially (Azuma et al. 2006) through the 1980s and 1990s.

Formaldehyde Emission Behavior

Past research has established that the rate at which formaldehyde is emitted from some building products drops slowly as the materials ages after manufacture. This concept is often brought up when the topic of indoor formaldehyde emissions from materials is discussed. The fact is often used to indicate that indoor formaldehyde concentrations will decrease with time, lessening risk and health problems. However, the rate at which emissions drops is not well determined and will depend upon many factors. A recently released industry association report (SEFA 2008) concluded that emissions can drop by 25% within a month of manufacture and usually drop by half within six months.

A study of emission characteristics of pressed-wood products conducted by Oak Ridge National Laboratory (ORNL) for the U.S. Consumer Product Safety Commission (Matthews 1985) found that the time needed for emissions to drop to approximately 37% of initial rate was between 0.9 and 2.2 years depending on the material tested. These decay rates indicate a rate constant of 1.1 per year and 0.45 per year. The longer decay period (slower decay rate) was for a mixture of materials (particleboard underlayment, industrial particleboard, hardwood plywood paneling and medium density fiberboard). The shorter decay periods were associated with weaker board material at lower starting formaldehyde concentrations.

Using the slower decay period of 2.2 year determined in the ORNL study (Matthews 1985) for materials that are similar to THU materials, and assuming a starting formaldehyde concentration of 300 ppb with an air exchange rate of 0.5 h^{-1} (HUD standard for particle board), the required duration for the concentration in a new trailer to drop to a concentration of 10 ppb (similar to background, ASTDR 1999) is 7.5 years. For the lighter materials with the faster decay rate measure by ORNL, and assuming a starting concentration of 200 ppb, the time to reach 10 ppb is between three and five years.

Another key finding in the ORNL study was the effectiveness of vinyl flooring as a barrier in reduction of formaldehyde emission rates. This finding is salient to the THUs studied in this project in that much of the floor area had sheet vinyl covering and the walls, ceiling, cabinets, and doors were also covered with a polyvinyl chloride (PVC), photo-laminate or vinyl material. The ORNL report found through both modeling and measurements that carpet and cushion covering resulted in approximately a 2.5 fold reduction in formaldehyde emission rates while vinyl flooring reduced emission by approximately 30 fold (Matthews 1985).

Other building material studies have reported on the effectiveness or lack of effectiveness of coatings, layers, laminates, and other coverings showing that different coverings retard emissions differently. Some studies have shown that there can be significant sink effects with certain floor and wall covering materials when used in conjunction with other emitting sources highlighting more complex interactions and effects of flooring and wall assemblies including peak VOCs shifts with respect to time instead of simple decays (Won et al. 2001).

VOC Emissions from Building Materials - Background Information

Considerably less information is available on VOC emissions from construction materials other than formaldehyde. Key sources of new information are Hodgson et al. (1999, 2000, and 2004),

Hodgson and Levin (2003), the California Integrated Waste Management Board (CIWMB 2003), Hipellein (2004) and Won et.al. (2004). For the purposes of this report we are able to make comparisons of residential concentrations and to whole structure VOC emission factors on a perfloor area basis. The sparse data on VOC emissions at the material level make comparisons more tenuous, however, enough data exists to make some qualitative conclusions regarding individual materials' contributions to indoor VOC concentrations in the THUs.

The objectives of this study are to 1) directly measure indoor concentrations and whole trailer emission factors in four unoccupied THUs for a range of VOCs and 2) determine materials specific emission factors from individual surface materials collected directly from the THUs providing initial information into the magnitude and potential sources of indoor pollutant loadings in the tested THUs.

METHODS

Overview of Experimental Approach

Four unoccupied THUs, each produced by a different manufacturer, were selected for study from stock at the FEMA staging yard in Purvis, Mississippi. For each THU, indoor and outdoor air concentrations were determined under steady-state ventilation conditions for a range of pollutants at two separate time points and ventilation rates were measured. After completion of the whole trailer measurements, representative surface materials were cut directly from each THU, packaged and shipped to LBNL for testing in small chambers to determine material-specific VOC emission factors. The projected surface areas of the materials in the THUs were measured and used along with the emission factors to characterize the relative contributions of the materials to total pollutant loads in the THUs.

Description of Study Units

The THUs selected for study included a Thor Industries Dutchmen manufactured September 2005, a Pilgrim International manufactured October 2005, a Coachmen's Spirit of America manufactured October 2006 and a Gulfstream Coach Cavalier manufactured March 2006. The units were unoccupied and in excellent condition. The approximate floor areas ranged from 221 – 240 square feet. The Dutchman trailer was equipped with an additional pullout section approximately 14 feet long by 3 feet (~42 ft²) that was not opened up during sampling.

The trailer dimensions and specifications are summarized in Table 1. The Pilgrim and Cavalier trailers were built to FEMA specification while the Dutchmen and Coachmen were built to HUD standards. The units tested were all travel trailer designs that had either not been previously occupied or had been reconditioned and made ready for re-deployment. The projected surfaces areas of each surface material in the THUs are summarized in Table 2. A description of the individual building material types is provided in Table 3 and the surface covering or finishes are summarized in Table 4.

The trailers were moved to a central staging area at the storage yard on November 9, 2007 and were parked in approximately the same directional orientation. A series of small holes (~6 mm) were drilled in the entrance door of each trailer (Figure 1) to allow insertion of rigid stainless steel sampling tubes for sample collection (Figure 2). Rigid sampling tubes were extended approximately 1 meter into a trailer and elevated 1 meter from the floor to facilitate sampling of VOCs, aldehydes, acetic acid, temperature, relative humidity, and tracer gas concentrations without opening the trailer. Mixing fans were installed in each trailer for use only in mixing the

injected tracer gas to determine each THU's characteristic air exchange ventilation rates. These fans were not otherwise operated during VOC sampling.

After initial setup, the trailers were closed and remained closed to allow the ambient ventilation rates to come to steady-state. Sampling was conducted on November 14, 2007. Temperature, relative humidity and CO₂ concentrations were monitored in each trailer and at a central location outdoors during the experiments using calibrated indoor air quality monitors (Q-Trac Plus; TSI).

Air Sampling and Analysis

Volatile Organic Chemicals

VOC samples were collected and analyzed following USEPA Methods TO-1 and TO-17 (USEPA 1999). VOCs were collected onto multibed sorbent tubes (P/N 012347-005-00; Gerstel or equivalent) with primary bed of Tenax-TA® sorbent backed with a section of Carbosieve®. Prior to use, the sorbent tubes were conditioned by helium purge (~10 cc/min) at 275 °C for 60 minutes and sealed in Teflon capped tubes. VOC samples were collected through a rigid stainless steal tube inserted through the trailer door, directly into the tube for outdoor samples, and directly from the exit port in the small emission chamber. A vacuum pump (Model DOA-P104-AA; Gast) with electronic mass flow controllers (lab), or calibrated personal sampler pumps (field) were used to pull air through the sample tubes at ~100 cc/min. Approximately 6 liters were collected from the whole-trailers and 3 liters from the emission chambers. Flows were verified using a separate calibrated flow meter prior to the emission chamber experiments. The personal sampler pumps used in the field were calibrated prior to use and checked after use. Sorbent tubes were sealed with Teflon lined caps after use and either analyzed the same day or stored on ice or in a freezer until analysis. Sample stability over freezer storage times of more than 2 months have been confirmed previously in our lab for many of the VOCs included in this study.

Sorbent tubes were thermally desorbed for analysis by gas chromatography/mass spectrometry (TD-GC/MS) using a thermodesorption auto-sampler (Model TDSA2; Gerstel), a thermodesorption oven (Model TDS3, Gerstel) and a cooled injection system (Model CIS4; Gerstel). The cooled injection system was fitted with a Tenax-packed glass liner (P/N 013247-005-00; Gerstel). Desorption temperature was 25 °C with a 0.5 minute delay followed by a 60 °C ramp to 250 °C and a 4 minute hold time. The cryogenic trap was held at -10 °C and then heated within 0.2 minutes to 270 °C at a rate of 12 °C/s, followed by a 3-minute hold time. Compounds were resolved on a GC (Series 6890Plus; Agilent Technologies) equipped with a 30 meter HP-1701 14% Cyanopropyl Phenyl Methyl column (Model 19091U-233; Agilent Technologies) at an initial temperature of 1 °C for 0.5 minutes then ramped to 40 °C at 25 °C/min, to 115 °C at 3 °C/min and finally to 250 °C at 10 °C/min holding for 10 minutes.

The resolved analytes were detected using an electron impact MS system (5973; Agilent Technologies). The MS was operated in scan mode. One sample from each trailer was analyzed and all compounds over the detection limit (< 1 to several ng) were identified by library search using the National Institute of Standards and Technology (NIST) spectral library followed by comparison to reference standards. Multipoint calibrations were prepared from pure standards for 43 VOCs that were common indoor pollutants and/or elevated in one or more of the whole trailer samples. All pure standards and analytes were referenced to an internal standard (~120 ng) of 1-bromo-4-fluorobenzene.

Low Molecular Weight Aldehydes

The target analytes in the aldehyde analysis included formaldehyde, acetaldehyde and acetone. Higher carbon-number aldehydes were quantified using the VOC method described above. Samples of these low molecular weight carbonyl compounds were collected and analyzed following ASTM Test Method D 5197-92 (ASTM, 1997). As with the VOCs, the air samples were drawn directly from the small emission chamber or through a short rigid tube inserted though holes in the trailer door. Samples were collected on commercially available silica gel cartridges coated with 2,4-dinitrophenyl-hydrazine (XPoSure Aldehyde Sampler; Waters corporation). An ozone scrubber (P/N WAT054420; Waters) was installed upstream of the silica cartridge in the field samples. Samples were collected from the trailers for 60 minutes at ~ 1 lpm using personal sampling pumps that were calibrated before use and checked after use. Samples were collected from the laboratory emission chambers using a vacuum pump (Model DOA-P104-AA; Gast) with sample flow rates regulated by electronic mass flow controllers. Sample cartridges were capped and stored on blue ice or in the freezer until extraction.

Cartridges were eluted with 2 mL of high-purity acetonitrile into 2 ml volumetric flasks and the eluent was brought to a final volume of 2 ml before analysis. Extracts were analyzed by high-performance liquid chromatography (HPLC) (1200 Series; Agilent Technologies) using a C_{18} reverse phase column with 65:35 H₂O:Acetonitrile mobile phase at 0.35 ml/minute and UV detection at 360 nm. Multipoint calibrations were prepared for the target aldehydes using commercially available hydrazone derivatives of formaldehyde, acetaldehyde and acetone.

Acetic Acid

Acetic acid was collected in the same way as the carbonyl samples but collected on silica gel sorbent tubes (P/N 22655; SKC) and extracted using 5 mL of 18 mOhm deionized water, filtered through a 0.22 micron membrane. Samples were collected from the trailers for 60 minutes at \sim 1 lpm using personal sampling pumps that were calibrated before use and checked after use. Samples were collected from the emission chambers using a vacuum pump (Model DOA-P104-AA; Gast) with sample flow rates regulated by electronic mass flow controllers. Samples were stored in sealed plastic bags at -15°C until extraction and analysis.

Extracts were analyzed by ion chromatography (IC) (ICS 2000; Dionex) equipped with an autosampler (AS40; Dionex), hydroxide ion generator (EluGen cartridge, P/N 058900; Dionex) and a conductivity detector. Samples were separated on an AS11 column (P/N 044076; Dionex) at a flow rate of 1.0 ml/min. The column was not heated. An injection loop of 25 μ L was used to inject samples. A gradient of hydroxide ions was generated starting at 0.20 mM for 2.3 min. before increasing to 15.00 mM at 12.0 min, then to 35.00 mM at 15.0 min. A multipoint calibration ranging from 0.287 mg/L (of extract) to 52.363 mg/L was prepared from a 1.000g/L acetate ion chromatography standard (P/N 13669; Fluka) and was used to quantify the instrument response. The approximate instrumental limit of quantitation is 0.287 mg/L with a limit of detection of 0.05 mg/L. A typical calibration curve has a relative standard deviation of 4.53% and a coefficient of determination of 99.80%.

Quality Assurance

All samples were quantified with multipoint calibration curves prepared from pure chemicals. For the VOCs that did not have pure standard available or that were a mixture of compounds (i.e., alkylbenzenes), the compounds were tentatively identified by National Institute of Standards and Testing (NIST) library spectrum search and quantified as toluene equivalent values. Analytical blanks were included in all analyses. Trip blanks were prepared, transported to the field sampling site, stored and analyzed along with the whole trailer samples. Method blanks for the small chamber emission experiments including backing plate and tape in the chamber represented more than 10% of all samples collected and chamber blanks representing only the background in the chamber represented an additional 10% of samples collected.

Measurement of Whole Trailer Concentrations

Air concentrations were measured under pseudo steady-state conditions on November 14, 2007 after the THUs had been closed for several days. No attempt was made to control the ambient wind or temperature that the THUs were subjected to during this period. All THUs were setup with samplers and pumps so that all three samples (VOC, aldehyde and acetic acid) could be collected simultaneously in all THUs. A morning sampling event and an afternoon sampling event were conducted for each trailer and at a central outdoor location. The first sample collection started between 11:00 and 11:30 AM and continued for approximately one hour during which time the ambient temperature, relative humidity and wind speed were $25.1 \pm 2.6\%$ (C) and $49 \pm 6.5\%$ (%) and $2.8 \pm 41\%$ (m/s), respectively. The second sampling event started between 14:00 and 14:30 and again lasted about an hour during which time the ambient temperature, relative humidity and wind speed were $26.4 \pm 1.5\%$ (C), $48 \pm 3.2\%$ (%) and $2.6 \pm$ 43% (m/s), respectively. Start and stop times were recorded for each sample along with flow rates. Each sample pump was checked against a calibrated flow meter before and after the sampling event. All samples including two trip blanks for each sample type were sealed and placed on ice for transport back to LBNL. Upon arrival at LBNL the samples were stored in a freezer until analysis.

Measurement of Steady-State Ventilation Rates

The THUs did not include mechanical forced air ventilation systems and operable windows remained closed throughout the study period. Ventilation rates were determined after collection of air samples using a tracer gas decay method. Externally controlled circulation fans were switched on in each trailer and pure carbon dioxide (CO₂) was injected from a Tedlar bag into each unit to achieve an initial concentration that was significantly elevated over ambient conditions. The concentration of CO₂ was measured continuously using Q-Trac Indoor Air Quality monitors through the sample ports in the trailer doors. Mixing fans were run for 15 - 20 minutes after dumping CO₂ into trailers allowing the air concentration of tracer gas was measured.

The ventilation rate is determined from the decay of the tracer gas concentration in the trailer. When using a chemical like CO_2 as a tracer gas, the background level can influence the clearance rates. The equation for decay or clearance of the tracer gas from a trailer after elevating and mixing the CO_2 tracer gas is

$$C_{t} = C_{ss} + \left(C^{*} - C_{ss}\right) \times \exp^{-\mathcal{Q}\left(t-t^{*}\right)}$$

$$\tag{1}$$

where C_t (ppm) is the measured concentration in the unit at time *t*, C* is the maximum at the start of the stable decay period, C_{ss} is the background or ambient concentration, and Q (h⁻¹) is the

rate constant for removal of the tracer from the system, which for a non-reactive chemical that does not significantly interact with surfaces, is the ventilation rate in terms of air changes per hour, ACH (h⁻¹). Equation 1 can be rearranged to the form

$$\ln\left(C_{t}-C_{ss}\right)=-Q\left(t-t^{*}\right)$$
⁽²⁾

so the slope of the natural log of the difference between measured concentration and the ambient concentration against elapsed time is the -ACH as illustrated in Figure 3.

Collection and Characterization of Indoor Materials

The total projected surface area of each material in the trailer was measured and recorded in the field when the material samples were collected for testing after the whole trailer measurements were collected. A representative piece (> 15 cm on a side) of each material was cut directly from the trailer, triple wrapped in foil, placed in a labeled manila envelope and boxed for shipment to LBNL. A total of 45 samples representing 16 different materials were collected from the four trailers. The materials were inventoried upon arrival at LBNL and stored at room temperature in their original packing. Prior to testing, the materials (excluding the fabric and cushions) were cut to size using a dry table saw with sharp blade and returned to their original packing. The fabric and cushion materials were cut to size with a razor or knife. Each material was either pressed into a stainless steel tray to expose only the face or the back was covered with a stainless steel plate and the edges sealed with aluminum tape. When tape was used to seal the edges, the final exposed face was measured and recorded. The individual material samples had already aged in the trailers prior to collection of the test materials so we did not include an additional conditioning period beyond what was required to achieve steady state concentrations in the chambers prior to testing.

Measurement of Material Specific Emission Factors

Material specific emission factors were determined using small chambers generally following the ASTM Standard Guide D-5116-97. Because the goal was to reconstruct whole-trailer emission rates and the trailers were well aged in the field, the individual materials were not conditioned prior to testing. Also, the air-sampling period in the small chambers started after approximately six air changes rather than the recommended 96 hour pre-test period used for new materials. This approach was taken to provide emission factors that were more closely linked to the actual emission rates measured in the whole trailers.

The emission tests were conducted in four 10.5 liter stainless steel chambers that were maintained at 23 ± 1 °C in a controlled environmental chamber with a 0.06 m³/h inlet flow of carbon filtered preconditioned air at 50% ± 5% relative humidity supplied continuously to each test chamber. The relative humidity within the test chambers was controlled by a flow of mixed streams of dry- and water-saturated air. After sealing the backs and raw edges of the material as described above, the materials were placed face up on screens resting slightly below the center of the test chambers. The emitting area, A, (m²) was 0.023, the loading factor, L, (m²/m³) was 2.2 and the area specific flow rate (m³/m²/h) was typically 2.6 for each material. The collection of air samples was initiated after at least six air changes and the VOC, aldehyde and acetic acid samples were all collected from the chamber exhaust stream at a total rate less than 90% of the inlet air stream.

Data Analysis

The whole trailer emission rates normalized to floor area and the material specific emission factors normalized to projected surface area were calculated assuming that the systems (trailer or test chamber) were at pseudo steady-state and were well mixed. The steady-state form of the mass balance equation for calculating area-specific emission rates, *ER*, ($\mu g/m^2/h$) in a well-mixed system is

$$ER = \frac{f \times (C - C_0)}{A} \tag{3}$$

where $f(m^3/h)$ is the ventilation flow rate, $A(m^2)$ is the exposed surface area of the material or the floor area of the whole trailer, $C(\mu g/m^3)$ is the measured steady state concentration in the chamber or trailer and $C_0(\mu g/m^3)$ is the background concentration in the chamber or the outdoor concentration for the whole trailer experiments. Ventilation rate in terms of air flow are not readily available for the whole trailer measurements but given that ACH is equal to the ventilation rate divided by the volume (f/V) and the loading factor is equal to the exposed area divided by the volume, Eq. 3 can be rearranged to give

$$ER = \frac{ACH \times (C - C_0)}{L} \tag{4}$$

where $L (\text{m}^2/\text{m}^3)$ is the loading factor in the chamber or trailer. To relate the material specific emission factors to the whole trailer emission rates we multiply the material specific emission rates by the projected surface area of the material and divide by the floor area of the THU. Normalizing to floor area facilitates comparison among units of different size. To get the floor area normalized emission rate for the whole trailer experiments we note that *ACH* is equal to f/Vas indicated above and that V is the floor area multiplied by the height, h (m) so that Eq. 3 may also be written as

$$ER = ACH \times h \times (C - C_0) \tag{5}$$

for estimating the area normalized emission factors for the whole trailers. The formaldehyde emission rates were compared across trailers and differences between the morning samples and afternoon samples were tested in Excel using the TTest function with two tailed distribution and assuming the samples were of unequal variance. A probability associated with a Student's paired t-Test with a two-tailed distribution less than 0.05 is considered significant.

RESULTS

Material specific loading ratios

The loading ratio for the different composite wood categories in the THUs are compared to the recommended loading ratios in the HUD standard and the ASTM E6007 Standard in Table 5. The recommended loading ratios in the HUD standard are for consistency in the emission measurements and not necessarily intended to represent actual home designs. The loading ratios are calculated from the total amount (surface area) of each composite wood type found in each THU and the approximate internal volume of the THU where volume includes the entire indoor space. No attempt was made to determine readily exchangeable volume where some of the

internal volume of the THUs is taken up by closed cabinet and storage space that may not interact rapidly with the bulk air in the THU so the actual loading ratio of materials in the trailers may be greater than reported in Table 5.

Additionally the ratio of air flow (f) to projected surface area of each wood type in each THU is calculated and compared to the values defined in the HUD standard (Section 408). As with the loading ratios, these airflows are only for standardization and not meant to represent actual conditions in the indoor environment. The air flow is estimated as the product of the internal volume and air exchange rate. Again, no attempt was made to determine readily exchangeable internal volume so the f/A values reported in Table 5 might be biased high, i.e., actual flows are likely to be lower than what is calculated in Table 5.

These calculations show that the loading ratios for Hardwood plywood range from between 2 to 3 times the loading ratio used in the HUD standard for which the concentration limits are established. The ratios of air flow to projected material surface areas in the THUs do not match those used in the standard either. Using HUD compliant hardwood plywood (HWPW) at the loading ratio found in the four different manufactured THUs would be expected to result in a steady-state room concentration 2 to 3 times higher the HUD concentration limit as determined under standardized conditions with all other things being equal.

Whole trailer Ventilation and VOC measurements

When determining ventilation rates, the linear region of the decay curves in the tracer experiment were monitored for approximately 2 hours after the CO_2 concentration had stabilized in each THU. The duration of the decay curves and the correlation coefficient (r^2) from the calculation of *ACH* are included in Table 1. The outdoor CO_2 concentration during the ventilation measurements was 366 ppm \pm 1.6% and the indoor starting concentration for the decay curves were a factor of 9.3, 6.5, 6.8 and 6.6 greater than outdoors for the Dutchmen, Pilgrim, Coachmen and Cavalier, respectively. The minimum tracer concentration indoors relative to outdoor levels at the end of the CO_2 decay period was greater than a factor of 3.4 for all units. The temperature, relative humidity and wind speed (average \pm the percent coefficient of variation (CV)) measured during the two VOC sampling periods and during the tracer gas experiment are summarized in Table 6. Wind speed and indoor/outdoor temperature gradient were similar for the morning (AM) and afternoon (PM) air sampling events. The tracer gas-sampling period had calm wind conditions and the indoor/outdoor temperature gradient was elevated compared to the air sampling times.

In the initial qualitative analysis of VOC samples from the four THUs, approximately 80 individual compounds were tentatively identified in the four different units. Forty-five of the compounds were positively identified and quantified. These target compounds were selected because they were toxicologically important (i.e., benzene) and/or their concentrations were relatively high. Thirty-three of the 45 chemicals that were quantified had steady-state concentrations above $0.4 \ \mu g/m^3$ in one or more of the units. The 33 VOCs are listed in Table 7 sorted by chemical class and increasing boiling point.

A number of higher molecular weight alkyl-benzenes were detected in one THU. These alkylbenzenes had analytical retention times between 36 and 40 minutes in the GC analysis and were combined and quantified as toluene equivalents. The 2,2,4-Trimethyl-1,3-pentanediol diisobutyrate (TMPD-DIB, TXIB) was quantified as 2,2,4-Trimethyl-1,3-pentanediol monisobutyrate (TMPD-MIB, Texanol) although the toluene equivalent quantification gave similar results. The steady-state concentration for each compound in the morning and afternoon samples is given in Table 8 along with the morning and afternoon outdoor concentrations. The indoor concentrations are converted to whole trailer indoor emission rates normalized to the floor area for each unit and presented in Table 9.

Material Specific VOC measurements

All THUs had a significant fraction of the internal surface area (walls, ceiling, cabinet walls) constructed of 1/8-inch plywood with a vinyl or PVC skin or simulated wood finish. All units had sheet vinyl flooring while the Dutchmen and Pilgrim also had carpeted areas. All countertops were particleboard surfaced with high-pressure laminate. A variety of wood products were used for the sub-floor and for the bench and bed platforms.

Material specific emission factors were measured for the same target chemicals that were identified in the whole trailer measurements. The emission factors for each material are first summarized by individual THU in Tables 10 through 13. These values are converted to whole trailer emission factors for each THU by multiplying the measured emission factor by the projected surface area for each material in the trailer then dividing by the total floor area of the trailer. These results are presented in Tables 14 through 17.

Percent Contribution of Material Specific Emissions to Whole Trailer Measurements

The values in Table 14 through Table 17 are transformed to the approximate percent contribution to total pollutant load in each THU and reported in Tables 18 through 21. The chemicals are sorted in order of decreasing emission for each THU. As an example, the total emissions of formaldehyde from all materials in the Pilgrim is reported in Table 19 as 493 ($\mu g m^{-2} h^{-1}$) with 55% of the emission stream from "cabinet" material, 14% from walls, 8% from seat cushion and small fractions from curtain, bed deck, carpet, and ceiling. These percentages should be treated as approximations given the limited number of samples tested and the differences between the test conditions and the actual whole trailer conditions. In addition, the results cannot distinguish between primary sources and secondary sources that are re-emitting chemicals that have been sorbed over time in the THU.

The total material specific emission factors across all materials normalized to the THU floor area are compared to the average of the two field measurements for the whole trailer emission factors for each THU in Table 22. These results further illustrate that the dynamics in the whole THU likely suppress emissions from the primary sources given the long-term mixing of pollutants among the indoor sources and competitive emissions in the whole trailer that do not exist in the small chamber experiments with individual materials.

DISCUSSION

Building material emission measurements for formaldehyde and other VOCs have been published in the literature over recent years. These emission factors may be used for comparison to those measured in the THUs. However, it is important to note that both the whole THU emission rates and the material specific measurements represent materials that have been exposed to the entire mixture of VOCs in the indoor environment for the life of the THU. The values from the chamber experiments and the whole trailer measurements do not necessarily reflect primary emissions that are typically measured in standard protocols where new, unexposed materials are tested after a specific aging period. In addition, the conditions used to collect the whole trailer measurements represented a lower bound ventilation scenario because the units had been closed for several days prior to testing. Nevertheless, we can tentatively identify likely sources of the VOCs in the THUs based on other studies in combination with the material-specific measurements from this study.

VOC Emissions from Building Materials

Hodgson et. al. (2000) measured VOC concentrations under pseudo-steady state conditions in four new manufactured homes and seven new site-built houses. The geometric mean (GM) concentrations (ppb) are reported for each housing type along with geometric standard deviation (GSD). Hipelein (2004) measured indoor air VOC concentrations in 79 rooms in 39 private dwellings in Germany. The homes were not associated with health complaints but 27% of the rooms investigated were occupied by smokers. No information is provided about the ages of the dwellings but the objective of the study indicates that the homes were representative of background conditions so we assume the values represent the existing housing stock. The concentrations (ug/m³) were reported for a number of VOCs in the original manuscript as the GM. These values are transformed to units of ppb using conversion factors reported by Hodgson and Levin (2003). Approximately half of the chemicals that were quantified in the THUs (17 of 33) had concentration values reported for the new manufactured and site built homes and/or the German residences. These measurements are compared in Figure 4. The results from Hipelein (2004) are generally lower than the values reported for the new dwellings by Hodgson et. al. (2000). Although many of the VOCs measured in the THUs are similar to reported residential values, several are in excess. Dodecane, tridecane, p-xylene, alpha-pinene, beta-pinene and hexanal are all above values reported in the German homes while phenol, TMPD-DIB, acetic acid and formaldehyde are even in excess of values measured in the new dwellings.

A number of the other chemicals that were measured in the THUs but did not have literature values for comparison were also elevated as compared to the values that are included in Figure 4. The concentrations of four chemicals (styrene, tetradecane, pentadecane and the alkylbenzenes) were greater than the median concentration of the chemicals reported for the THUs in Figure 4. Another four (2-ethyl-1-hexanol, benzaldehyde, hexadecane and dimethyl methylphosphonate) were greater than the 25th percentile of the THU data reported in Figure 4. And all but two chemicals had measured concentrations in at least one unit that was greater than the 5th percentile of the values reported in Figure 4.

In addition to reporting indoor concentrations of VOCs, Hodgson et.al., (2000) also report whole unit emission rates normalized to floor area. These values for whole building emission factors in new site-built and manufactured houses are compared to the VOC emission factors measured in the four THUs in Figure 5. As with the concentration data described previously, the compounds presented for emission factor comparison were selected from the available data because they were included in both the THUs studied and in the Hodgson et.al., (2000) paper. These compounds represent a wide range of functional groups including terpenes, alcohols, ester alcohols, aldehydes, and organic acids. Two of the compounds (styrene and acetaldehyde) had similar emission factors in the THUs were lower than from the newly constructed dwellings for twelve of the eighteen compounds compared. All of the alkane and terpene compound emissions were lower in the THUs than in the new houses, as were TMPD-MIB, and most of the aldehydes. The lower emission factors in the THUs may be due to aging where the four THUs tested were more than 1 ½ years old while the site-built and manufactured homes were all approximately 6 months

old or less. In contrast, the emission factors for phenol, TMPD-DIB, acetic acid and formaldehyde remained higher in the THUs than the new homes. In particular, the median whole trailer formaldehyde emissions from the four units was more than five times greater than the values reported for new manufactured and site build residences.

These results provide a general focus for discussion of the VOC and aldehyde emissions within the THUs studied. The measured emissions of the ester alcohol TMPD-DIB are large, likely due to the relatively large amounts of vinyl flooring and other vinyl materials in the THUs. A large number of individual materials were found to emit TMPD-DIB in the small chamber experiments and we were not able to distinguish between primary emitters and secondary emission of sorbed chemical. Interestingly the Coachmen had far lower whole trailer TMPD-DIB emissions than the other three units; possibly due to the lower emissions of the compound from the vinyl floor in that THU. Elevated levels of the high molecular weight alkyl-benzenes in that THU indicates the use of these chemicals in place of TMPD-DIB in some vinyl flooring.

The acetic acid concentrations/emissions were also large in the THUs and the material specific measurements indicate the source of acetic acid was predominantly the subfloor but some of the units also had bed deck and bench seat materials that emitted high levels. Again, the experiments were not designed to distinguish between primary and secondary emission sources but secondary sources are not expected to be as pronounced for the more volatile chemicals like acetic acid as compared to the lower volatility chemicals like TMPD-DIB.

The fact that the levels summed across all materials exceeded that which was measured in the whole THUs for a number of the chemicals is likely an indication of secondary emission sources where chemicals released from the primary source in the whole trailer are being sorbed into/onto other materials in the THUs. This can occur where some of the surfaces act as sinks and secondary re-emission sources that compete with the primary emission source of any individual chemical and material. For example, if a material emits a particular chemical into the indoor environment, that chemical can partition into other materials until it approaches a steady-state concentration that is a function of the thermodynamic equilibrium partition coefficient between the material and the overlying air. These sorbed chemicals can be released back into the air as secondary emission sources when the primary source is removed or when conditions in the trailer change such as an increase in the ventilation rate or change in temperature.

Aldehyde Emissions from Building Materials

Hodgson et al. (2002) measured material specific emissions of aldehydes and terpenes for a single new manufactured house. The study selected materials from a newly constructed modular home and collected the materials direct from the factory that fabricated the dwelling. The new materials were tested in small emission chambers to determine material specific emission factors. Indoor house measurements were also collected in the newly manufactured home and the material emission factors were used to reconstruct whole house emission rates. Table 23 and Table 24 provide the aldehyde and terpene emissions, respectively, from material samples. This work used fresh materials not previously used in the house so the measurements represented primary emissions. The study, along with an earlier report (Hodgson, 1999) identified composite woods made with urea-formaldehyde resin as important formaldehyde and terpene hydrocarbon sources in buildings.

The State of California has conducted studies and has initiated various programs and regulations intended to lower material emissions of formaldehyde since the California Air Resource Board

(CARB) identified the compound as a Toxic Air Contaminant (TAC) in 1992 (CalEPA, 1992). Part of this effort included a survey of emissions from composite wood products on the market in California, conducted by Battelle Labs during 1995 (Battelle 1996; Kelly et al. 1999). The results, which are summarized by CARB (CARB 2008) and reproduced in Table 25, also indicate that composite wood material are the main contributors to formaldehyde emissions from building materials.

HUD Standard 24 CFR Ch. XX Part 3280, *Manufactured Home Construction and Safety Standards* (HUD 2001) specifies test chamber concentrations (*Css*, ppm) for plywood and particle board of 0.2 and 0.3 formaldehyde, respectively, based on the ASTM emission testing method E-1333. The standard specifies a modified loading (L, m²/m³) ratio of 0.95 and 0.43 for plywood and particle board, respectively, and an air exchange rate of 0.5 h⁻¹ for the test. Using these values with the equation specified in ASTM E-1333 (Eq. 6) we can estimate an initial emission rate (*ER*, mg/m²/h) from these materials as

$$ER = 1.23 \times Css \times ACH/L$$
 Eq. 6

The calculation results in an initial ER for plywood and particle board of 130 μ g/m²/h and 430 μ g/m²/h, respectively.

These numbers represent fresh material. As discussed in the introduction, emission rates are expected to decrease as the materials age. If we use the slower decay rate discussed in the introduction based on the work presented by Mathews (1985), the expected emission rates for materials that have aged more than 1 $\frac{1}{2}$ years would be approximately 65 μ g/m²/h and 220 $\mu g/m^2/h$ for the plywood and particle board, respectively. Comparing the material specific emissions to the values for fresh and aged material, none of the materials in the Dutchman exceeded either the aged or fresh emission factors. The bed deck (oriented strand board) in the Pilgrim had an emission rate of 136 μ g m⁻² h⁻¹ which exceeded the fresh plywood standard but was below the criteria for particle board (either fresh or aged). However, the plywood cabinet wall in the Pilgrim (419 μ g m⁻² h⁻¹) exceeded even the fresh emission criteria. Two plywood materials in the Coachman exceeded the fresh material criteria (bench seat at 233 μ g m⁻² h⁻¹ and cabinet wall at 174 μ g m⁻² h⁻¹). One material in the Caviler (plywood cabinet wall at 490 μ g m⁻² h^{-1}) exceeded the fresh material criteria and two more exceeded that for the aged material (particle board seat bottom at 292 μ g m⁻² h⁻¹ and particle board sub floor at 416 μ g m⁻² h⁻¹). It is unclear whether the subfloor, which is covered with vinyl or carpet, contributes significantly to the indoor air concentrations. Several materials that do not have test criteria were also found to emit relatively high levels of formaldehyde. These include the seat cushion in the Duchman (70 μ g m⁻² h⁻¹), the curtain and seat cushion in the Pilgrim (323 μ g m⁻² h⁻¹ and 409 μ g m⁻² h⁻¹ respectively), and the seat cushion and walls in the Coachman (151 μ g m⁻² h⁻¹ and 60 μ g m⁻² h⁻¹. respectively) although it is unclear whether these are secondary emission or primary sources.

The whole building formaldehyde emissions in previously studied site-built and manufactured homes were 31 and 45 μ g m⁻² h⁻¹, respectively (Figure 5 and Hodgson 2000). These values are up to an order of magnitude lower than those measured in the THUs. Whole THU formaldehyde emissions (per floor area) ranged from 173 to 266 μ g m⁻² h⁻¹ in the AM and 257 to 347 μ g m⁻² h⁻¹ in the PM. When the THUs are occupied, differences in ventilation rates, temperatures, relative humidity and indoor air movement may influence the steady-state concentration of VOCs and

emission rates but information is currently lacking on the magnitude and direction of this influence.

Thus, the whole trailer formaldehyde emission factors are high. Many of the material specific formaldehyde emission factors for composite wood materials used in the THUs appear to be within the range reported for fresh materials in previously published research (Tables 23 - 25), but significant differences can be expected due to the aging of the material in the THUs. It is difficult to conclude that the materials would have been within previously reported ranges had the tests been conducted using fresh materials but a number of the materials in the THUs are higher than would be expected even if a slow emission decay rate is assumed for aging. If the materials used in the THUs were in fact within the acceptable range for emission when new, then the results from the whole trailer measurements indicate a difference in the construction/design that may lead to elevated concentrations and whole trailer emission rates.

Three features of material application in the THUs may lead to elevated whole trailer concentrations relative to expected concentrations based on test criteria. These include 1) the extensive use of lightweight composite wood products, 2) high surface loading of composite wood products and 3) low fresh air per unit surface area of composite wood products in the THUs. Much of the projected surface area in the THUs (wall, ceiling, and cabinetry) use 1/8" plywood. Although the formaldehyde emission from these materials were generally low, the large surface area can results in significant contributions to overall emissions (e.g., the Coachmen). Given that almost all surfaces in these structures are wood, the wood product loading factor of the THU is far higher than in housing that uses gypsum for walls and ceilings. In addition, the surface area of cabinetry relative to volume of residence is high. Considering this in terms of the area-specific clean air flow rates, the high material loading ratio in the units combined with relatively low fresh air ventilation rates results in area-specific air flow rates that are quite low relative to other housing types. With all other factors being equal, the steady-state concentrations indoors are inversely proportional to the air exchange rates as indicated in Eq. 4. It is unclear how the low ventilation relative to surface area will affect the measured formaldehyde emission rates presented in this report, but these factors can influence the exposure concentrations experienced by occupants.

CONCLUSIONS

The results of this study are not statistically representative of the entire fleet of FEMA THUs because the study was based on a convenience sample of four THUs. Nonetheless, the measured material-specific emission factors for volatile organic compounds, including formaldehyde, were not atypical relative to the literature for new materials. However, it is important to consider that the materials in this study were both aged and allowed to interact with emissions from other materials. Formaldehyde and some of the other VOCs measured in the unoccupied THUs and the associated whole trailer emission factors were found to be higher, sometimes much higher, than what is typically found in residential environments. The difference between these THUs and other housing appears to be the very high composite wood surface area relative to room volume and the low ventilation rates in terms of low area-specific fresh air flow rates relative to internal surface area in the THUs.

Recommendations for future work

This report provides a preliminary assessment into the effect of THU design and material choices on indoor VOC and aldehyde concentrations. It is by no means definitive as a convenience sample of only four THU models produced by four manufactures was evaluated. Additionally, the focus of this study was on the travel trailers, while a significant portion of THUs are park trailer models, and manufactured homes. A systematic assessment across a wider range of THU makes and models including a better characterization of fresh air ventilation rates under occupied conditions could provide a better understanding of the time integrated exposure concentrations in occupied units.

The results in this report do not yet address temperature and humidity effects on material emissions within the studied units. It is hypothesized that at the higher temperature and relative humidity conditions found in the summertime in the southeastern portions of the US, emissions of formaldehyde from the urea-formaldehyde composite woods will increase. Chamber experiments and a seasonal study designed to investigate the potential effects of temperature and humidity should be completed.

Recommendations for formaldehyde mitigation approaches for the THUs have not been provided; nonetheless, the results of this study may be useful to help identify mitigation strategies for reducing VOCs including formaldehyde in THUs. An assessment of the literature for information on the effectiveness of material coverings should be considered. As part of this effort to explore the influence of diffusion resistance at the material surface, the resistance to diffusion on the air/surface interfaces as influenced by airflow and boundary layer effects should be considered. It would be informative to explore the effect of material aging and the role of different material types as surface sinks and sources of secondary emissions of indoor pollutants and how this impacts the primary emission source material.

This study was not intended to assess health effects that may be related to the measured emission factors and whole trailer concentrations in the THUs. The experimental conditions were not necessarily representative of occupied THUs. An estimate of the time-history concentration of each chemical along with time activity patterns of the THU occupants would be needed to characterize exposures in a way that is relevant to health effects assessments.

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TABLES:

	Trailer				
Manufacturer	Thor Industries	International	Spirit of America	Gulfstream	
Model	Dutchmen	Pilgrim	Coachmen	Coach Cavalier	
VIN	47CTDER256G520549	5L4TF332563013658	1TC2B970861308196	1NL1VTR2661064665	
Manufactured	Sep 2005	Oct 2005	Oct 2006	Mar 2006	
FEMA Specs.	No	Yes	No	Yes	
Floor area (m ²)	20.2	20.3	22.4	19.9	
Internal height (m)	2.08	1.98	2.06	1.98	
Internal Volume (m ³)	41.9	40.3	46.2	37.9	
Ventilation					
Characteristics					
ACH ^a (h ⁻¹)	0.25	0.15	0.39	0.21	
Duration of linear tracer					
decay (min)	153	135	142	118	
r ² for linear region of					
tracer decay	0.998	0.999	0.998	0.998	
Apparent Air Flow ^b (m ³ /h)	10.5	6.0	18.0	8.0	

Table 1. Specifications and Ventilation Characteristics of the Temporary Housing Units

^a ACH, air changes per hour measured after the final sampling event of the day. ^b Apparent Air Flow is the product of internal volume and ACH and represents the fresh air flow through the THU

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Material	Trailer				
	Dutchmen	Pilgrim	Coachmen	Cavalier	
Ceiling	23.6	19.4	24.1	19.0	
Walls	60.1 ^ª	40.7	63.1	60.5	
Subfloor	23.6	20.3	22.4	19.9	
Carpet	8.3	7.4			
Vinyl floor	17.7	13.0	22.4	19.9	
Cabinet Walls	30.0	13.2	6.9	17.8	
Cabinet Ends	2.6		0.9	0.16	
Countertop	2.72	1.56	1.79	1.14	
Interior Door	2.79	2.04	0.98	1.86	
Exterior Door	1.02	1.02	1.02	1.02	
Trim board	1.18				
Fabric	7.18	6.84	6.58	7.04	
Fabric Divider			3.40		
Bed Platform MDF	6.09			6.78	
Bed Platform Plywood			5.42		
Bed Platform OSB		3.89	2.70		
Tub surround	3.20	3.24	3.20	3.74	
Windows	5.44	1.76 ^b	2.55	1.76	
Vinyl seat ^c	2.50	2.01	1.96	2.06	

Table 2. Projected Surface Area of Indoor Materials (m²)

^a numbers written in bold text indicate that the material was included in set for determination of emission factors; ^b window area used to represent fabric curtain material; ^c vinyl seat areas used to represent projected surface area of seat/cushion material

	Trailer			
	Dutchmen	Pilgrim	Coachmen	Cavalier
ceiling	1/8 inch plywood	1/8 inch plywood	1/8 inch plywood	1/8 inch plywood
wall	1/8 inch plywood	1/8 inch plywood	1/8 inch plywood	1/8 inch plywood
sub floor	5/8 inch plywood	5/8 inch plywood	9/16 inch OSB ^a	5/8 particle board/OSB finer fiber and darker resin
carpet	Low pile with backing	Low pile with backing		
vinyl floor	vinyl with slight residue of glue on back	vinyl does not have indication of glue	vinyl does not have indication of glue	vinyl does not have indication of glue
cabinet wall	1/8 inch plywood	1/8 inch plywood	1/8 inch plywood	1/8 inch plywood
cabinet wall thick	$\frac{1}{2}$ inch HB ^b or MDF ^d			
cabinet door				1/2 inch HB or MDF
countertop	5/8 inch PB ^c	5/8 inch PB	½ inch PB	5/8 inch PB
door	Hollow core 1/8 inch HB panels (1 smooth back, 1 textured back) with cardboard fill	Hollow core 1/8 inch HB panels (both textured back) with cardboard fill	Hollow core 1/8 inch HB panels (both smooth back) with cardboard fill	Hollow core 1/8 inch HB panels (1 smooth back, 1 textured back) with cardboard fill
trim	3/8 inch MDF			
curtain/door fabric			fabric	plastic impregnated fabric
seat cushion	fiber fill material (white) with fabric cover	Polyurethane foam dense and light in color covered with 2 layers plastic film and fabric	Polyurethane foam dense and light in color covered with simulated fiber, 2 layers plastic film and fabric	fiber fill material (white) with fabric cover
seat bottom/bed platform	3/8 inch PB	3/8 inch OSB	1/2 inch OSB	3/8 inch MDF
bench-seat bunk bed platform			3/8 inch plywood	
bench end			1/2 inch MDF	

Table 3. Description of Surface Materials Harvested from Trailers and Tested for Emissions

^a OSB, oriented strand board; ^b HB, hardboard or high density fiber board; ^c PB, particle board; ^d MDF, medium density fiber board

	Trailer				
Material	Dutchmen	Pilgrim	Coachmen	Cavalier	
ceiling	textured white vinyl finish with unfinished veneer backing	textured white vinyl finish with unfinished veneer backing	textured white vinyl finish with unfinished veneer backing	textured white vinyl finish with unfinished veneer backing	
wall	vinyl laminant with unfinished veneer backing	vinyl laminant with unfinished veneer backing	vinyl laminant with unfinished veneer backing	vinyl laminant with unfinished veneer backing	
sub floor carpet	unfinished low pile	unfinished low pile	unfinished	unfinished	
vinyl floor	vinyl	vinyl	vinyl	vinyl	
cabinet wall	simulated wood photo laminate front with veneer backing simulated wood	simulated wood photo laminate front with veneer backing	simulated wood photo laminate front with veneer backing	simulated wood photo laminate front with veneer backing	
	photo laminate finish both sides				
cabinet door				simulated wood photo laminate finish both sides	
countertop	HP Laminate with backing covered with a slightly thicker layer of Formica	HP Laminate with backing cover of dense brown paper.	HP Laminate with backing cover of dense brown paper.	HP Laminate with backing cover of dense brown paper.	
door	Simulated wood photo laminate each side (oak & maple)	Simulated wood photo laminate (maple)	Simulated wood photo laminate (oak)	Simulated wood photo laminate each side (oak & maple)	
trim	simulated wood photo laminate front and sides, back unfinished				
curtain/door fabric		loose weave polyester fabric	loose weave fabric pleated and impregnated with plastic		
seat cushion	fabric with vinyl material for back (vinyl not tested)	Fabric with vinyl material for back (vinyl not tested)	Fabric with vinyl material for back (vinyl not tested)	fabric with vinyl material for back (vinyl not tested)	
seat bottom/bed platform	simulated wood photo laminant on one surface and unfinished on back	unfinished	unfinished	unfinished	
bench-seat bunk bed platform bench end			unfinished veneer both sides simulated wood photo laminate		

Table 4. Surface Coverings and Finishes on Tested Materials

Hardwood Plywood (HWPW)	Dutchmen	Pilgrim	Coachmen	Coach Cavalier
HWPW Surface Area (m ²)	137	94	99	97
HWPW Loading Ratio (m ² /m ³)	3.28	2.33	2.15	2.46
ASTM E1333 Loading Ratio (m ² /m ³)	0.95	0.95	0.95	0.95
HWPW flow/area (m/h)	1.22	2.87	1.19	1.93
ASTM E1333 flow/area (m/h)	0.53	0.53	0.53	0.53
Particleboard (PB)				
PB Surface Area (m ²)	5	2	2	21
PB Loading Ratio (m ² /m ³)	0.12	0.04	0.04	0.53
ASTM E1333 Loading Ratio (m ² /m ³)	0.43	0.43	0.43	0.43
PB flow/area (m/h)	32	172	66	9
ASTM E1333 PB flow/area (m/h)	1.17	1.17	1.17	1.17
Medium Density Fiberboard (MDF) and Hardboard (HB)				
MDF,HB surface Area (m ²)	10	0	2	9
MDF, HB Loading Ratio (m ² /m ³)	0.24	0.00	0.04	0.23
ASTM E1333 Loading Ratio (m ² /m ³)	0.26	0.26	0.26	0.26
MDF flow/area (m/h)	17	0	60	21
ASTM E1333 MDF flow/area (m/h)	1.91	1.91	1.91	1.91
Particleboard Door Core				
Door Core Surface Area (m ²)	4	3	2	3
Door Core Loading Ratio (m ² /m ³)	0.09	0.08	0.04	0.07
ASTM 6007 Loading Ratio (m ² /m ³)	0.13	0.13	0.13	0.13
Door Core flow/area (m/h)	44.0	87.7	59.2	65.3
ASTM 6007 flow/area Ratio (m/h)	3.85	3.85	3.85	3.85

Table 5. Surface Loading Ratios and Area-Specific Clean Air Flow Rates

	Temperature (C)	RH	Wind speed (m/s)
AM sample period			
(11:05 – 12:23)			
Dutchmen	24.3 ± 1.1%	55 ± 1.2%	
Pilgrim	22.8 ± 1.1%	55 ± 2.9%	
Coachmen	25.1 ± 1.0%	55 ± 1.6%	
Cavalier	21.9 ± 1.1%	58 ± 2.1%	
Outdoors	25.1 ± 2.6%	49 ± 6.5%	2.8 ± 41%
PM sample period			
(14:00 – 15:20)			
Dutchmen	28.4 ± 1.0%	48 ± 4.4%	
Pilgrim	27.2 ± 1.6%	46 ± 7.9%	
Coachmen	29.6 ± 2.6%	46 ± 8.9%	
Cavalier	25.7 ± 1.9%	49 ± 10%	
Outdoors	26.4 ± 1.5%	48 ± 3.2%	2.6 ± 43%
Tracer gas sample period			
(16:00 – 19:00)			
Dutchmen	26.8 ± 2.3%	55 ± 3.6%	
Pilgrim	25.6 ± 2.9%	55 ± 4.7%	
Coachmen	26.1 ± 2.6%	59 ± 3.8%	
Cavalier	25.0 ± 2.7%	66 ± 3.9%	
Outdoors	21.6 ± 4.2%	80 ± 10%	0.8 ± 50%

Target Compound	CAS#	Chemical Class ^e	BP (°C)
Acetic acid ^a	64-19-7	Acid	118
Phenol	108-95-2	Alc	182
2-ethyl-1-hexanol	104-76-7	Alc	183
Formaldehyde ^b	50-00-0	Ald	-19
Acetaldehyde ^b	75-07-0	Ald	20
Hexanal	66-25-1	Ald	128
Octanal	124-13-0	Ald	174
Benzaldehyde	100-52-7	Ald	179
Nonanal	124-19-6	Ald	195
Dodecane	112-40-3	Alka	216
Tridecane	629-50-5	Alka	236
Tetradecane	629-59-4	Alka	252
Pentadecane	629-62-9	Alka	270
Hexadecane	544-76-3	Alka	287
Benzene	71-43-2	Arom	80
Toluene	108-88-3	Arom	111
Ethylbenzene	100-41-4	Arom	136
p-Xylene	106-42-3	Arom	139
Styrene	100-42-5	Arom	145
Propyl-benzene	103-65-1	Arom	159
1,3,5-trimethyl-benzene	108-67-8	Arom	165
1,2,3-trimethyl-benzene	526-73-8	Arom	175
AlkylBenzenes (36 min - 40 min) ^c		Arom	
TMPD-MIB ^d	25265-77-4	Estr	244
TMPD-DIB	6846-50-0	Estr	280
Acetophenone	98-86-2	Ket	202
Hexamethyl cyclotrisiloxane,	541-05-9	Misc	134
Octamethyl cyclotetrasiloxane,	541-02-6	Misc	175
Dimethyl methylphosphonate	756-79-6	OP	181
α-Pinene	7785-70-8	Terp	155
3-Carene	13466-78-9	Terp	165
β-Pinene	18172-67-3	Terp	166
d-Limonene	5989-27-5	Terp	177

^a acetic acid originally qualitatively assessed by GCMS as toluene equivalents then quantified by ion chromatography (IC); ^b low molecular weight aldehydes were analyzed by HPLC; ^c the series of alkyl-benzenes eluting between 36 and 40 minutes are combined and quantified as toluene equivalents; ^d TMPD-DIB was quantified as TMPD-MIB (Texanol); ^e the definition of chemical classes are Alc = alcohol, Ald = aldehyde, Alka = alkane, Arom = aromatic chemical, Estr = ester, Ket = Ketone, Misc. = miscellaneous, OP = organophosphate and Terp = terpene.

	Out	Outdoors	Duc	Duchmen	piiq	Pilgrim	Coac	Coachmen	Cav	Cavalier
Target Compound	AM	M	AM	Mq	AM	Md	AM	Mq	AM	M
Acetic acid (GCMS as toluene) ^a			69.2	105	89.6	153	10.4	22.9	88.8	121
Acetic acid (IC method)	44.9	43.6	939	1080	1250	1430	577	553	1090	1260
Phenol	5.76	4.29	24.5	36.1	40.2	58.6	18.3	23.7	31.8	49.7
2-ethyl-1-hexanol	0.91	0:30	1.38	2.70	2.00	4.08		0.49	4.77	7.85
Formaldehyde	2.06	3.34	378	487	601	926	331	433	632	764
Acetaldehyde	2.03	2.32	12.1	12.6	13.3	15.3	7.18	6.16	10.06	9.28
Hexanal		0.23	12.5	18.4	22.7	31.3	7.42	7.79	34.8	44.3
Octanal	0.29	0.40	4.93	7.51	5.22	7.67	1.97	2.63	6.39	8.79
Benzaldehyde	5.79	4.30	1.62	2.80		1.39			1.52	3.70
Nonanal	0.37	0.52	6.97	12.4	9.86	11.88	5.23	6.84	9.48	16.1
Dodecane	0.04		0.27	0.42	11.2	15.1	0.33		1.26	1.89
Tridecane	0.10	0.04	11.3	23.7	132	178		1.24	40.7	60.8
Tetradecane	0.38	0.25	25.7	37.8	95.2	125	30.9	41.1	38.2	56.9
Pentadecane	0.31	0.20	5.99	9.49	8.35	12.1	20.4	27.0	5.56	9.26
Hexadecane			1.50	2.29	1.97	3.38	4.40	6.11	1.82	3.08
Benzene	0.68	0.64			0.13					
Toluene	0.16	0.20	2.79	1.57	1.23	1.04	1.15	0.50	1.46	1.33
Ethylbenzene		0.06	0.37	0.20	0.43	0.37	0.31	0.15	0.61	09.0
p-Xylene		0.07	0.40	0.29	0.44	0.37	0.43	0.23	0.35	0.28
Styrene	0.02	0.01	9.16	4.15	14.4	11.2	4.52	1.56	20.1	17.2
Propyl-benzene							0.56			
1,3,5-trimethyl-benzene			0.09	0.13	0.45	0.55	0.09			
1,2,3-trimethyI-benzene				0.21	1.01	1.25				
AlkylBenzenes (36 min - 40 min)							184	242		
TMPD-MIB	0.96	0.92	5.36	8.13	7.09	10.2	1.58	1.53	17.7	26.7
TMPD-DIB	4.15	5.19	406	519	307	390	9.59	14.2	429	565
Acetophenone	5.77	4.35								
Hexamethyl-cyclotrisilox ane		0.04	0.38	0.28	0.83	0.74	0.82	0.85	1.95	0.84
Octamethyl-cyclotetrasiloxane	0.16	0.12	0.06	0.05	0.13	0.24	0.19	0.18	0.35	0.21
Dimethyl methylphosphonate									8.84	5.00
α-Pinene	0.53	0.49	90.4	82.2	103	103	28.7	19.4	69.1	73.0
3-Carene			2.09	2.17	4.91	5.59	3.28	2.72	9.98	11.1
β-Pinene	0.28	0.29	9.54	9.40	13.9	14.9	3.78	2.84	10.8	11.8
d imperator			200	000	2 74	A 50	1 10	70.0	200	44

Table 8. Measured Steady-state VOC Concentrations ($\mu g/m^3$) in Field Samples

^a Acetic acid was initially identified and tentatively quantified as toluene during the GCMS analysis of <u>VOCs</u>. These values are much lower than the concentrations determined using the IC method because of chromatographic peak broadening on the GC column. The results are include for comparison with the IC result but only the IC results are used in subsequent data analysis and emission factor estimations.

				Trai	lers			
-	Duch	men	Pilg	rim	Coacl	nmen	Cav	/alier
Target Compound	AM	PM	AM	PM	AM	PM	AM	РМ
Acetic acid (IC method)	405	467	287	330	336	321	353	413
Phenol	12.9	19.1	11.6	16.9	14.7	19.0	12.6	19.7
2-ethyl-1-hexanol	0.73	1.43	0.58	1.18		0.39	1.89	3.11
Formaldehyde	200	257	174	267	266	347	261	315
Acetaldehyde	6.38	16.2	3.83	9.62	5.77	15.5	4.16	12.4
Hexanal	6.59	9.70	6.56	9.03	5.96	6.26	13.8	17.6
Octanal	2.60	3.97	1.51	2.21	1.58	2.11	2.53	3.48
Benzaldehyde	0.86	1.48		0.40			0.60	1.46
Nonanal	3.69	6.56	2.85	3.43	4.20	5.49	3.76	6.38
Dodecane	0.14	0.22	3.23	4.36	0.27		0.50	0.75
Tridecane	5.96	12.5	38.2	51.3		1.00	16.1	24.1
Tetradecane	13.6	20.0	27.5	36.1	24.8	33.0	15.1	22.5
Pentadecane	3.17	5.02	2.41	3.50	16.4	21.7	2.20	3.67
Hexadecane	0.79	1.21	0.57	0.98	3.53	4.91	0.72	1.22
Benzene			0.04					
Toluene	1.47	0.83	0.35	0.30	0.92	0.41	0.58	0.53
Ethylbenzene	0.19	0.10	0.13	0.11	0.25	0.12	0.24	0.24
p-Xylene	0.21	0.16	0.13	0.11	0.35	0.18	0.14	0.11
Styrene	4.84	2.19	4.15	3.23	3.63	1.25	7.95	6.80
Propyl-benzene					0.45			
1,3,5-trimethyl-benzene	0.05	0.07	0.13	0.16	0.07			
1,2,3-trimethyl-benzene		0.11	0.29	0.36				
AlkylBenzenes (36 min - 40 min)					148	195		
TMPD-MIB	2.84	4.30	2.05	2.96	1.27	1.23	7.00	10.6
TMPD-DIB	215	275	88.7	113	7.70	11.4	170	224
Acetophenone								
Hexamethyl-cyclotrisiloxane	0.20	0.15	0.24	0.21	0.66	0.68	0.77	0.33
Octamethyl-cyclotetrasiloxane	0.03	0.03	0.04	0.07	0.16	0.15	0.14	0.08
Dimethyl methylphosphonate							3.50	1.98
α-Pinene	47.8	43.4	29.8	29.7	23.1	15.6	27.4	28.9
3-Carene	1.10	1.15	1.42	1.62	2.63	2.19	3.95	4.38
β-Pinene	5.04	4.97	4.01	4.31	3.03	2.28	4.27	4.65
d-Limonene	1.63	1.70	1.07	1.31	0.90	0.70	2.01	2.44

Table 9. Whole Trailer Emission Rates Normalized to Floor Area (µg m⁻² h⁻¹)

Target Compound	bed	cabinet	cabinet	carpet	ceiling	counter	door	seat	sub	trim	vinyl	wall
Acatic acid	Nan	Wall	eria			do	ILICELIOL	CUSHIOL				
Acelic acid				0.00				0.00	999			
Phenol	0.93	0.74	0.28	32.2	5.70	0.69	4.48	33.8	3.03	0.48	25.5	4.42
2-ethyl-1-hexanol	0.56		0.77	0.99	0.87		0.24	9.19	0.98	0.48	1.47	0.85
Formaldehyde	4.11	17.7	5.21	42.4		11.2	45.6	69.2	4.31	11.0		10.9
Acetaldehyde				4.98		2.34	0.12	6.13	11.4	4.69		
Hexanal	0.85		0.47	10.9	0.59	0.74	1.86	1.08	8.69	0.83	3.22	2.21
Octanal				8.18								
Benzaldehyde	0.60	0.47		24.0	0.28	1.79	0.23	2.35	0.26		1.02	0.82
Nonanal	1.53	0.05	6.84	13.2	1.90	0.73	1.40	25.2	2.63		3.84	10.8
Dodecane									0.86		0.93	
Tridecane	0.25		0.86	16.5	1.07		3.33	10.8	16.8		37.7	0.92
Tetradecane	0.36			44.3	6.57		8.34	31.3	45.2	0.69	96.4	2.12
Pentadecane				9.84	3.02		2.87	9.08	37.0		78.5	1.02
Hexadecane	0:30	0.31		1.93	1.37		1.63	2.00	6.86		12.7	0.41
Benzene	0.42		0.25	1.18		0.28	0.13	0.40		0.06		1.70
Toluene	1.36	0.74	0.81	2.26	1.42	0.69	0.80	0.09	2.46	0.73	0.61	1.16
Ethylbenzene				0.59			0.01	0.10				
p-Xylene				1.41			0.03	0.35				0.59
Styrene				0.69			0.02	0.15				
Propyl-benzene							0.03	0.08				
1,3,5-trimethyl-benzene				0.17				0.14				
1,2,3-trimethyl-benzene												
AlkylBenzenes (36 – 40 min)												
TMPD-MIB		0.32		6.92	2.64	2.11	69.7	10.1	3.11		16.9	1.37
TMPD-DIB	6.05	30.6	2.44	663	468		369	845	263	2.64	2080	144
Acetophenone	0.94	1.12		7.10	0.30	1.41	0.21	0.43			0.41	0.46
Hexamethyl-cyclotrisiloxane			0.11	7.25	7.39	5.63		1.44	5.38		7.87	3.48
Octamethyl-cyclotetrasilox ane			0.36	1.01	1.70	0.64		0.38	1.61	0.09	1.56	0.21
Dimethyl methylphosphonate												
α-Pinene	1.65	1.04	0.99	3.49	2.07	0.85	0.79		8.36	1.79	2.95	1.00
3-Carene				0.34								
β-Pinene	1.15	0.85		0.69	1.61	1.11	0.43		5.19	1.55	1.06	0.86
d-l imonene				1.30				0.49			0.46	7 7 2

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Target Compound	bed	cabinet	carpet	ceiling	counter	curtain	door	seat	sub	vinyl	wall
	deck				top		interior	cushion	floor	floor	
Acetic acid	2400			11.1			24.0	0.25	911		
Phenol	26.0	7.24	69.6	6.91	0.34	97.2	0.33	164	1.99	26.9	4.34
2-ethyl-1-hexanol	0.58	0.20	0.71	1.26	0.60		0.01	3.24	0.81	0.74	0.80
Formaldehyde	136	419	57.6	22.1	4.97	323	14.3	410	14.8	1.69	33.7
Acetaldehyde	3.16	8.09	0.16				6.38		9.53		
Hexanal	27.8	0.64	1.89	0.49	0.73	0.46	0.78	9.52	3.32	3.73	0.37
Octanal			12.5								
Benzaldehyde	1.60	0.91	6.60	0.12	0.30	0.93	0.01	4.14	0.01	0.29	0.72
Nonanal	5.18	0.92	12.5	3.93	2.09	1.51		10.9	1.33		1.21
Dodecane	1.94		9.23	0.62				16.1	2.86	14.1	0.62
Tridecane	31.4	11.5	149	15.3	0.26	3.10	5.37	296	42.3	105	10.6
Tetradecane	41.7	17.9	122	30.6		14.2	10.6	235	34.3	62.1	13.5
Pentadecane	4.98	2.76	14.1	5.11	0.29	4.41	1.60	18.1	3.82	4.48	2.61
Hexadecane	1.29	2.40	2.60	2.29	0.23	2.41	0.60	3.84	1.12	1.46	1.30
Benzene	0.43	0.33	0.12		1.55	0.21	0.40	1.21	0.58		0.24
Toluene	2.61	1.73	0.06	1.21	0.85		0.83	3.71	1.66	0.48	1.18
Ethylbenzene			0.08		0.02		0.03	0.50			
p-Xylene			0.20		0.08		0.04	1.11			
Styrene			0.39		0.22	0.05	0.03	1.66			
Propyl-benzene							0.02				
1,3,5-trimethyl-benzene			0.13					0.39			
1,2,3-trimethy I-benzene			0.38					1.07			
AlkylBenzenes (36 min - 40 min)											
TMPD-MIB	2.30	1.69	6.44	4.13	9.51	4.98	0.58	12.26	5.89	5.38	1.63
TMPD-DIB	143	241	326	503	9.47	422	35.9	880	435	925	221
Acetophenone	1.04	1.28	1.98	0.21	0.12	0.52	0.13	1.07		0.52	0.97
Hexamethyl-cyclotrisiloxane	2.53	3.28		2.95	13.2			10.7	3.09	3.22	
Octamethyl-cyclotetrasiloxane	1.23	0.62		0.94	1.70			2.12	1.13	0.79	
Dimethyl methylphosphonate			0.33					0.42			
α-Pinene	7.80	3.74	2.58	2.90	1.89		0.11	10.16	4.27	1.34	1.90
3-Carene			1.56					1.89			
β-Pinene	8.24	3.64	0.68	2.03	1.14			4.24	2.59	0.78	1.33
d-Limonene	1.27		2.02				0 13	2 06			

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Target Compound	peq	bench	cabinet	ceiling	counter	curtain	door	end	seat	aub	vinyl	wall
	deck	seat	wall		top		interior	bench	cushion	floor	floor	
Acetic acid		575						34.5	584	4510		
Phenol	14.1	2.98	0.50	11.1	0.36	24.23	1.79	0.86	63.4	197	17.9	2.16
2-ethyl-1-hexanol	0.30	0.61	0.32	0.99	0.91		2.90	1.23	2.00	0.48	0.57	0.54
Formaldehyde	41.5	233	175	25.1	7.86	14.4	35.9	33.3	151	3.61		59.7
Acetaldehyde	12.8	2.43				2.47	0.75	1.66	4.50	18.3		
Hexanal	10.9	3.05	0.56	0.49	2.50	0.16	9.45	5.02	5.24	107	1.02	0.29
Octanal							0.57					
Benzaldehyde	0.82		0.05	0.34	1.48	0.16	0.02		1.60	14.6	1.42	0.60
Nonanal		3.40	0.96	7.60	1.07	0.04	29.7	1.43	5.36	1.46	0.86	0.71
Dodecane							1.47			6.54		
Tridecane		0.79		0.23		0.07	54.9		3.53	138	4.09	
Tetradecane	58.1	10.8	3.60	24.1		0.33	5.28	6.74	87.7	2060	82.8	3.79
Pentadecane	41.2	9.74	3.53	27.4		0.59	6.62	5.76	61.6	1210	52.6	3.78
Hexadecane	4.77	2.21	1.27	8.26		0.43	1.69	1.47	12.4	240	10.98	1.93
Benzene	0.21	0.47	0.64	0.22	1.52	13.3	0.54	0.04	0.57	0.08		0.08
Toluene	2.39	1.27	1.87	1.18	2.04		2.15	2.34	2.72	2.19	0.68	0.80
Ethylbenzene							0.02		0.41			
p-Xylene						0.02	0.07		1.00	1.14		
Styrene						0.00	0.02		0.63		0.42	
Propyl-benzene												
1,3,5-trimethy I-benzene												
1,2,3-trimethy I-benzene												
AlkylBenzenes (36 min - 40 min)												
TMPD-MIB	2.12	1.32		1.69	2.14			0.34	4.62	16.0	1.75	0.32
TMPD-DIB	11.3	4.70	1.26	26.4		1.59	1.97	1.22	26.2	1030	102	3.79
Acetophenone	0.22			0.49	1.08	0.14	0.26		0.46	1.15	0.19	0.71
Hexamethyl-cyclotrisiloxane			2.31						3.74	0.58	4.13	
Octamethyl-cyclotetrasiloxane		0.48	0.32				0.19	0.43	1.66	1.25	0.83	
Dimethyl methylphosphonate												
a-Pinene	4.92	2.90	3.46	2.37	1.17		1.51	4.99	7.89	31.7	2.06	1.37
3-Carene									2.10	1.34		
β-Pinene	5.56	1.92	2.29	2.63	1.48		0.92	2.94	2.73	15.7	1.00	1.15
d-Limonene	0.75					010	0.22		1 70	9 50		

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Target Compound	cabinet	cabinet	ceiling	counter	door	seat	seat	sub	vinyl	wall
	door	wall		top	interior	bottom	cushion	floor	floor	
Acetic acid							99.4	1220		
Phenol	0.79	1.15	5.70	0.43	1.19	15.3	85.2	47.8	21.9	7.16
2-ethyl-1-hexanol	0.52		0.75	0.23	0.00	3.08	17.3	0.76	0.80	1.26
Formaldehyde	91.8	488	20.1	9.69	35.7	293	30.4	416	13.9	26.0
Acetaldehyde	7.14	1.57		2.12	6.51	3.11		16.4	3.41	
Hexanal	3.24	1.08	0.48	1.27	3.83	11.7	1.56	81.6	8.10	0.98
Octanal								2.32		
Benzaldehyde		0.07	0.29	0.85	0.75	2.75	4.02	3.46	1.78	0.47
Nonanal	1.37		2.75	0.54	0.26	8.61	13.0	6.33	2.93	5.25
Dodecane						1.24	1.54	5.79	1.11	
Tridecane	3.43	0.79	5.48	0.02	0.96	28.1	87.4	238	46.6	3.20
Tetradecane	6.01	3.34	14.7		5.46	44.6	144	231	46.9	6.08
Pentadecane	1.49	0.86	4.12		2.01	9.14	22.5	22.3	6.90	1.98
Hexadecane	0.75		2.22	0.12	0.74	3.72	6.29	6.49	3.31	1.61
Benzene	0.51	0.15	0.44	0.28	0.44	3.86	0.76	0.20		0.13
Toluene	0.92	0.99	0.79	0.62	1.13	5.10	0.80	1.50		0.93
Ethylbenzene					0.02	0.99	0.36			
p-Xylene					0.02	2.58	1.16			
Styrene				0.36			0.27			
Propyl-benzene				0.11			0.19			
1,3,5-trimethy I-benzene							0.24			
1,2,3-trimethy I-benzene										
AlkylBenzenes (36 min - 40 min)										
TMPD-MIB	2.53	1.97	10.5	4.42	0.74	11.5	87.7	47.1	20.3	6.33
TMPD-DIB	81.3	7.77	771	0.26	139	939	1813	2920	2250	481
Acetophenone			0.29	0.17	0.32	1.01	2.81	0.90	0.44	0.48
Hexamethyl-cyclotrisiloxane	10.6			10.5	6.97		11.4		5.29	
Octamethyl-cyclotetrasiloxane	1.70	0.33		1.30	0.72	0.36	1.36	0.49	0.76	
Dimethyl methylphosphonate										
α-Pinene	1.10	1.52	1.13	1.09	0.63	6.17		50.2	1.67	1.55
3-Carene							0.06	0.81		
β-Pinene	0.81	0.91	1.17	0.48		4.65		11.4	0.63	1.99
d-Limonene						2.33	0.89	1.46		

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	bed	cabinet	cabinet	carpet	ceiling	counter	door	seat	sub	trim	vinyl	wall
	deck	wall	end			top	interior	cushion	floor		floor	
Acetic acid									1170			
Phenol	0.28	1.10	0.04	13.2	6.68	0.09	0.62	4.20	3.55	0.03	22.4	13.2
2-ethyl-1-hexanol	0.17		0.10	0.41	1.02		0.03	1.14	1.15	0.03	1.29	2.54
Formaldehyde	1.24	26.3	0.68	17.4		1.51	6.31	8.58	5.05	0.64		32.5
Acetaldehyde				2.05		0.32	0.02	0.76	13.4	0.27		
Hexanal	0.26		0.06	4.48	0.70	0.10	0.26	0.13	10.2	0.05	2.83	6.60
Octanal				3.37								
Benzaldehy de	0.18	0.71		9.89	0.33	0.24	0.03	0.29	0:30		0.89	2.45
Nonanal	0.46	0.08	0.00	5.44	2.22	0.10	0.19	3.12	3.08		3.37	32.2
Dodecane									1.00		0.82	
Tridecane	0.08		0.11	6.77	1.25		0.46	1.34	19.7		33.1	2.74
Tetradecane	0.11			18.2	7.69		1.15	3.88	52.9	0.04	84.6	6.34
Pentadecane				4.05	3.54		0.40	1.13	43.3		69.0	3.04
Hexadecane	0.09	0.47		0.79	1.61		0.23	0.25	8.03		11.1	1.23
Benzene	0.13		0.03	0.49		0.04	0.02	0.05				5.06
Toluene	0.41	1.10	0.11	0.93	1.67	0.09	0.11	0.01	2.88	0.04	0.54	3.46
Ethylbenzene				0.24				0.01				
p-Xylene				0.58				0.04				1.77
Styrene				0.28				0.02				
Propy -benzene								0.01				
1,3,5-trimethyl-benzene				0.07				0.02				
1,2,3-trimethy I-benzene												
AlkylBenzenes (36 min - 40 min)												
TMPD-MIB		0.48		2.85	3.10	0.28	9.64	1.25	3.65		14.8	4.10
TMPD-DIB	1.83	45.5	0.32	272	549		51.0	105	307	0.15	1820	431
Acetophenone	0.28	1.67		2.92	0.36	0.19	0.03	0.05			0.36	1.39
Hexamethyl-cyclotrisiloxane			0.01	2.98	8.65	0.76		0.18	6.31		6.91	10.4
Octamethyl-cyclotetrasiloxane			0.05	0.41	1.99	0.09		0.05	1.89	0.01	1.37	0.63
Dimethyl methylphosphonate												
α-Pinene	0:50	1.55	0.13	1.43	2.42	0.11	0.11		9.79	0.10	2.59	2.97
3-Carene				0.14								
β-Pinene	0.35	1.26		0.28	1.88	0.15	0.06		6.08	0.09	0.93	2.55
d-Limonene				0.53				0.06			0.41	82.7

Table 14. Material Emission Factors Normalized to Whole Trailer Floor Area (µg m⁻² h⁻¹) for the Dutchmen trailer

Target Compound	bed	cabinet	carpet	ceiling	counter	curtain	door	seat	aub	vinyl	wall
	deck				top		interior	cushion	floor	floor	
Acetic acid	460			10.6			2.41	0.02	911		
Phenol	4.97	4.69	25.2	6.61	0.03	8.41	0.03	16.2	1.99	17.2	8.68
2-ethyl-1-hexanol	0.11	0.13	0.26	1.21	0.05			0.32	0.81	0.48	1.61
Formaldehyde	26.1	271	20.9	21.1	0.38	28.0	1.44	40.5	14.8	1.08	67.5
Acetaldehy de	0.60	5.24	0.06				0.64		9.53		
Hexanal	5.31	0.42	0.68	0.47	0.06	0.04	0.08	0.94	3.32	2.38	0.74
Octanal			4.54								
Benzaldehyde	0.31	0.59	2.39	0.11	0.02	0.08		0.41	0.01	0.18	1.43
Nonanal	0.99	0.60	4.54	3.75	0.16	0.13		1.07	1.33		2.42
Dodecane	0.37		3.34	0.59				1.59	2.86	9.03	1.24
Tridecane	6.00	7.45	53.9	14.6	0.02	0.27	0.54	29.3	42.3	67.2	21.3
Tetradecane	7.98	11.6	44.0	29.3		1.23	1.07	23.2	34.3	39.6	26.9
Pentadecane	0.95	1.79	5.12	4.89	0.02	0.38	0.16	1.78	3.82	2.86	5.22
Hexadecane	0.25	1.56	0.94	2.19	0.02	0.21	0.06	0.38	1.12	0.93	2.59
Benzene	0.08	0.22	0.04		0.12	0.02	0.04	0.12	0.58		0.48
Toluene	0.50	1.12	0.02	1.16	0.07		0.08	0.37	1.66	0.30	2.36
Ethylbenzene			0.03					0.05			
p-Xylene			0.07		0.01			0.11			
Styrene			0.14		0.02			0.16			
Propyl-benzene											
1,3,5-trimethyl-benzene			0.05					0.04			
1,2,3-trimethy I-benzene			0.14					0.11			
AlkylBenzenes (36 min - 40 min)											
TMPD-MIB	0.44	1.10	2.33	3.95	0.73	0.43	0.06	1.21	5.89	3.43	3.26
TMPD-DIB	27.4	156	118	481	0.72	36.5	3.61	87.0	435	590	443
Acetophenone	0.20	0.83	0.72	0.20	0.01	0.05	0.01	0.11		0.33	1.95
Hexamethyl-cyclotrisiloxane	0.48	2.12		2.82	1.01			1.06	3.09	2.06	
Octamethy I-cyclotetrasilox ane	0.24	0.40		0.90	0.13			0.21	1.13	0.50	
Dimethyl methylphosphonate			0.12					0.04			
α-Pinene	1.49	2.42	0.93	2.77	0.14		0.01	1.00	4.27	0.86	3.80
3-Carene			0.57					0.19			
β-Pinene	1.58	2.36	0.24	1.94	0.09			0.42	2.59	0.50	2.65
d-Limonene	0.24		0.73				0.01	0.29			

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Target Compound	bed	bench	cabinet	ceiling	counter	curtain	door	end	seat	sub	vinyl	wall
	deck	seat	wall		top		interior	bench	cushion	floor	floor	
Acetic acid		139						1.40	51.1	4510		
Phenol	1.69	0.72	0.15	11.9	0.03	3.67	0.08	0.03	5.54	197	17.9	6.07
2-ethyl-1-hexanol	0.04	0.15	0.10	1.06	0.07		0.13	0.05	0.18	0.48	0.57	1.51
Formaldehyde	5.00	56.3	53.5	26.9	0.63	2.17	1.56	1.35	13.2	3.61		168
Acetaldehyde	1.55	0.59				0.37	0.03	0.07	0.39	18.3		
Hexanal	1.31	0.74	0.17	0.52	0.20	0.02	0.41	0.20	0.46	107	1.02	0.81
Octanal							0.02					
Benzaldehyde	0.10		0.02	0.36	0.12	0.02			0.14	14.6	1.42	1.69
Nonanal		0.82	0.29	8.15	0.09	0.01	1.29	0.06	0.47	1.46	0.86	2.01
Dodecane							0.06			6.54		
Tridecane		0.19		0.24		0.01	2.39		0.31	138	4.09	
Tetradecane	7.00	2.61	1.10	25.8		0.05	0.23	0.27	7.66	2060	82.8	10.7
Pentadecane	4.97	2.35	1.08	29.3		0.09	0.29	0.23	5.38	1210	52.6	10.6
Hexadecane	0.58	0.53	0.39	8.85		0.07	0.07	0.06	1.08	240	11.0	5.44
Benzene	0.02	0.11	0.20	0.24	0.12	2.01	0.02		0.05	0.08		0.22
Toluene	0.29	0.31	0.57	1.26	0.16		0.09	0.10	0.24	2.19	0.68	2.26
Ethylbenzene									0.04			
p-Xylene									0.09	1.14		
Styrene									0.06		0.42	
Propyl-benzene												
1,3,5-trimethyl-benzene												
1,2,3-trimethyl-benzene												
AlkylBenzenes (36 min - 40 min)												
TMPD-MIB	0.26	0.32		1.81	0.17			0.01	0.40	15.95	1.75	0.91
TMPD-DIB	1.36	1.14	0.39	28.3		0.24	0.09	0.05	2.29	1030	102	10.7
Acetophenone	0.03			0.53	0.09	0.02	0.01		0.04	1.15	0.19	2.01
Hexamethyl-cyclotrisiloxane			0.71						0.33	0.58	4.13	
Octamethyl-cyclotetrasiloxane		0.12	0.10				0.01	0.02	0.15	1.25	0.83	
Dimethyl methylphosphonate												
α-Pinene	0.59	0.70	1.06	2.55	0.09		0.07	0.20	0.69	31.72	2.06	3.86
3-Carene									0.18	1.34		
ß-Pinene	0.67	0.47	0.70	2.82	0.12		0.04	0.12	0.24	15.71	1.00	3.24
d-Limonene	0.09					0.01	0.01		0.15	6.58		

Table 16. Material Emission Factors Normalized to Whole Trailer Floor Area (µg m⁻² h⁻¹) for the Coachmen trailer

Target Compound	cabinet	cabinet	ceiling	counter	door	seat	seat	aub	vinyl	wall
	door	wall		top	interior	bottom	cushion	floor	floor	
Acetic acid	-						10.3	1220		
Phenol	0.01	1.03	5.42	0.02	0.11	5.20	8.80	47.8	21.9	21.7
2-ethyl-1-hexanol			0.71	0.01		1.05	1.79	0.76	0.80	3.84
Formaldehyde	0.76	435	19.2	0.56	3.32	99.6	3.14	416	13.9	79.0
Acetaldehyde	0.06	1.40		0.12	0.61	1.06		16.4	3.41	
Hexanal	0.03	0.96	0.46	0.07	0.36	3.99	0.16	81.6	8.10	2.97
Octanal								2.32		
Benzaldehyde		0.06	0.27	0.05	0.07	0.93	0.42	3.46	1.78	1.43
Nonanal	0.01		2.62	0.03	0.02	2.93	1.34	6.33	2.93	15.95
Dodecane						0.42	0.16	5.79	1.11	
Tridecane	0.03	0.70	5.22		0.09	9.54	9.03	238	46.6	9.70
Tetradecane	0.05	2.98	14.0		0.51	15.2	14.9	231	46.9	18.4
Pentadecane	0.01	0.77	3.92		0.19	3.11	2.33	22.3	6.90	6.00
Hexadecane	0.01		2.11	0.01	0.07	1.26	0.65	6.49	3.31	4.88
Benzene		0.14	0.42	0.02	0.04	1.31	0.08	0.20		0.38
Toluene	0.01	0.88	0.75	0.04	0.11	1.73	0.08	1.50		2.84
Ethylbenzene						0.34	0.04			
p-Xylene						0.88	0.12			
Styrene				0.02			0.03			
Propyl-benzene				0.01			0.02			
1, 3, 5-trimethy I-benzene							0.03			
1,2,3-trimethy I-benzene										
AlkylBenzenes (36 min - 40 min)										
TMPD-MIB	0.02	1.76	9.96	0.25	0.07	3.92	90.06	47.1	20.3	19.2
TMPD-DIB	0.67	69.4	734	0.01	13.0	319	187	2920	2250	1460
Acetophenone			0.28	0.01	0.03	0.34	0.29	0.90	0.44	1.47
Hexamethyl-cyclotrisiloxane	0.09			09.0	0.65		1.17		5.29	
Octamethyl-cyclotetrasiloxane	0.01	0.29		0.07	0.07	0.12	0.14	0.49	0.76	
Dimethyl methylphosphonate										
α-Pinene	0.01	1.35	1.08	0.06	0.06	2.10		50.3	1.67	4.71
3-Carene							0.01	0.81		
β-Pinene	0.01	0.81	1.12	0.03		1.58		11.4	0.63	6.05
d-Limonene						0.79	0.09	1.47		

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Target Compound	Total	bed deck	cabinet wall	cabinet	carpet	ceiling	counter	door, interior	seat	sub floor	trim	vinyl floor	wall
		1000		2			42					8	
TMPD-DIB	3590		1%		8%	15%		1%	3%	%6		51%	12%
Acetic acid	1170									100%			
Tetradecane	175				10%	4%			2%	30%		48%	4%
Pentadecane	124				3%	3%				35%		55%	2%
Formaldehyde	100	1%	26%		17%		2%	6%	%6	5%			32%
d-Limonene	84												%66
Tridecane	66				10%	2%			2%	30%		50%	4%
Phenol	65		2%		20%	10%			6%	5%		34%	20%
Nonanal	51			2%	11%	4%			%9	%9		%2	63%
TMPD-MIB	40		1%		7%	8%		24%	3%	%6		37%	10%
Hexamethyl-cyclotris ilox ane	36				8%	24%	2%			17%		19%	29%
Hexanal	26				17%	3%		1%		40%		11%	26%
Hexadecane	24		2%		3%	7%			1%	34%		47%	5%
α-Pinene	22	2%	2%		%2	11%				45%		12%	14%
Acetaldehyde	17				12%		2%		5%	80%	2%		
Benzaldehyde	15	1%	5%		65%	2%	2%		2%	2%		%9	16%
β-Pinene	14	3%	%6		2%	14%	1%			45%		%2	19%
Toluene	1	4%	10%		8%	15%				25%		5%	30%
2-ethyl-1-hexanol	8	2%		1%	5%	13%			14%	15%		16%	32%
Acetophenone	7	4%	23%		40%	5%	3%					5%	19%
Octamethyl-cyclotetrasiloxane	9				%9	31%	1%			29%		21%	10%
Benzene	9	2%			8%								87%
Octanal	e				100%								
p-Xylene	2				24%				2%				74%
Dodecane	2									55%		45%	
Styrene	٢				94%				%9				
Ethylbenzene	ř				95%				5%				
3-Carene	v				100%								
1,3,5-trimethyl-benzene	v				80%				20%				
Propy -benzene	<b>~</b>								100%				

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Target Compound	Total	bed deck	cabinet	carpet	ceiling	counter top	curtain	door, interior	seat cushion	sub floor	vinyl floor	wall
TXIB-DIB	2380	1%	7%	5%	20%		2%		4%	18%	25%	19%
Acetic acid	1380	33%								%99		
Formaldehyde	493	5%	55%	4%	4%		6%		8%	3%		14%
Tridecane	243	2%	3%	22%	6%				12%	17%	28%	%6
Tetradecane	219	4%	5%	20%	13%				11%	16%	18%	12%
Phenol	<u>94</u>	5%	5%	27%	7%		6%		17%	2%	18%	%6
Pentadecane	27	4%	7%	19%	18%		1%		7%	14%	11%	19%
TXIB-MIB	23	2%	5%	10%	17%	3%	2%		5%	26%	15%	14%
Dodecane	19	2%		18%	3%				8%	15%	47%	%2
α-Pinene	18	8%	14%	5%	16%				%9	24%	5%	21%
Acetaldehyde	16	4%	33%					4%		59%		
Nonanal	15	%2	4%	30%	25%	1%			7%	%6		16%
Hexanal	14	37%	3%	5%	3%				7%	23%	16%	5%
Hexamethyl-cyclotrisiloxane	13	4%	17%		22%	8%			8%	24%	16%	
β-Pinene	12	13%	19%	2%	16%				3%	21%	4%	21%
Hexadecane	10	2%	15%	%6	21%		2%		4%	11%	%6	25%
Toluene	8	7%	15%		15%			1%	5%	22%	4%	31%
Benzaldehyde	9	%9	11%	43%	2%		1%		7%		3%	26%
2-ethyl-1-Hexanol	5	2%	3%	5%	24%				%9	16%	10%	32%
Octanal	5			100%								
Acetophenone	4	5%	19%	16%	2%		1%		2%		8%	44%
Octamethyl-cyclotetrasiloxane	4	%2	11%		26%	4%			%9	32%	14%	
Benzene	2	5%	13%	2%		7%	1%	2%	7%	34%		28%
d-Limonene	-	19%		57%				1%	23%			
3-Carene	-			75%					25%			
Styrene	٧			43%		5%	1%	1%	50%			
1,2,3-trimethyl-benzene	٧			57%					43%			
p-Xylene.	v			37%		3%		2%	57%			
Dimethyl methylphosphonate	v			74%					26%			
1,3,5-trimethy I-benzene	٧			26%					44%			
Ethylbenzene	v			36%		2%		4%	58%			
Propy-benzene	v							100%				

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Target Compound	Total	bed deck	bench seat	cabinet wall	ceiling	counter top	curtain	door, interior	end of bench	seat cushion	sub floor	vinyl floor	wa
Acetic acid	4500		3%							1%	%96		
Tetradecane	2190				1%						94%	4%	
Pentadecane	1310				2%						92%	4%	
TMPD-DIB	1170				2%						88%	%6	
Formaldehyde	332	2%	17%	16%	8%					4%	1%		51%
Hexadecane	268				3%						%06	4%	2%
Phenol	245				5%		2%			2%	80%	2%	2%
Tridecane	145							2%			95%	3%	
Hexanal	113	1%									95%		
α-Pinene	44	1%	2%	2%	%9					2%	73%	5%	%6
β-Pinene	25	3%	2%	3%	11%						63%	4%	13%
TMPD-MIB	22	1%	1%		8%					2%	74%	8%	4%
Acetaldehyde	21	7%	3%				2%			2%	86%		
Benzaldehyde	18				2%						%62	8%	%6
Nonanal	16		5%	2%	53%			8%		3%	%6	%9	13%
Toluene	8	4%	4%	7%	15%	2%		1%	1%	3%	27%	8%	28%
D-Limonene	7	1%								2%	%96		
Dodecane	7										%66		
Hexamethyl-cyclotrisiloxane	9			12%						%9	10%	72%	
2-ethyl-1-hexanol	4		3%	2%	25%	2%		3%	1%	4%	11%	13%	35%
Acetophenone	4				13%	2%					28%	5%	49%
Benzene	e		4%	6%	8%	4%	65%			2%	3%		7%
Octamethyl-cyclotetrasiloxane	2		5%	4%						%9	51%	34%	
3-Carene	2									12%	88%		
p-Xylene	-									7%	92%		
Styrene	۷									12%		88%	
Ethylbenzene	v							2%		88%			
Octanal	v							100%					

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Table 21. Total (µg m⁻² h⁻¹) and Percent Contribution of Each Material to Area Normalized Whole Trailer Emission Rates for the Cavalier

Target Compound	Total	cabinet door	cabinet wall	ceiling	counter top	door, interior	seat bottorn	seat cushion	sub floor	vinyl floor	wall
TMPD-DIB	7960			%6			4%	2%	37%	28%	18%
Acetic acid	1230								%66		
Formaldehyde	1070		41%	2%			%6		39%	1%	%2
Tetradecane	344			4%			4%	4%	67%	14%	5%
Tridecane	319			2%			3%	3%	75%	15%	3%
Phenol	112			5%			5%	8%	43%	20%	19%
TMPD-MIB	112		2%	%6			4%	8%	42%	18%	17%
Hexanal	66						4%		83%	8%	3%
α-Pinene	61		2%	2%			3%		82%	3%	8%
Pentadecane	46		2%	%6			%2	5%	49%	15%	13%
Nonanal	32			8%			6%	4%	20%	6%	50%
Acetaldehyde	23		%9			3%	5%		71%	15%	
β-Pinene	22		4%	5%			%2		53%	3%	28%
Hexadecane	19			11%			%/	3%	35%	18%	26%
2-ethyl-1-hexanol	6			8%			12%	20%	%6	%6	43%
Benzaldehyde	8			3%			11%	5%	41%	21%	17%
Toluene	8		11%	%6		1%	22%	1%	19%		36%
Hexamethyl-cyclotrisiloxane	8	1%			8%	8%		15%		68%	
Dodecane	7						%9	2%	%11	15%	
Acetophenone	4			%2			%6	8%	24%	12%	39%
Benzene	e		5%	16%		2%	51%	3%	8%		15%
D-Limonene	2						34%	4%	62%		
Octanal	2								100%		
Octamethyl-cyclotetrasiloxane	2		15%		4%	3%	%9	%2	25%	39%	
p-Xylene.	-						88%	12%			
3-Carene	-								%66		
Ethylbenzene	٧						%06	10%			
Styrene	v				42%			58%			
Propyl-benzene	v				24%			76%			
1.3.5-trimethyl-benzene	v							100%			

Table 22. Comparison of Sum Material Specific Emission* (µg m⁻² h⁻¹) with Measured Whole Trailer Emission

	Ducl	hmen	Pilg	grim	Coac	hmen	Cav	alier
Target Compound	Material	Whole trailer	Material	Whole trailer	Material	Whole trailer	Material	Whole trailer
Acetic acid		46.0	473	35.0	191	13.3	10	41.6
Phenol	62	16.0	92	14.3	48	16.9	64	16.1
2-ethyl-1-hexanol	7	1.08	4	0.88	4	0.39	8	2.50
Formaldehyde	95	230	478	220	329	306	655	288
Acetaldehyde	3	11.3	7	6.72	3	10.6	7	8.29
Hexanal	15	8.15	11	7.80	6	6.11	17	15.8
Octanal	3	3.29	5	1.86		1.85		3.01
Benzaldehyde	15	1.17	6	0.40	4		5	1.03
Nonanal	48	5.12	14	3.14	14	4.85	26	5.07
Dodecane	1	0.18	16	3.79		0.27	2	0.62
Tridecane	46	9.25	200	44.8	7	1.00	81	20.1
Tetradecane	122	16.8	185	31.8	138	28.9	113	18.8
Pentadecane	81	4.09	23	2.96	107	19.0	23	2.93
Hexadecane	16	1.00	9	0.77	28	4.22	12	0.9
Benzene	6		1	0.04	3		2	
Toluene	8	1.15	6	0.33	6	0.66	6	0.5
Ethylbenzene		0.15		0.12		0.18		0.24
p-Xylene	2	0.18		0.12		0.27	1	0.1
Styrene		3.52		3.69		2.44		7.3
Propyl-benzene						0.45		
1,3,5-trimethyl-benzene		0.06		0.14		0.07		
1,2,3-trimethyl-benzene		0.11		0.33				
TMPD-MIB	36	3.57	17	2.50	6	1.25	65	8.78
TMPD-DIB	3280	245	1940	101	147	9.56	5040	197
Acetophenone	7		4		3		3	
Hexamethyl-cyclotrisiloxane	30	0.17	10	0.23	5	0.67	8	0.5
Octamethyl-cyclotetrasiloxane	5	0.03	2	0.05	1	0.15	1	0.1
Dimethyl methylphosphonate								2.74
a-Pinene	12	45.6	13	29.7	12	19.3	11	28.1
3-Carene		1.12	1	1.52		2.41		4.1
β-Pinene	8	5.01	10	4.16	9	2.66	10	4.4
d-Limonene	84	1.67	1	1.19		0.80	1	2.2

*The sum of the material normalized to floor area excluding the contribution from sub floor.

		Em	ission factor	(µg m⁻²	h⁻¹)	
		Cabinetry N	/laterials*		Passage	Plywood
Compound	РВ Тор	PB case	Hardboard	Stile	door	subfloor*
Formaldehyde	92, 82	470	10	330	153	11, 8
Acetaldehyde	38, 40			20	11	19, 10
Pentanal	51, 42			36	8	28, 25
Hexanal	249, 199			260	42	169, 161
2-Furaldehyde	6, 5		72	7		
Heptanal	12, 9			7		4, 3
2-Heptenal	8, 5			9		5, 5
Benzaldehyde	16			42	3	5
Octanal	22, 18			28		8, 8
2-Octenal	19, 12			29		12, 11
Nonanal	19, 16			40		21, 22

Table 23. Material specific aldehyde emissions from cabinetry, passage door, and subfloor used to fabricate a new manufactured house

*Values are presented for duplicate specimens separated by a comma. The data are for new material direct from factory as reported in Hodgson et. al. 2002

Table 24. Material specific emission factors of terpene hydrocarbons from indoor sources used to
fabricate a new manufactured house

Compound	PB countertop ^a	Cabinet frame lumber	Plywood subfloor ^a
	(µg m⁻² h⁻¹)	(µg m⁻² h⁻¹)	(µg m⁻² h⁻¹)
$\alpha$ -Pinene	19, 26	14	114, 278
β-Pinene	7,7	17	29, 69
d-Limonene	6, 6	<3	29, 113

a Values are presented for duplicate specimens separated by a comma. The data are for new material direct from factory as reported in Hodgson et. al. 2002

Material		Emission (µg m ⁻²	Factor ² h ⁻¹ )		Sample Size	Notes
	Mean	Median	Min	Max	Ν	
Hardwood Plywood	87	74	6.8	170	12	¼"-¾" stock HWPW and ½" HWPW-VC
Medium Density Fiberboard	293	288	210	385	6	5∕8"-¾" MDF
MDF Cabinet Doors	420		364	535	2	
Particleboard	189	161	104	508	22	5⁄8'-3/4" industrial PB, 5⁄8" PB underlayment, and 5∕8" mobile home decking
¼" Particleboard Wallpaper	1375 27		1170	1580	2	Ű
Coated MDF Cabinet Doors	880		460	1300	2	
Coated PB – Paper Laminated	60	52	26	120	6	
Coated PB – Mobile Home Decking	44		35	52	2	
Coated PB – Melamine Laminated	20	11	2.2	86	12	
Coated PB – Rigid Vinyl	24		16	31	2	
Coated PB – Vinyl or Acrylic	4	2.7	1.3	8.6	8	
Interior Door – PB Core	11		7.0	15	2	

## Table 25. Reported Formaldehyde Emission Factors from CARB's Battelle (1996) study^a.

^a Source: Appendix D Basis for Formaldehyde Emission Factors, Rulemaking to Consider Adoption of the Proposed Airborne Toxic Control Measure (ACTM) to Reduce Formaldehyde Emissions From Composite Wood Products, California Air Resources Board. April 2007.

http://www.arb.ca.gov/regact/2007/compwood07/compwood07.htm

## **FIGURES:**



Figure 1. Preparation for indoor sampling in a THU. Half inch holes were drilled into the THU door for insertion of ¹/₄" stainless steel sampling tubes. A sampling tube and sample pump are seen in the foreground.



Figure 2. Collection of indoor sample through the THU door.

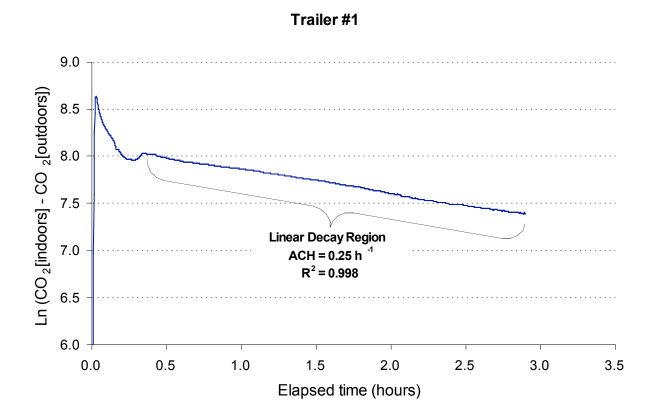


Figure 3. Example of tracer gas experiment determining ventilation rate in trailer showing initial stabilization period followed by the linear decay region. The ventilation rate is determined from the slope of the decay curve in the linear region as described in the text. The response shown here for Trailer 1 is typical of all the units tested.

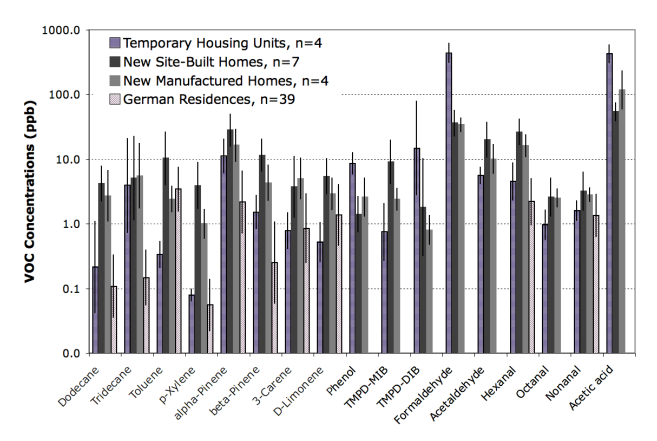


Figure 4. Comparison of measured indoor air concentration (ppm) data for new site-built and manufactured homes (Hodgson et. al., 2000), German residences (Hippelein, 2004) and the four THUs. The data are reported as geometric mean (GM) with error bars representing one geometric standard deviation (GSD). The GSD for the Hippelein (2004) data were calculated from the arithmetic mean and standard deviation.

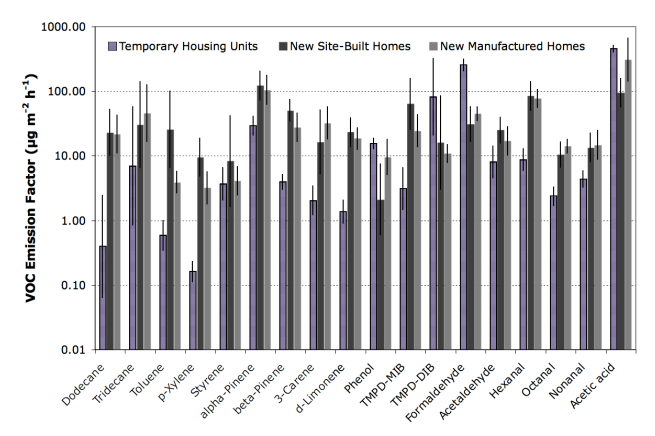


Figure 5. Comparison of GM (GSD error bars) measured whole building VOC emission factors (emissions per floor area) for seven new site built houses, four new manufactured houses (Hodgson et. al. 2000), and the four THUs studied in this project. Note that this chart is plotted on a logarithmic scale.

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