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Multiscale Design And Life-Cycle Based Sustainability Assessment Of Polymer Nanocomposite Coatings

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MULTISCALE DESIGN AND LIFE-CYCLE BASED SUSTAINABILITY ASSESSMENT OF POLYMER NANOCOMPOSITE COATINGS

by

ROHAN G. UTTARWAR

DISSERTATION

Submitted to the Graduate School

of Wayne State University,

Detroit, Michigan

in partial fulfillment of the requirements

for the degree of

DOCTOR OF PHILOSOPHY

2013

MAJOR: CHEMICAL ENGINEERING

Approved by:

____________________________________
Advisor

____________________________________
Date
DEDICATION

To my father Ganesh,

my mother Madhuri

and

my sister Ruchita
ACKNOWLEDGEMENTS

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CHAPTER 1
INTRODUCTION

Paint is a type of polymeric material made of four main ingredients: pigment, resin, solvent and additives. It is well known that paint have numerous applications over wide industries which include automotive, aerospace, ship-making, military, etc. Thus, paint material has been studied continuously to obtain improved properties and performance of the final coating. Out of all the ingredients, it is the resin/binder that leads to most of the physical and chemical properties of the paint material. Most common resin materials are acrylic, alkyd, epoxy, phenolic resin, unsaturated polyester and polyurethane. With the advancement of polymer engineering and paint technology, various inorganic fillers (clays, metals, minerals, etc.) are being added into coating formulations to improve some performance properties. It was around three decades ago when technological advances allowed addition of decreasing sizes of various fillers at the nanometer scale. It brought us to the realization of significant improvement of coating film’s performance that is achievable with “nanocomposite coatings”.

1.1 Challenges for Nanocoating Technology

Nanocomposite coating technology has become a rapidly expanding area of research. It encompasses a tremendous variety of systems with substantially improved properties and novel functionalities. Some of the already developed nanocoatings are claimed to have enhanced barrier and mechanical properties such as tensile strength, stiffness, elongation at break, impact
strength, etc. (Chow and Ishak, 2007; Nobel et al., 2007). Some exhibit improved flame retardancy and thermal resistance (Fang et al., 2011). These materials also have become increasingly important because of their electrical and magnetic properties (Dincer et al., 2012). One of the obvious benefits of using nanocomposite coatings over conventional coatings is that all the superior properties can be achieved with typically 5-10% (by weight) loading of nanomaterials, while the conventional coatings may require 10-50% (by weight) loading of inorganic fillers into coating compositions. Most of the superior properties achievable with nanoparticles addition into coating matrix seem to depend not only on the properties of individual entities but also on the interfacial interaction between organic and inorganic phases (Baur and Silverman, 2007). As a result of all advanced properties, nanoparticle coatings are finding numerous applications in automotive, aerospace, ship-making, security, chemical, electronics, steel, construction, and many other industries (Khanna, 2008).

Table 1.1 summarizes some known research on polymer nanocomposites. It provides general information about various types of nanoparticles leading to specific superior or smart functionalities. There are also more combinations of polymer systems and nanoparticles which deliver improved properties of the final nanocomposite. These nanocomposites not only have applications in paints and coatings but also in plastic industry.
Table 1.1. Summary of R&D on polymer-nanocomposites in past decade

<table>
<thead>
<tr>
<th>Publication</th>
<th>Polymer type</th>
<th>Nanoparticle type</th>
<th>Performance improvement type</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bharadwaj et al., 2002</td>
<td>Polyester</td>
<td>MMT platelet</td>
<td>Corrosion prevention</td>
</tr>
<tr>
<td>Nobel et al., 2007</td>
<td>Acrylic resin</td>
<td>Boehmite, laponite disk, MMT platelet</td>
<td>Stiffness</td>
</tr>
<tr>
<td>Dastjerdi et al., 2012</td>
<td>Polysiloxane emulsion</td>
<td>Colloidal TiO$_2$, Silver</td>
<td>Stain repellency</td>
</tr>
<tr>
<td>Teng, 2011</td>
<td>Acrylic resin</td>
<td>TiO$_2$, SiO$_2$</td>
<td>Self-cleaning, hydrophobicity</td>
</tr>
<tr>
<td>Refat et al., 2009</td>
<td>Polyethylene</td>
<td>Nano silicate (MMT clay)</td>
<td>High thermal stability</td>
</tr>
<tr>
<td>Mirabedini et al., 2011</td>
<td>Acrylic coating</td>
<td>TiO$_2$, SiO$_2$</td>
<td>Enhanced photocatalytic activity</td>
</tr>
<tr>
<td>US 6387519, 2002 (PPG)</td>
<td>Acrylic-melamine</td>
<td>Ceramic</td>
<td>Scratch resistance</td>
</tr>
<tr>
<td>US 20090081373, 2009 (Nanovere)</td>
<td>Dendrimer</td>
<td>Zinc oxide</td>
<td>Scratch and chemical resistance, self-cleaning</td>
</tr>
<tr>
<td>US 20090136441, 2009 (Taiwan)</td>
<td>Oligomer</td>
<td>Fluoroalkylsilane modified metal oxide</td>
<td>Anti-fouling</td>
</tr>
</tbody>
</table>

Although PNC coatings have these numerous advantages, they also come with some drawbacks. The study of these nanocomposite materials requires a multi-disciplinary approach (Judeinstein and Sanchez, 1996). In spite of the fact that these materials have been investigated for several years, no profound knowledge about their physicochemical attributes that lead to the superior properties is available. Due to the existence of vast design parameters and experimental complexity, nanopaint design optimality is extremely difficult to address. The challenges to the development of these materials come in the production stages, which include,

Processing. The compatibility between nanoparticles and polymer resin is usually poor. It is challenging to control a uniform dispersion of nano-fillers inside the coating matrix. Only a few thermoplastic polymer resins are easily compatible with selective inorganic filler systems.
In many cases, the intercalation of nanoclays can change the functionality of the polymeric material and inhibit certain coating film properties.

**Property Optimization.** The properties of the final coating film depend largely on polymer morphology, nanoparticles chemistry, size, shape and dispersion within the coating film. In order to obtain optimal properties of the coating material, a thorough understanding of the formulation is needed. The current research and scientific knowledge is insufficient to develop nanocoating materials with various combinations of resin and nanoparticle.

The above mentioned challenges prevent nanopaint-based products from getting accepted for large scale manufacturing. Thus, it becomes essential to build a thorough understanding of the correlation between nanocomposite behavior on molecular scale and its effects on the performance. An integrated computational-experimental approach for synthesis, development and analysis of PNCs should be very useful for studying the fundamentals of molecular behavior leading to their superior or entirely new functional properties. It is known that computational material design can provide impressive freedom and control over the investigation of material parameters and product properties through virtually any number of *in-silico* experiments. Moreover, multiscale modeling and simulation can greatly facilitate identification of important correlations among material, structure, property, and performance. Recent studies on computational modeling and simulation of smart PNCs provided some deep understanding of the science behind the superior functionalities of scratch resistance and corrosion prevention (Xiao and Huang, 2009; Xiao *et al*., 2010; Chen *et al*., 2008; Khanna, 2008). The model-based approach should be very valuable for optimally designing experiments, but the computational findings must be validated experimentally for material development. On other hand,
experimental study will provide reliable information for more advanced computational design of new properties of smart PNC.

In order to ascertain the potential of PNC coatings, it is important to look at the entire product life-cycle for the impact of nanoparticles addition to conventional paint formulations. The two key stages of the life-cycle of nanocoating are paint manufacturing and coating formation. There have been serious concerns about the environmental and health issues associated with nanomaterials (Lewisky, 2008). These environmental risks depend on the type and concentration of nanoparticles and exposed surrounding. Although there have been many studies on toxicology of nanoparticles (Schrand et al., 2010; Napierska et al., 2010), the effects of nanopaint exposure and application have not been studied in detail. There is a serious knowledge gap existing between nanocomposite coatings potential and sustainability issues, which include nanoparticles-related environmental threats as well as life cycle performance of nanocoating products and the overall performance in terms of energy use, safety, water use, waste emission, etc.

For automotive bodies, the paint material is applied via paint-spray technique. It is the most common application practice which contributes to significant energy consumption and environmental emission of toxic chemicals. In production cycle, paints-spray technique decides several key issues, such as transfer efficiency, wet film topology and VOC/nanoparticle emission, which correspond to economic, quality and environmental performances, respectively.

1.2 Motivation

Nanocoating technology, in-spite of showing tremendous potential for delivering superior
properties, faces various challenges to get accepted in a commercial market. Paint manufacturers (e.g. BASF, DuPont and AkzoNobel) and end users (e.g. automotive, industrial, and architectural industries) are scrutinizing this evolving category of paints for all the possible benefits and risks involved. There have been several experimental techniques existing for the synthesis of nanopaints with different types of nanoparticles (Tigli and Evren, 2005; Chow and Mohd Ishak, 2007; Nobel et al., 2007; Chen et al., 2008). However, it is extremely difficult to address the issues involved with the optimization of their formulations solely using experiments due to huge design space and experimental complexity. In a cradle-to-grave life cycle of nanocoatings, there are innumerable key parameters which need to be examined before this technology can be accepted for production of commercial paints. The first and most important concern is the risk to environment and human health that this nanoparticle-based technology can pose. The release of toxic nanoparticles during various stages of life-cycle, which include extraction/mining, paint manufacturing, application and end use needs to be assessed and analyzed through experiments and computational modeling if the empirical data is not available. The second concern is the lack of scientific knowledge regarding the correlation among coating’s material formulation, microstructure, mesoscopic properties and final macroscopic film performance. This issue brings-in the need for multiscale computational modeling and simulation. The simulation of nanocomposite material on a molecular level can provide key insights into structure-property correlations which can help experimentalist in developing optimal formulations of nanocoatings. The macroscopic simulation of paint application process can generate crucial information related to nanoparticles emission and exposure to the surrounding atmosphere. Such findings can assist in making the paint application phase of the life-cycle of nanocoating technology more sustainable.
The life cycle data from all the stages can be used effectively to generate recommendations for the future research only if it is assessed in a comprehensive and methodological way. There are several sustainability matrixes available for the assessment of chemical processes and manufacturing industries (e.g. AIChE, IChemE, etc.). However, a complete framework of sustainability matrixes for all the pillars of sustainability (economic, environmental and social) and for all the stages of life cycle of a nanocoating technology is not available. Such life-cycle-based sustainability assessment framework is useful for determination of the right parameters and indicators and collection of appropriate data for the analysis of this nanotechnology. This research frontier can also provide guidance to nanocoating researchers and end users for integrating life cycle assessment and sustainability criteria into nanopaint design, which becomes an urgent need in nanomaterial development.

1.3 Multiscale Modeling and Simulation

The multiscale modeling and simulation approach has become increasingly important over the last decade. It has found applications in various fields, such as, plasma and thermal spray, crystallization, material design, chemical reactor designing, etc. (Vlachos, 1997; Kramer et al., 1999; Fauchais and Vardella, 2000; Maroudas, 2000; Li and Kwauk, 2001). It has helped in generating many new understandings related to the system’s final product or process. It can provide a pathway in addressing all the necessary information about the whole spectrum of coating quality parameters (nano- to macro- scale), material efficiency parameters and energy efficiency parameters at the same time. Such type of comprehensive study is not possible with existing mono-scale modeling approaches. However, the field of multiscale modeling and
simulation requires further exploration. Since, sustainable nanocoating technology development needs a complete understanding of microstructure, material, product and process behavior throughout a wide range of length (10⁹ – 10 m) and time (10⁻⁶ – 10³ s) scales, multiscale modeling and simulation becomes a must.

The modeling of microstructure of polymer nanocomposites and the correlation with material properties is done on micro- and meso- scale using molecular simulation techniques. The simulation at macro scale for the paint application process is performed using computational fluid dynamics (CFD) based modeling technique.

**Molecular simulation of polymer nanocomposite coatings.** Molecular modeling and simulation is a very useful tool of research especially in the field of nano scale sciences due to the compatibility at all the time and length scales involved. In general, molecular simulations are used to calculate or support properties determined through experiments; but more importantly, this approach is very effective for systems that have not been studied through experiments to guide the development of novel material functionalities and properties. With the aid of advanced technology and computer power today, the modeling and simulation of complex systems, such as polymers, has become possible within an acceptable time range. Classical molecular simulation approaches involve modeling of material systems via functional forms that define inter-molecular and intra-molecular interactions among various entities of the molecule. These functional forms and corresponding parameter sets form a ‘force field’ that is used to describe the potential energy of the system. A force field is derived and optimized by validating simulation results with experiments and high-level quantum mechanical calculations. With reliable force field and modeling technique, molecular simulations can assist in predicting
various material properties and guiding experimental efforts for synthesis and characterization of new materials.

Two most popular approaches for molecular simulation are: stochastic approach (Monte Carlo) and deterministic approach (Molecular Dynamics). Additionally, there are several other techniques that work at different length and time scales and sometimes combine the features of MD and MC. The most distinctive feature of these molecular simulation techniques is their potential to evaluate material’s macroscopic thermodynamic properties such as internal energy, shear and tensile modulus, pressure, coefficient of thermal expansion, heat capacities, etc. Macroscopic properties at equilibrium are calculated using the time average of MD simulation and the ensemble average of MC simulation. The obvious benefit of MD over MC is that it gives route to the evaluation of transport (macroscopic) properties- also known as dynamical properties such as transport coefficient, rheology of the system, etc. using time correlation functions of corresponding microscopic variables. MD simulation makes use of optimized force field parameters to predict bulk properties of the system, whereas in MC the parameters are assumed. Hence, the simulation results obtained from MD can be applied to make comparisons with proposed theories and experiments directly, while it is not possible with the results from MC simulation.

Molecular dynamics simulation was first applied by Alder and Wainwright to study the phase transition of fluids (Alder and Wainwright, 1957). Theoretically in molecular modeling, the trajectories of particles/atoms are related to physical properties of the system by statistical thermodynamics. MD allows the prediction of time evolution of the material. The trajectory of positions and momenta of particles at different times can be used to evaluate all bulk physical properties of the system. Primarily, MD consists of three constituents: (1) initial configuration
and velocities of all the particles of the system, (2) the interaction potentials among all the particles, (3) trajectory of positions of the particles at different times calculated by solving standard Newtonian equation of motion (Zeng et al., 2008). The equation of motion is usually given as,

$$\vec{F}_i(t) = m_i \frac{d^2 \vec{r}_i}{dt^2} \quad (1.1)$$

where $\vec{F}_i$ represents the force acting on the particle $i$ at time $t$ and it can be derived from the potential energy $U(\vec{r})$; $\vec{r}_i$ is the position of particle $i$ having mass $m_i$. A physical simulation consists of the total potential energy that is a combination of all the interaction potentials among particles of the system, periodic boundary conditions and control of appropriate temperatures and pressures to satisfy conditions of different thermodynamic ensembles. All the interaction potentials together with corresponding parameter set which contribute towards total energy of the system form a force-field. The selection of appropriate force field depends on transferability, accuracy of parameters and total computational time. The total interaction potential can be broadly classified into two categories of bonded interactions and nonbonded interactions (Allen, 2004). Several terms involved in these two categories that sum up to total energy can be represented as:

$$U(\vec{r}) = \sum U_{\text{bond}}(\vec{r}_a, \vec{r}_b) + \sum U_{\text{angle}}(\vec{r}_a, \vec{r}_b, \vec{r}_c) + \sum U_{\text{torsion}}(\vec{r}_a, \vec{r}_b, \vec{r}_c, \vec{r}_d) + \sum U_{\text{vdw}}(\vec{r}_a, \vec{r}_b) + \sum U_{\text{electrostatic}}(\vec{r}_a, \vec{r}_b) \quad (1.2)$$

where $U_{\text{bond}}$ is the bond stretching energy; $U_{\text{angle}}$ is angle bending energy; $U_{\text{torsion}}$ is the dihedral angle energy; $U_{\text{vdw}}$ is the van der Waals energy; and $U_{\text{electrostatic}}$ is the electrostatic energy. The bond and angle terms, $U_{\text{bond}}$ and $U_{\text{angle}}$, do not allow covalent bonds to break. The torsion energy term, $U_{\text{torsion}}$, usually consists of two types of potentials: dihedral angle potential and improper potential. The former term is included to constrain the rotation of molecule around the specific
bond and the later term is used to maintain the planarity of atoms. The non-bonded terms, $U_{vdw}$ and $U_{electrostatic}$, are more computationally expensive because of the higher number of interactions involved. At the start of the simulation, atoms/particles are assigned initial velocities which contribute towards total kinetic energy of the system. This kinetic energy is dictated by the temperature of the system. Using the forces on each particle and temperature of the system, the acceleration of each particle can be determined.

Integration of all the equations of motion for the system yields to the formation of a trajectory having detailed information about positions and velocities of all the particles, total internal energy and temperatures and pressures of the system at different points of time. The integration of equations of motion over long time may be tedious and very complex at times. These equations can be integrated through many available algorithms using finite difference methods. All the available integration algorithms assume that the positions, velocities and accelerations of the particles can be approximated using Taylor series expansion.

The selected algorithm must comply with the characteristics of the equations of motion as given below.

(i) The selected algorithm must deal with short or long time-scales involved in the simulation.

(ii) The calculation of forces is mathematically expensive and time consuming. Thus, it should be performed at a frequency as low as possible.

(iii) The updating of atomic coordinates must follow calculation of dynamic properties accurately and assist sampling of correct ensemble.

(iv) The algorithm should also favor large time steps and thus they should not involve storage of large derivatives of positions and velocities.
The algorithm of Verlet satisfies most of these requirements. Commonly used algorithms in MD simulations are velocity verlet and leapfrog (Rapaport, 2001; Hairer et al., 2003). The Verlet algorithm calculates the position \( \vec{r}(t+\Delta t) \) and acceleration \( \vec{a}(t+\Delta t) \) of the particle at a time \( t+\Delta t \) by using the position \( \vec{r}(t) \) and acceleration \( \vec{a}(t) \) at present time \( t \) and the position of particles \( \vec{r}(t-\Delta t) \) at time \( t-\Delta t \). Whereas, in Leapfrog algorithm, the velocities are calculated at first at time \( t+\frac{1}{2}\Delta t \) and these are used to calculate positions of particles at time \( t+\Delta t \).

In case of polymers, the system size is very large. Most of the times the surface effects of the polymer with simulation box surface are neglected. In such cases periodic boundary conditions (PBC) are employed on the simulation box. It considers an infinite space filled with an array of exact replicas of the constructed simulation region. PBC comes with some consideration to satisfy periodicity. It assumes that an atom leaving the simulation box through a particular wall immediately re-enters the region from the opposite end. The interaction of one atom with another atom at a distance \( r_c \) within the box is same as that of the atom present at the same distance in the adjacent replica of the box. These constraints must be taken into account while dealing with equations of motion and new positions of atoms after each integration step. After each step, the atoms moving out of the boundary of the box must be brought inside from the opposite side and the coordinates must be adjusted accordingly. Evidently, PBC are most easy to handle in case of regions of rectangular dimensions; although it is not an essential requirement.

MD simulation can be carried out in many different ensembles. Ensemble is a collection of systems with different microscopic states but same macroscopic and thermodynamic states. Most common ensembles include microcanonical (NVE), canonical (NVT) and grand-canonical (\( \mu VT \)) where \( \mu \) is a chemical potential. Sometimes in practice, simulations need to be performed
at a constant temperature or pressure condition. The methods to incorporate these isothermal and isobaric conditions consist of restructuring of Lagrangian equations of motion. Some of the recognized methods for temperature control are Nosé-Hoover thermostat, the Berendson thermostat and Langevin dynamics. The methods for pressure control are the Berendson pressure bath coupling, Langevin piston method, etc.

**Macroscale modeling using computational fluid dynamics.** Nanopaint application using paint-spray technique is a complex multiphase system. The paint material is composed of resin, pigment, additives, solvents and nanoparticles. During the spray application substantial amount of solvents are released into the atmosphere in the form of volatile organic content (VOC) and the nanoparticles are released along with the paint droplets. Inside a spray booth, the excess of paint which does not get deposited on the substrate panel is removed by ventilation air passing through the booth ambience. In order to model this paint-spray process, multiple components and fluid phases need to be analyzed carefully and with precision. A successful design of the process depends on the accurate prediction of the interactions (chemical, mechanical and thermal) between these phases. Since this process is impossible to observe at a micro level, one has to rely on numerical modeling and simulation to gain insights into improving the process performance, environmental emissions, safety and reliability.

Numerical simulation can play an important role in analyzing such complex system problems. A discipline of computational fluid dynamics studies the numerical simulations and the solutions for the equations of motion of fluids. Computational Fluid Dynamics (CFD) is the science of predicting fluid flow, heat and mass transfers, chemical reactions, and corresponding phenomena by solving the equations governing these processes using a numerical process. It
provides a wide variety of methods for simulating the fluid flow problems. Through CFD, a qualitative and quantitative prediction of fluid flows can be made by means of

- Mathematical modeling (partial differential equations),
- Softwares and tools (various types of solvers and tools for pre- and post-processing of the system),
- Numerical methods (discretization and solution techniques).

CFD calculations are based on the first principles of mass, momentum and energy conservation. CFD modeling is capable of providing a detailed description of fluid flow variables, velocity, temperature and mass concentration profiles anywhere in the flow regime. Such details may not be possible to obtain from physical models and systems. In CFD modeling, the fluid flow region is divided into numerous small elements within which the flow is either kept fixed or varies smoothly. The equations of mass, momentum and energy balances are represented in terms of variables at the predetermined positions inside the elements. The solutions of these equations are evaluated until it reaches the required accuracy.

The working and use of CFD requires a basic set of steps to be followed. These steps are given below.

1. The physical problem to be studied should be defined.
2. The physical system should be designed by defining its geometry in either 2D or 3D space.
3. The conservation of mass, momentum and energy should be satisfied throughout the system’s region under consideration.
4. The properties of fluids involved in the system are modeled empirically.
5. Assumptions are made to simplify the problem and make it tractable (e.g. steady-state, transient, compressible, 2 dimensional, etc.).

6. Appropriate initial and final boundary conditions should be provided.

7. The system domain is discretized into a finite number of volumetric regions, called cells or grids. The discretized area is called the “mesh”.

8. The governing equations for mass, momentum and energy conservation are applied on the mesh and individual cells through numerical methods of discretization.

9. The post-processing of the solution is carried out to obtain the results for desired quantities (e.g. heat flow, mass fraction, temperature, drag, separation, pressure gradient, etc.).

The foundation for modeling of the fluid flow is provided by Navier-Stokes and continuity equations. The Navier-Stokes equations can be derived by the dynamic equilibrium of fluid elements. For compressible flow, the governing equations are the continuity equation, momentum equation (Navier-Stokes) and the energy equation.

The continuity equation is given as,

\[
\frac{\partial \rho}{\partial t} + \frac{\partial (\rho u)}{\partial x} + \frac{\partial (\rho v)}{\partial y} + \frac{\partial (\rho w)}{\partial z} = 0
\]  \hspace{1cm} (1.3)

The Navier-Stokes equations are given as,

\[
\rho \left( \frac{\partial u}{\partial t} + u \frac{\partial u}{\partial x} + v \frac{\partial u}{\partial y} + w \frac{\partial u}{\partial z} \right) = -\frac{\partial p}{\partial x} + \mu \left( \frac{\partial^2 u}{\partial x^2} + \frac{\partial^2 u}{\partial y^2} + \frac{\partial^2 u}{\partial z^2} \right) + F_x
\]  \hspace{1cm} (1.4)

\[
\rho \left( \frac{\partial v}{\partial t} + u \frac{\partial v}{\partial x} + v \frac{\partial v}{\partial y} + w \frac{\partial v}{\partial z} \right) = -\frac{\partial p}{\partial y} + \mu \left( \frac{\partial^2 v}{\partial x^2} + \frac{\partial^2 v}{\partial y^2} + \frac{\partial^2 v}{\partial z^2} \right) + F_y
\]  \hspace{1cm} (1.5)

\[
\rho \left( \frac{\partial w}{\partial t} + u \frac{\partial w}{\partial x} + v \frac{\partial w}{\partial y} + w \frac{\partial w}{\partial z} \right) = -\frac{\partial p}{\partial z} + \mu \left( \frac{\partial^2 w}{\partial x^2} + \frac{\partial^2 w}{\partial y^2} + \frac{\partial^2 w}{\partial z^2} \right) + F_z
\]  \hspace{1cm} (1.6)
The energy equation is given as,

\[
\rho c_p \left( \frac{\partial T}{\partial t} + u \frac{\partial T}{\partial x} + v \frac{\partial T}{\partial y} + w \frac{\partial T}{\partial z} \right) = \Phi + \frac{\partial}{\partial x} \left[ k \frac{\partial T}{\partial x} \right] + \frac{\partial}{\partial y} \left[ k \frac{\partial T}{\partial y} \right] + \frac{\partial}{\partial z} \left[ k \frac{\partial T}{\partial z} \right] \\
+ u \frac{\partial p}{\partial x} + v \frac{\partial p}{\partial y} + w \frac{\partial p}{\partial z}
\]

(1.7)

where \( \Phi \) is the dissipation function, \( u, v \) and \( w \) are the velocity components in \( x, y \) and \( z \) directions, \( \rho \) is the density, \( p \) is the pressure, \( T \) is the temperature, \( c_p \) is the specific heat at constant pressure and \( \mu \) is the viscosity.

The continuity equation represents the law of conservation of mass and thus must be satisfied at each point in the fluid region. In Navier-Stokes equations (Eqs. 1.4-1.6) the terms on the left hand side represent the inertial term which arise from momentum change. This term is compensated by the pressure gradient \( \left( \frac{\partial p}{\partial x} \right) \), the viscous forces and the body forces \( (F_b) \).

The importance of accuracy in multiphase modeling of sprays and droplets in various engineering applications is well known (Sirignano, 1999; Sazhin, 2006). Especially in the case of paint spray system, the model needs to take care of complicated fluid dynamics, multiple phases, heat/mass transfer, evaporation and deposition etc. In order to study all the characteristics of the coating, the application process of the paint material must be reproducible and well-controlled. With all the functionalities and abilities of CFD-modeling, it can provide a detailed view of the process, and study the effects of different operating conditions and geometries by simulating the corresponding flow behavior. It can offer significant insights into the coating process and show how the changes in workplace geometry, operating conditions, material type and application pattern can affect the quality of coating and performance of the system. With accurate modeling of this painting process, one can expect a number of
advancements in the understanding of the spray painting related issues, such as environmental emissions, paint transfer efficiency, coating quality, uniform film deposition, solvent evaporation, defects, and safety measure by assessing workers’ exposure to overspray paint, VOC, nanoparticles, etc. as a function of spray booth structure and ventilation system.

1.4 Towards Sustainable Nanocoating Technology Development

Nanotechnology, as a result of all the possible opportunities it provides for innovation, is finding innumerable ways to enter human’s life. It offers a promise to provide breakthrough technologies for a number of industries and consumer sectors with improved and novel functionalities and with a reduction in consumption of hazardous materials, consumption of energy as well as generation of wastes. Paint technology is no exception in accepting the ever increasing dominance of nanotechnology based products over conventional products. In fact, a number of nanotechnology based consumer products of paints and inks are already available in the market. However, the implications of nanomaterials and products on the environmental safety and human health are often either ignored or not highlighted. There is a major knowledge gap existing between the applicability of nano-size materials into consumer products and their effects on health and environment.

Nanomaterial is referred as a material with at least one component in the order of 1-100 nanometers. These materials can be individual nanoparticles of different shapes or aggregates of several nanoparticles together. In case of coating application there are numerous types of nanoparticles that are incorporated into polymer resin to synthesize final nanocomposite coatings. These nanoparticles include TiO₂, SiO₂, Ag, MMT clay, aluminium oxide, zinc oxide,
etc. They introduce improved and newer characteristics into nanocomposites such as improved mechanical, thermal, dielectric properties, biodegradability, anti-corrosion, self-cleaning, dirt repellency, anti-bacterial, etc (Zhang et al., 2005; Nobel et al., 2007; Chen et al., 2008). Presumably, this type of revolutionary technology should be sustainable in terms of economy, resource and energy efficiency and health care. However, so far only the economic prospect of nanotechnology has been highlighted and a very little attention is given to its social and environmental implications (Linkov et al., 2007; Dhingra et al., 2010). Its potential to develop systems with smart and newer functionalities significantly inspires competitiveness among different companies which use nanotechnology based coatings to avail all its economic benefits. Currently, the economic growth of the nanocoatings market and corresponding research and development gives a very little attention to assessment of social and ecological risks which are a part of complete holistic sustainability assessment of nanocoating products. Thus, it is important to stress on benefits and risks of this technology during the life cycle to detect all hidden short and long term adverse effects and to support all the decisions related to its future development. A holistic view for a sustainable development of nanocoating technology is represented in Figure 1.1.

Figure 1.1. Holistic view of sustainability.
A comprehensive study on the life cycle of nanoproducts can analyze, evaluate and address all the issues related to the environmental and health effects of nanoparticle induced coating materials. The life cycle based study, as proposed by EPA, considers all the stages of nanocoating’s life from “cradle-to-grave” which include, (1) material selection, (2) system’s design, (3) use and maintenance, (4) recycling and reuse, and (5) disposal.

Figure 1.2 broadly classifies these various stages of nanopaint life with applicable EPA regulations.

Figure 1.2. Life cycle of a nanopaint-nanocoating system

In general, it is very challenging to perform complete sustainability assessment of emerging or developing technologies (e.g. nanocomposite coatings) due to insufficient data availability for inputs and outputs of the system at each stage of life cycle. However, if succeeded, it can provide significant amount of supplementary information to support decisions related to the future development. LC based sustainability assessment can provide answer to the questions such as,
• How the performance of nanocoating is compared with conventional coating over its life-cycle?
• How much energy efficiency is accomplished by incorporating nanoparticles into conventional products?
• Which stage of life cycle dominates the energy consumption?
• Which stage of life cycle is most prone to the release of nanomaterials in the environment?
• What are the toxicity issues involved with released nanomaterials at different phases of life cycle?
• What is the impact of nanoparticle coatings compared to those of conventional paint products on geographical parameters?
• How is the end-of-life management of nanopaints compared to that of conventional paints? Is there a way for reuse or recycling?

The answers to these questions and development of a comprehensive life-cycle based sustainability assessment methodology for analysis of nanocoating technology can significantly assist in directing the research and sustainable development of these paint products.

1.5 Main Goals and Scientific Contributions

The objective of this research is to develop a comprehensive multiscale design and development tool to aid the development of sustainable nanocoating technology which can deliver products with multiple functionalities. The challenge to this research comes from the lack of thorough knowledge about the vast and complex microstructure, huge design space and
substantially limited data availability. The existence of vast design parameters and experimental complexity makes nanopaint design optimality extremely difficult to address if not impossible. Thus, the focus of research on nanocoating technology is being shifted towards the optimal design formulation of nanocoatings, where the need for computational modeling and simulation methodologies is inevitable. *In silico* experiments using computational material designs of nanocoating resins have immense potential to facilitate identification of important correlations among material, structure, property and performance. It can assist the scientific research for the development of more sustainable multifunctional nanocoatings, the application methodologies and product design.

In order to meet the goal of developing a methodology to aid the research on sustainable nanocoating materials, we integrate the top-down, goals/means, inductive systems engineering and bottom-up, cause and effect, deductive systems engineering approaches, as shown in Figure 1.3 (McDowell and Olson, 2008), and develop a complete multiscale framework which can define a correlation among material-structure-property-performance. Previously, through top-down approach, the nanocoating system was studied by using modeling and simulation at meso- and macro-scales of length and time (Xiao *et al.*, 2007; Xiao *et al.*, 2010; Xiao and Huang, 2009). This research focuses on bottom-up approach to develop nano- and micro-scopic models and connect it to the already developed mesoscopic modeling methodology to complete the multiscale materials development framework.
For the process, product design and mesoscale development of paint material, the models based on CFD simulation technique and multiscale Monte-Carlo based simulation technique have been developed (Xiao et al., 2008; Xiao and Huang, 2009; Xiao et al., 2010; Xiao et al., 2010). The CFD based modeling provides crucial knowledge about paint film application technique and operating parameters related to paint spray and curing, and the Monte Carlo based modeling provide intrinsic relationship among nanopaint components and bulk properties through development of coarse-grained (CG) freely jointed bead-spring structures. Although, the bead-spring model is useful to understand the correlation between coating microstructure and bulk properties, it lacks the details from specific chemistry of polymers. Thus, the properties of
nanoparticle-induced coatings with different materials of resin cannot be studied. This chemical specificity can be provided through structure and thermodynamics based coarse-grained modeling. In this work, via bottom-up approach, we develop chemistry-specific CG nanocoating material designs and integrate it with the previous accomplishments to produce a complete multiscale framework that can lead to the development of optimal multifunctional nanocoating formulation using assessment of the correlation of material-property-product-processing.

The introduction of unique and improved properties of the paint material through nanotechnology may also have environmental and societal implications which may lead to greater risks to human health and safety. Thus, the environment, health and safety impact assessment of existing and emerging nanocoating materials has become a serious issue. In this research, a life-cycle-based sustainability assessment framework is developed for the nanocoating technology which can highlight the potential problems that this technology may cause to the environment, society and economy. For the assessment, five stages of cradle-to-grave life-cycle of a paint material (material selection, paint manufacturing, paint spray/flash, curing and end use) have been considered. Based on the data available from literature, research papers, industries and computational models related to paint technology, individual sets of parameters have been identified for each stage of the life-cycle which aid to the overall sustainability assessment of each stage. The sustainability indicator matrices for each stage encompass all the aspects concerning the economic benefits, environmental issues and societal impact of that stage. Using case studies of automotive coating systems, the assessment results are combined together and analyzed to identify critical parameters which influence the overall nanopaint technology sustainability. The analysis of each stage of the life cycle using sustainability aspects and developing a correlation among different parameters and indicator
matrices related to environmental, social and economic aspects can significantly direct the future development of these products towards sustainability.

1.6 Organization of Dissertation

The dissertation body is mainly composed of two sections. The first section describes the multiscale computational modeling effort for polymer-nanocomposite coatings study and the second section focuses on the development of life-cycle based sustainability assessment framework and paint process modeling to obtain the data for the assessment.

In Chapter 2, the multiscale computational design of nanopaint is developed. The nanopaint is modeled on atomistic level by using CHARMM force-field. It is then mapped onto the coarse-grained structure which is developed by applying MARTINI force field protocols. The developed coarse-grained system is then used for the study of effects of different sizes, volume fractions and distributions of nanoparticles and the polydispersity of polymer resin on the structure-property correlations and interfacial behavior of resin molecules. In Chapter 3, the efforts made towards the experimental verification of multiscale computational design of nanopaint are described. As a case study, the polymer nanocomposite films are synthesized by layer-by-layer application of acrylic resin and TiO₂ nanoparticles dispersion. These films are characterized and tested through experimental analysis techniques such as XRR and AFM to determine the change in the mechanical stiffness of the films after addition of different dispersions of nanoparticles.

The later Chapters are focused on the development of life-cycle based sustainability assessment (LCSA) methodology and the macroscale modeling of paint application process to obtain the data for nanocoating technology assessment. In Chapter 4, a comprehensive LCSA
methodology is described. The parameter sets and corresponding sustainability indicator matrixes are developed for each of the 5 stages of life-cycle which include material selection and preprocessing, paint manufacturing, paint application and curing, use and disposal. The applicability of the methodology is demonstrated with a case study on a set of coating systems assessment. Chapter 5 and Chapter 6 provide the details of the modeling and simulation of paint application process (automotive paint-spray). The modeling of paint-spray technique is performed in order to study the effects of nanoparticles addition into coating matrix on the environmental emissions inside the spray-booth and coating film quality parameters. The topological characteristics of the paint film are also studied by developing case studies of various paint spray patterns and application parameters. The data obtained in these chapters could be used for the quantification of some of the sustainability indicators described in Chapter 4.

Finally, the concluding remarks and possible directions to extend this work in the future are outlined in Chapter 7.
CHAPTER 2
MULTISCALE MODELING AND ANALYSIS OF POLYMER NANOCOMPOSITE SYSTEMS

With the advancement of computing power, it is evident that computational modeling tools are going to accelerate the development of nanocomposite coating technology for all the superior properties achievable. The ability of having a complete control on nanocoating microstructure and complex behavior of polymer matrix makes the research using computational methods more significant. However, it is the selection of appropriate modeling techniques and mathematical algorithms that ensures the generation of meaningful information out of simulation. Nanopaints being materials with applications at continuum level and properties defined by microstructures, it is critical to generate intrinsic correlations among coating’s material, structure, property and performance.

2.1 Objective and Significance

It has been proven that the interface between polymer and nanoparticle plays a key role in introduction of superior and newer functionalities into nanocoatings. Besides the interface, the chemical nature of polymeric material, type and morphology of nanoparticles are amongst other important parameters to bring along the property enhancement. An extensive experimental study is underway to determine all critical factors responsible for these superior properties as well as
newer functionalities of nanocomposite coating matrices. The enhancement of mechanical and rheological properties of polymer nanocomposites is demonstrated by addition of different types of nanoparticles such as MMT clay, silica, TiO$_2$, etc. into polymer resins (Van Hamersvelt et al., 1999; Tigli and Evren, 2005; Chow and Mohd Ishak, 2007; Nobel et al., 2007; Chen et al., 2008). A work is also in progress to bring-in newer functionalities like self-cleaning and smart corrosion resistance into coating materials (Zhang et al., 2005; Radhakrishnan et al., 2009).

Despite of growing knowledge about nanocomposite coatings through experiments, there are several structure and property correlations that are unanswered and still being investigated. With the development and availability of computing power, computational modeling and simulation has become a very useful tool to study and address these issues. Over the years, simulation techniques are developed and applied to various systems such as polymers and biomolecules at multiple scales of length and time ranging from nanoscopic to macroscopic continuum levels. Simulations on atomistic level provide chemical specificity to molecular models using empirically verified and generic force fields. But the design complexity and huge design space of polymer melts at atomistic level brings along several limitations of time and length scales. It is presently very difficult to model the behavior of polymer melt around nanoparticles at atomistic levels that requires tens of microseconds of time at tens of nanometers of length for equilibration.

Coarse-graining approach can help in overcoming this limitation of atomistic modeling. Kremer and Grest proposed the popular bead-spring model for simulation of long chain molecules (Kremer and Grest, 1990). It has been applied for CG modeling of large molecules like polymers by several research groups (Starr et al., 2002; Smith et al., 2003; Kalra et al., 2010; Xiao et al., 2010). Although the bead-spring model is applicable for understanding the
sciences behind polymer-nanoparticle surface interaction leading to superior properties, it lacks the details from specific chemistry of polymers. Thus, it cannot be applied for studying the properties of coating with different materials of polymers. This leaves a clear gap between two scales of simulation and makes it almost impossible to develop a correlation between atomistic and CG modeling. In order to incorporate chemical specificity into CG models, the development of a structure based or thermodynamics based coarse-grained models is essential. This type of CG modeling involves grouping of atoms from all-atom model into larger size beads. Thus, it decreases the total number of atoms/particles in the system to be simulated. As a result of reduced number of degrees of freedom, the CG models usually run much faster with larger time steps during molecular dynamics (MD) simulations compared to atomistic models.

In order to prepare a comprehensive multiscale modeling tool for the study of nanocomposite coating material, it is critical to develop a well-defined CG model of selected polymer resin system that can be transferrable over different scales of time and length. In our work, we adopt a thermodynamics based CG modeling technique called “The MARTINI force-field method” which was introduced by Marrink et al. (2004). It was originally developed for lipid bilayers and detergent molecules. Later it was extended for coarse-graining of organic solvents, proteins and carbohydrates (Marrink et al., 2007; Monticelli et al., 2008; Lopez et al., 2009). The applicability of the MARTINI approach for polymers has been tested and verified over specific cases of polyethylene glycol, polystyrene and thermoset polyester coatings (Lee et al., 2009; Rossi et al., 2011a,b). This approach consists of verification of bonded and non-bonded parameters involved to define macromolecular structures using the data from atomistic scale modeling. The accuracy of this model can be shown by comparing the structural and
thermodynamic properties such as radii of gyration, pair distribution functions, etc. with that from all-atom models.

A significant attention is being given towards the development of water-based nanopaint resin systems to meet continuously increasing demand for low VOC and environmentally friendly materials. Acrylic coatings are water-dispersible and have applications in many different types of coatings. Consequently, a research focus is gradually shifting towards developing optimal formulations of nanocoatings that have acrylic-based resins. Needless to say that the contribution from molecular simulation can be significant in the developmental studies of these types of nanocoating materials due to a large number of complex structural features involved. For this purpose, a multiscale model of an acrylic nanocomposite material is required. Previous accomplishments of Dr. Huang’s group covered the modeling and simulation of nanocoating material on the mesoscopic level. In order to build a complete multiscale modeling framework which can generate crucial information about nanocoating’s material-structure-product-process correlation, the simulation on the finer level is required. The nanoscopic and microscopic simulations then need to be bridged to the previously developed mesoscopic methodology to accomplish the objective of development of a comprehensive multiscale modeling framework for the study of nanocoating technology. This major task involves five distinct steps of simulation work which are to be covered.

1. Generation of the atomistic model of an acrylic polymer system using a wisely selected force-field,
2. Development of the MARTINI parameter set to model the acrylic polymer at CG level and verifying it using atomistic simulations and experimental data,
3. Incorporation of nanoparticles of selected size and morphology.
4. Use of the final polymer-nanocomposite model to predict structural features and bulk properties of the nanocoating material and to explore critical parameters that contribute towards enhancement of the material performance.

5. Building a bridge between MARTINI-based simulation model and previous Monte Carlo-based simulation model.

After accomplishing the objective to develop a multiscale molecular modeling methodology to simulate a polymer-nanocomposite coating, it can be very useful to learn and discover new knowledge about the microstructure-property-performance correlation among paint resin and dispersed nanoparticles. It can help experimentalists to prepare optimum formulations with controlled enhancement in target properties. The findings from life-cycle based sustainability assessment work can be employed easily to predict property changes of the coating material with accuracy and with lower cost and time.

2.2 Atomistic Modeling

The optimization of MARTINI force field parameters significantly relies on accuracy of atomistic modeling. The atomistic modeling uses data from generic force-fields for simulation of molecules. By selecting appropriate force field, atomistic simulation is capable of predicting a number of structural and thermodynamic properties of a molecular system. In case of polymeric material the size of the system, length and number of polymer chains and number of atoms in the system decide the time required for equilibration under given conditions of ensemble. Usually the time step for this simulation is very small. The validity of the atomistic model is justified by
reproducing empirical polymer melt densities and certain structural properties such as radii of gyration, mean squared distances, etc.

2.2.1 Force Field Selection and Optimization

The selection of appropriate force field depends upon several factors such as the quality, applicability to the selected molecular structure and ability to predict target properties with accuracy. The force field stores all the necessary information for the calculation of forces and energies. Typically the force field should consist of following information: (i) atom types, (ii) partial atomic charges, (iii) functional forms of equations to represent energies, (iv) corresponding parameter values needed for all functional equations, (v) rules to generate new parameters for molecules that are not explicitly defined, and (vi) ways of assigning different types of functional forms and corresponding parameters.

The basic functional form expressing total energy of the system forming the force field is the sum of bonded (covalently) interactions and nonbonded interactions (Van der Waal’s and electrostatic interactions). The bonded energy, $E_{\text{bonded}}$, generally accounts for bond stretching ($E_{\text{bond}}$), angle bending ($E_{\text{angle}}$) and dihedral torsion ($E_{\text{dihed}}$) terms.

$$E_{\text{bonded}} = E_{\text{bond}} + E_{\text{angle}} + E_{\text{dihed}}$$  \hspace{1cm} (2.1)

The nonbonded energy, $E_{\text{nonbonded}}$, accounts for Van der Waals ($E_{\text{vdw}}$) and electrostatic ($E_{\text{coulomb}}$) interaction terms.

$$E_{\text{nonbonded}} = E_{\text{vdw}} + E_{\text{coulomb}}$$  \hspace{1cm} (2.2)

The acrylic polymer selected for this work is polymethyl methacrylate (PMMA). There are several classical force fields that are suitable for the simulation of PMMA model. These
include CHARMM, AMBER, CVFF, COMPASS and PCFF. The force field selected for atomistic modeling of PMMA is an extension of CHARMM (Chemistry at HARvard Macromolecular Mechanics). CHARMM was originally developed by Dr. Alex MacKerell (Feller et al., 1997; Guvench and MacKerell, 2008). It was parameterized on the basis of *ab-initio* and experimental data on energies and geometries of small organic molecules, vibrational spectra and torsional rotational surfaces.

There are several different versions of CHARMM that are released till date. The most commonly used are CHARMM19 (Feller et al., 1997), CHARMM22 (MacKerell et al., 1998) and CHARMM27 (Foloppe and MacKerell, 2000). In CHARMM force-field, potential energy functions for bonded interactions are expressed as harmonic potentials. The total bonded energy term can be written as:

\[
E_{bonded} = \sum_{bonds} K_b (b-b_0)^2 + \sum_{angles} K_\theta (\theta-\theta_0)^2 + \sum_{dihedrals} K_\phi (1 + \cos (n\phi-\delta)) \\
+ \sum_{improper} K_\omega (\omega-\omega_0)^2 + \sum_{Urey-Bradley} K_u (u-u_0)^2
\]  

(2.3)

where the first term represents harmonic bond potential and \(K_b\) and \(b_0\) are the force constant and equilibrium bond length respectively. The second term in the equation stands for the harmonic angle potential where \(K_\theta\) and \(\theta_0\) denote angle constant and equilibrium angle in degrees. The third term accounts for the dihedral angle potential with torsional force constant \(K_\phi\), dihedral angle \(\phi\) in degree, \(n\) is the multiplicity and \(\delta\) is the phase shift. Impropers are used to restrict the molecular geometry. This fourth term corresponds to out of plane bending where \(K_\omega\) is the stiffness with equilibrium angle \(\omega_0\). The fifth Urey-Bradley potential term accounts for interaction between the 1\(^{st}\) and 3\(^{rd}\) atom, separated by two harmonic bonds, where \(K_u\) is the force constant and \(u_0\) is the equilibrium distance between 1, 3 atoms in the harmonic potential.
The nonbonded potential functions are represented by the standard 12-6 LJ potential and electrostatic interactions as shown below:

\[ E_{\text{nonbonded}} = \sum_{\text{nonbonded pairs}} \epsilon \left[ \left( \frac{R_{\text{min}_{ij}}}{r_{ij}} \right)^{12} - 2 \left( \frac{R_{\text{min}_{ij}}}{r_{ij}} \right)^{6} \right] + \sum_{\text{nonbonded pairs}} \frac{q_i q_j}{r_{ij}} \]  

(2.4)

where the first term accounts for van der Waals force of interaction between a pair of atoms represented by the 12-6 LJ potential. \( R_{\text{min}_{ij}} \) is the equilibrium distance between two atoms at a point where the LJ potential becomes zero. The last term stands for the electrostatic force of interaction between atoms with partial atomic charges (MacKerell et al., 1998).

Figure 2.1 shows the proposed PMMA repeating unit model which is used in all the atomistic simulations. The molecular weight of each repeating unit of PMMA is 100 g/cm\(^3\).

![PMMA repeating unit](image)

Figure 2.1. PMMA repeating unit.

**Partial atomic charges.** CHARMM27 does not have an optimized set of partial charges directly applicable to PMMA molecules. Thus, partial charge on each atom of a PMMA repeating unit is determined using the CHelpG (charges from electrostatic potentials using a grid based method) type of atomic charge calculation scheme. Each PMMA chain was constructed with three types of residues, which are the derivatives of a molecular structure formed by capping both ends of the methyl methacrylate monomer with methyl groups. The
geometry of the PMMA monomer was optimized using Hartree Fock (HF) theory and 6-31G+(d,p) basis set. Calculations were started from a variety of initial conformations to ensure the optimized state corresponded to the global energy minimum (Breneman and Wilberg, 1990). This optimized geometry was used for the determination of partial charges via the CHELPG method (Breneman and Wilberg, 1990). The partial atomic charges on each atom of a PMMA repeating unit formed by two different initial conformations are given in Table 2.1. Conformation 2 led to the minimum energy of the optimized state of the system as compared to that obtained from Conformation 1. Thus, the corresponding set of charges was selected for further calculations.

Table 2.1. Partial atomic charges on MMA unit obtained after Gaussian calculations

<table>
<thead>
<tr>
<th>Atom name</th>
<th>Partial charges with Conformation 1</th>
<th>Partial charges with Conformation 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>C1</td>
<td>-0.20</td>
<td>-0.12</td>
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<tr>
<td>C2</td>
<td>0.09</td>
<td>0.16</td>
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<tr>
<td>C3</td>
<td>-0.15</td>
<td>-0.23</td>
</tr>
<tr>
<td>C4</td>
<td>0.88</td>
<td>0.87</td>
</tr>
<tr>
<td>C5</td>
<td>-0.12</td>
<td>-0.02</td>
</tr>
<tr>
<td>H1</td>
<td>0.07</td>
<td>0.01</td>
</tr>
<tr>
<td>H2</td>
<td>0.07</td>
<td>0.01</td>
</tr>
<tr>
<td>H3</td>
<td>0.04</td>
<td>0.05</td>
</tr>
<tr>
<td>H4</td>
<td>0.04</td>
<td>0.05</td>
</tr>
<tr>
<td>H5</td>
<td>0.04</td>
<td>0.05</td>
</tr>
<tr>
<td>H6</td>
<td>0.05</td>
<td>0.08</td>
</tr>
<tr>
<td>H7</td>
<td>0.05</td>
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</tr>
<tr>
<td>H8</td>
<td>0.05</td>
<td>0.08</td>
</tr>
<tr>
<td>O1</td>
<td>-0.64</td>
<td>-0.65</td>
</tr>
<tr>
<td>O2</td>
<td>-0.51</td>
<td>-0.42</td>
</tr>
</tbody>
</table>

The partial atomic charges for the three residues of PMMA were derived from the charges determined for the reference molecular structure and with 2nd initial conformation of atoms (see Figure 2.1). One type of residue was used to represent MMA repeat units located
inside each polymer chain, whereas the other two types formed capping units. The final set of atomic charges for each residue was obtained through normalizing the atomic charges of the structure in Figure 2.2 and by keeping the total charge on each residue equal to zero.

Figure 2.2. Partial atomic charges obtained from Gaussian after optimization of MMA monomer capped with methyl groups.

**Dihedral optimization.** All the bonded and non-bonded potential parameters for PMMA modeling were obtained from CHARMM27 force field developed for lipid molecules. Getting the non-bonded parameters for new molecules with the use of analogy to previously parameterized molecules can often be well assigned. However, the bonded parameters are capable of posing more difficulty as the possibility of having unparameterized connectivity among atoms become larger for bonds, angles and dihedral angles (Guvench and MacKerell, 2008). In case of PMMA, though it was possible to get the parameterized connectivity of bonds and angles, the dihedral potentials still needed to be optimized. It was achieved by using CHARMM general FF optimization methodology (Guvench and MacKerell, 2008; Vanommeslaeghe et al., 2010). Typically the target data selected for fitting the dihedral angle
potentials are energies from QM (quantitative mechanical) adiabatic potential energy scans. For a dihedral angle A-B-C-D, a relaxed potential energy scan is done by rotation about the bond B-C over 360 degrees at fixed increments in QM representation. The same scan is also done in the MM (molecular mechanics) representation by using all-atom CHARMM force field dihedral parameters to be optimized. Rotational energy barrier plots are then obtained by subtracting the potential energy of the lowest energy conformation from all higher energy conformations at remaining angles. The set of force field parameters that would give an exact match between the plots of rotational energy barriers from QM and MM are chosen for the selected dihedral A-B-C-D.

For optimization of the PMMA structure, we selected two dihedral angles. One dihedral was selected along the main chain backbone (C2-C1-C2-C3) and the other along the side chain of PMMA (C2-C4-O2-C5) to ensure optimization over the complete PMMA structure. In order to obtain potential energy scans at QM and MM levels, a trimer of PMMA was selected with both chain ends capped with hydrogen atoms. The selection of a trimer structure instead of a dimer ensured mimicking the behavior of monomers located in long chains where the corresponding dihedral angles would encounter steric hindrance during rotation over 360 degrees. Thus, it was possible to extract accurate rotational energy barrier values which were higher as a result of steric hindrance from neighboring monomers. Note that rotational energy barrier values of dihedral angles from dimer structure were smaller compared to that of trimer structure. The potential energy scans at QM level were performed by optimizing the trimer structure at each dihedral increment with HF/6-31+G(d) level of theory calculations. Whereas at MM level, potential energy scans were obtained over both dihedrals by optimizing the trimer structure using the CHARMM force field set of parameters to be optimized. The molecular
The simulation package used for performing MM level scans was CHARMM, version 34. The trimer structure was equilibrated at each increment of rotation and final rotational energy barrier plots were obtained. Figure 2.3 shows both dihedrals of trimer structure selected for potential energy scans. Figures 2.4a and b represent the final match of rotational energy barrier plots from QM and MM level calculations for both dihedrals C2-C1-C2-C3 (along the backbone) and C2-C4-O2-C5 (along the side chain) using optimized parameters for PMMA.

The graphs of rotational energy barriers show that minimum energy conformations are observed at ±60 degrees of dihedral angle C2-C1-C2-C3 and ±180 degrees of dihedral C2-C4-O2-C5. Figure 2.5 shows these stable conformations of PMMA trimer which should appear with highest probability in atomistic simulation trajectories. The final optimized set of CHARMM force field parameters of bonded and nonbonded interactions for modeling of PMMA are given in Table 2.2.

Figure 2.3. PMMA trimer used for determination of rotational barriers for C2-C1-C2-C3 and C2-C4-O2-C5 dihedrals.
Figure 2.4. Rotational energy barriers predicted by QM and MM calculations for dihedrals (a) C2-C1-C2-C3, (b) C2-C4-O2-C5, (c) C2-C1-C2-C3 and (d) C2-C4-O2-C5.

Figure 2.5. Minimum energy conformations for the PMMA trimer with: (a) dihedral C2-C1-C2-C3 fixed at -60 degrees, and (b) dihedral C2-C4-O2-C5 fixed at 180 degrees.
Table 2.2. CHARMM force-field parameters for the modeling of PMMA.

(a) Bond parameters

<table>
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<tr>
<th>Bond</th>
<th>$K_0$</th>
<th>Bond length ($b_0$)</th>
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</tr>
<tr>
<td>C1-H2</td>
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</tr>
<tr>
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<td>1.111</td>
</tr>
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</tr>
<tr>
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</tr>
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(b) Angle parameters

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<td>n/a</td>
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<tr>
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<td>113.5</td>
<td>11.16</td>
<td>2.561</td>
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<tr>
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<td>113.5</td>
<td>11.16</td>
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<tr>
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### (c) Dihedral parameters

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<td>180</td>
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<tr>
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<td>C1-C2-C3-H4</td>
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### (d) Non-bonded LJ parameters

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<th>$\epsilon_{1-4}$</th>
<th>$R_{min/2,1-4}$</th>
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</thead>
<tbody>
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<td>-0.01</td>
<td>1.9</td>
</tr>
<tr>
<td>C2</td>
<td>-0.020</td>
<td>2.275</td>
<td>-0.01</td>
<td>1.9</td>
</tr>
<tr>
<td>C3</td>
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<td>1.9</td>
</tr>
<tr>
<td>C4</td>
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<td>n/a</td>
</tr>
<tr>
<td>C5</td>
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<td>-0.01</td>
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</tr>
<tr>
<td>O1</td>
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<td>1.700</td>
<td>-0.12</td>
<td>1.4</td>
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</table>
2.2.2 Polymer Matrix Simulation

All of the atomistic simulations of PMMA model were performed using optimized set of CHARMM force field parameters. The initial configuration consists of atomic coordinates for all of the atoms located in the simulation box. The polymer chains are placed in the simulation box at various places. The size of the box has to be adequate enough to obtain the density of the material comparable with the experimental value.

The PMMA initial configuration was constructed using molecular packaging software PACKMOL (Martinez et al., 2009). The molecular topology file needed for the construction of PMMA chains defines three residue types as explained in previous section. One residue type represents monomer repeating unit inside each polymer chain and the other two residues are used to patch each polymer chain at the ends. Initial model consisted of 15 atactic chains of PMMA having 50 repeating units in each. The total number of atoms in the system were 11,280. The chains were positioned randomly inside the simulation box. The size of the box was selected in a way to keep the initial density of the system to 0.80 g/cm³.
Simulation details. The amorphous PMMA structure is designed in the simulation box with periodic boundary conditions. The molecular dynamics package used for performing all simulations is NAMD, version 2.8. Each input file required for NAMD stores a specific information about the system and simulation conditions. Typically the files needed are PDB, PSF, configuration file and parameter file. PDB (Protein data bank) file stores all the atomic coordinates with their names and residue types. PSF file stores all topological information about the molecule. The parameter file stores all the force field parameters for a selected molecule in CHARMM format. The configuration file provides all the details required to carry out the simulation at given conditions of temperature and pressure and at specified time step for specified run time. The NAMD output stores all atomic coordinates and velocities in the DCD file. All the information about forces and energies of the system are collected in the “log” file at each specified timestep.

In case of PMMA melt simulation, the system was equilibrated at temperature of 500 K and pressure of 1 atm for 40 ns in the NPT ensemble. The timestep for equilibration was 2 fs. A cutoff of 12 Å was used for the calculation of all nonbonded interactions during simulation. Switching was kept on with switching distance fixed at 10 Å. The pairlist distribution was set at 14 Å. The temperature during NPT ensemble was controlled with Langevin thermostat (damping coefficient 5 ps⁻¹) and the pressure was kept constant by applying Noé-Hoover Langevin piston pressure control. The values of piston period and piston decay were set at 100 fs and 50 fs respectively. The simulation was run for 20 ns further to extract data for measurement of all the properties.

Atomistic simulations provided validation to the optimized set of dihedral parameters by calculating the distributions of both dihedral angles from equilibrated trajectories. Figures 2.4 c
and d give the plots of normalized probability distribution of both selected dihedral angles. For dihedral C2-C1-C2-C3, the distribution is highest at angles ±60 degrees and for dihedral C2-C4-O2-C5 the distribution was highest ±180 degrees. These distributions indicate the appearance of conformations having lowest rotational energy barriers with highest probability.

### 2.2.3 Atomistic Model Verification

The atomistic model works as a benchmark to validate CG simulation. Thus, it is important to have this model accurately reproduce structural and thermodynamic properties of the polymer. For the modeling of PMMA structure, the nonbonded parameters are selected through analogy with similar molecular structures. The verification of these nonbonded LJ parameters for PMMA model is done by calculating thermodynamical properties of PMMA and comparing it with the empirical data.

The target properties selected for the verification were density of PMMA at room temperature and glass transition temperature. PMMA system was initially equilibrated under melt conditions at temperature of 500 K. It was then cooled gradually at intervals of 50 K. After cooling, the system was equilibrated under NPT ensemble for 5 ns and further for 2 ns to generate the data for analysis. The cooling was carried out until 150 K. The density of the system was measured after every temperature interval by averaging over the trajectory of last 2 ns of simulation.

The density of PMMA at 300 K was found to be 1.154 g/cm³ which was in good agreement with the bulk amorphous PMMA density of 1.17 g/cm³. The glass transition temperature of the simulated amorphous PMMA model was calculated using the density data at
each temperature of simulation. The increment in the specific volume is less pronounced below $T_g$ compared to above $T_g$ due to the onset of glassy state of the polymer. The point of intersection of two lines representing two rates of increments of specific volumes is marked as $T_g$ (Han et al., 1994). From the Figure 2.6, it was measured to be 370 K and it is in good agreement with the literature data value of 378 K (Condo and Johnston, 2004; Goldstein et al., 1993).

![Figure 2.6. Determination of the glass transition temperature ($T_g$).](image)

2.3 **Coarse-Grained Modeling of Polymer Resin**

Simple CG models of polymers with freely jointed, bead-spring structures have been used successfully at many cases to understand the general features of polymers and study its structure and dynamics. Xiao et al., 2010 used this type of model for the development of CG thermoset polymer nanocomposite system using Monte-Carlo simulation scheme to study their mechanical properties. It has been the only effort to model a thermoset type of polymer
nanocomposite system till date. The structure of polymer beads around platelet shaped nanoparticles exhibited the formation of structured pattern leading to substantial increase in its number density on the nanoparticle surface (Xiao et al., 2010). Through molecular dynamics simulation technique, Starr et al. studied structural changes and dynamic properties of polymer chains as a function of surface interaction between icosahedral shaped nanoparticles and polymer beads (Starr et al., 2002). Polymer chain length also plays a crucial role in regulating surface interactions. An application of shear force alters the rate of aggregation of nanoparticles and is a strong function of polymer chain length, as discussed by Kalra et al. (2010).

The objective of the project is to develop a comprehensive computational methodology to study microstructural behavior of polymer nanocomposites. In order to construct the model applicable to polymeric materials with specific chemistry, a CG model with structural and thermodynamical basis is required. It is accomplished by adopting the MARTINI force-field approach to develop the CG model. The applicability of this approach on polymer systems has been proven to be accurate for several examples of polymeric systems. The CG parameters optimized for polyethylene oxides model are transferable over different polymer molecular weights and chain lengths. The polystyrene model developed by Rossi G. et al. (2011) is in good agreement with empirical data and atomistic simulations in terms of structural properties, temperature transferability and compatibility with different solvents. In our case of acrylic polymer system, the optimization of interaction parameters among polymer beads are also performed by comparing with atomistic simulation results.

2.3.1 Methodological Framework
As explained earlier, the reliability of coarse-grained models is highly dependent on atomistic level simulations. CG model generally reproduces only the important degrees of freedom related to molecular structure. The accuracy of CG model in reproducing structural data from all-atom model is decided by the bonded and non-bonded parameters to define the system under consideration. A pathway which is followed to accomplish CG modeling of any molecular system is shown in the following figure.

![Coarse-graining algorithm](image)

Figure 2.7. Coarse-graining algorithm.

The interactions among ‘super-atoms’ from CG models are mainly classified into two categories as, bonded and non-bonded. The bonded interaction parameters are obtained by comparing the distributions of these interactions from both atomistic and CG simulations. Similarly, there are a number of objective functions which can be used to obtain non-bonded
parameters depending upon the type of CG model to be developed. In case of thermodynamics based CG models, the target properties for optimization of CG parameters are generally density, radius of gyration and mean square displacement of polymer chains. Whereas, in case of structure based CG modeling it is the radial distribution function. In developing MARTINI model, the target properties in the parameterization of bonded interactions are distributions of bonds and angles from atomistic simulations. Nonbonded interactions are optimized using melt density and radius of gyration of the polymer melt.

2.3.2 Model Development and Optimization

According to MARTINI protocols, the method of coarse graining consists of three steps; mapping, parameterization of bonded and non-bonded interactions (Lee et al., 2009; Rossi et al., 2011).

Mapping. Mapping step of coarse-graining consists of identification of groups of atoms to be represented as beads or “super-atoms”. In order to keep the PMMA model simple, each all-atom repeat unit is mapped onto two interaction sites in CG model. The interaction sites are represented as A and B. The selection of bead types is based on chemical similarity between MARTINI beads and groups of atoms from PMMA structure. When compared to the nomenclature of the MARTINI force-field, beads A and B are analogous to the bead types C1 and Na (Marrink et al., 2007). The bead type C1 is used to represent a site with very low polarity. Similarly the bead type Na represents a non-polar site and it can be used to map on the chemical group –COOCH₃. This analogy is critical for the selection of initial guess for non-bonded potential parameters for each bead. The schematic of the mapping of PMMA repeating
unit is shown in Figure 2.8. The size of the bead type B having four-to-one mapping is selected as 4.7 Å. The bead type A consists of three-to-one mapping where only three heavy atoms of carbon are mapped onto a single bead. The selection of masses and sizes of these beads are dependent on the performance of the model in terms of densities and structural properties.

![Figure 2.8. Schematic representation for mapping atoms in (a) to coarse-grained beads in (b) for PMMA.](image)

**Parameterization of bonded interactions.** The CG system of PMMA contained 15 chains with 50 repeating units in each chain. The polymer was in the melt state at temperature of 500 K and pressure of 1 atm. The cutoff of 12 Å was used in the calculation of nonbonded interactions. The switching distance was set to 14 Å. The time step of 12 fs was used for CG level simulations of polymer resin. The temperature and pressure were controlled during all simulations at NPT ensemble with Langevin thermostat and Noé-Hoover Langevin barostat. The system was run for total simulation time of 150 ns. MARTINI FF uses a cosine-based harmonic angle potential (Marrink et al., 2004). In NAMD-2.8, cosine-based angle potential can be employed by switching “cosangles” and “martiniswitching” functions “ON”.

The bond and angle interactions between CG beads are represented by means of simple harmonic potentials. In MARTINI force field, the chain stiffness is dictated with weak cosine type harmonic angle potential. For simplicity torsional angle potentials are ignored in this work. The parameters for bonded interactions are optimized using distribution of distances and angles from atomistic simulation trajectories. Following the MARTINI conventions, the bond and angle distributions are evaluated from trajectories of center of masses (COM) of groups of atoms in the all-atom model mapped onto CG beads (Lee et al., 2009). For harmonic potentials of CG simulations, the average distances and angles from COM trajectories are selected as equilibrium lengths and angles. The selection of force constants is based on reproduction of comparable widths of all distributions. The final set of bonded parameters for CG modeling of PMMA is reported in Table 2.3.

<table>
<thead>
<tr>
<th>Bond</th>
<th>$R_{\text{bond}}$ (nm)</th>
<th>$K_{\text{bond}}$ (kJ/mol)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A-A</td>
<td>0.282</td>
<td>14600</td>
</tr>
<tr>
<td>A-B</td>
<td>0.282</td>
<td>46000</td>
</tr>
<tr>
<td>Angle</td>
<td>$\theta_0$ (degree)</td>
<td>$K_{\angle}$ (kJ/mol)</td>
</tr>
<tr>
<td>A-A-B</td>
<td>71</td>
<td>85</td>
</tr>
<tr>
<td>NB pairs</td>
<td>$\sigma$ (nm)</td>
<td>$\varepsilon$ (kJ/mol)</td>
</tr>
<tr>
<td>A-A</td>
<td>0.425</td>
<td>3.56</td>
</tr>
<tr>
<td>A-B</td>
<td>0.473</td>
<td>2.7</td>
</tr>
<tr>
<td>B-B</td>
<td>0.473</td>
<td>3.7</td>
</tr>
</tbody>
</table>

At atomistic level, the trajectory from last 20 ns after equilibration was converted into COM trajectory to obtain all distributions of bonds and angles. All bond and angle parameters were optimized by comparing distributions of equilibrium bonds and angles extracted from
trajectories of 40 ns of CG simulation with the COM trajectories of atomistic simulation. The match among distribution plots of bonds and angles using optimized parameter sets are shown in Figure 2.9.

![Normalized probability distribution of bond length in (a) A-A and (b) A-B, and that of the angles in (c) A-A-A and (d) A-A-B.](image)

The distributions of bondlengths for bonds A-A and A-B obtained from atomistic and CG simulations match very well with each other. The plot for angle A-A-A also matches with atomistic COM trajectory. However, angle A-A-B does not seem to give comparable distribution for lower degrees. It is reasonable because of the larger size of bead type B than bead type A. The trajectory of COM of angle A-A-B is at highest probability for angles around
70°. In a CG model, values lower than 80° for angle A-A-B lead to bead B coming in close proximity to the backbone chain. Under such proximity, the beads encounter high steric hindrance from the backbone chain. Thus, it prevents the angle A-A-B to reach smaller values than 80°.

Parameterization of nonbonded interactions. The nonbonded interactions consist of contribution from Van der Waals forces and electrostatic forces between polymer beads. In the CG model of PMMA, both bead types are assigned with neutral charges. Thus, there are no electrostatic interactions among the polymer chains. According to the MARTINI protocol, the nonbonded interaction potentials among the neutral beads are represented by LJ function. The optimization of LJ potential parameters is based on the reproduction of selected target thermodynamic and structural properties. In this work, PMMA melt density and radius of gyration are set as target properties for optimization of nonbonded parameters. The CG model of PMMA considers two different types of beads with four-to-one and three-to-one mapping schemes. The size and mass for four-to-one mapped bead B is straightforward as 4.7 Å and 72 amu respectively. The size and mass of bead A with three-to-one mapping can be chosen as same or smaller depending upon the performance of the model in terms of target properties. It was found that the density obtained using smaller size of A was in better agreement (4.06% error) than the density using same size of A (12.3% error) with that of the density from atomistic simulation of PMMA melt.

In simulation, when the initial size (σ) and strength of interaction (ε) values from the original MARTINI force field were used for both the bead types, the density of the system was 1.161 g/cm³, which was significantly higher than that in the atomistic simulation (1.034 g/cm³), and the radius of gyration was also 18% larger. A larger radius of gyration indicated attractive
forces between backbone beads which were too weak. Thus, the force of attraction was increased by adjusting the value of $\varepsilon$ between A-A to 3.56 kJ/mol and B-B to 3.7 kJ/mol. Using these new values, the polymer melt density of the CG model at 500 K was found to be 1.076 g/cm$^3$ (with an error of 4.06%), and the radius of gyration of 1.236 nm was also in good agreement (with an error of 9.4% error) with that predicted by atomistic simulations. The performance of the CG model of PMMA was slightly better than the MARTINI model developed for polyethylene oxide (PEO) chains, where the densities of low molecular weight PEO are reproduced with an error of 6-8% (Lee et al., 2009). The optimized set of LJ parameters for all nonbonded pairs of beads are listed in Table 2.2, where $R_{\text{bond}}$ and $K_{\text{bond}}$ are the bond length and force constant at equilibrium, respectively, and $\theta_0$ and $K_{\text{angle}}$ are the angle and the force constants at equilibrium, respectively.

2.3.3 CG Model Verification

One of the important features of thermodynamics-based CG model for PMMA is its ability to retain molecular integrity at different temperatures. The developed CG model is verified through reproducing accurately the thermodynamic and structural features that are obtained from the atomistic simulation.

**Density.** The system characterized by the CG model consisted equal number of polymer chains as that in the atomistic model. The chains were allowed to equilibrate for 150 ns under melt condition at 500 K, and then cooled down at the interval of 100 K and equilibrated for 80 ns at each interval boundary under NPT ensemble. The densities were calculated by averaging over last 40 ns of trajectory. As shown in Table 2.4, the densities ($\rho$) evaluated by the CG model
and that through atomistic simulations have less than 5% of difference over the temperature range of 200 K to 500 K.

Structure. In general, CG models tend to lose structural features of a molecule at changing temperatures. Table 2.4 also shows that the radius of the gyration of PMMA chains ($R_{gp}$) seems to match well with that from atomistic simulation over the entire temperature range. A closer look at the data indicates that at lower temperatures, the density of polymer chains become larger, which leads to a smaller radius of gyration. Carbone et al. (2008) performed a comparative study on the transferability of the models that were obtained through coarse-graining of polystyrene (structure-based) and polyamide (thermodynamics-based). They demonstrated that the RDF’s of only polyamide model showed good comparison and thus the model was transferable at different temperatures. The CG model of PMMA could retain most of the structural features at all temperatures of simulation. As shown in Figure 2.10, the RDF curves of PMMA chains at the simulated temperatures are very close to each other. The first peak is increasing as the temperature is decreased, this is due to the stiffness of polymer chains at higher density.

<table>
<thead>
<tr>
<th>T (K)</th>
<th>$\rho$ (g/cm$^3$)</th>
<th>$R_{gp}$ (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>CG</td>
<td>Atomistic</td>
</tr>
<tr>
<td>500</td>
<td>1.076±0.012</td>
<td>1.034±0.006</td>
</tr>
<tr>
<td>400</td>
<td>1.121±0.015</td>
<td>1.107±0.009</td>
</tr>
<tr>
<td>300</td>
<td>1.162±0.009</td>
<td>1.154±0.005</td>
</tr>
<tr>
<td>200</td>
<td>1.198±0.007</td>
<td>1.191±0.008</td>
</tr>
</tbody>
</table>

Table 2.4. Comparison of the temperature-dependent density and radius of gyration of PMMA by the atomistic model and the CG model.
2.4 Modeling of Polymer Nanocomposite Coating

A nanocomposite coating consists of nanosize filler particles dispersed randomly in a coating matrix. There are several different ways to model nanoparticles inside a polymer matrix. Different types of nanoparticles (e.g. TiO$_2$ (Radhkrishnan et al., 2009), layered silicates (Nobel et al., 2007), MMT clay (Chow and Mohd Ishak, 2007), etc.) are generally added into polymer matrices to form nanocoatings with newer functionalities such as self-cleaning, high abrasion resistance, anti-fogging, super-scratch resistance, anti-corrosion resistance etc. Smart functionalities such as self-cleaning, abrasion resistant, pathogen killing coatings etc. can be obtained with several hydrophilic nanoparticles (Radhkrishnan et al., 2009; Menno et al., 2011). In order to mimic the behavior of these hydrophilic nanoparticles in nanocoatings, it is important to model nanoparticles with controlled hydrophilicity. According to the MARTINI FF, the sizes...
of beads which represent groups of atoms are predefined. These beads are assigned with different values of $\varepsilon$ of LJ interaction potential function. The values of $\varepsilon$ represent different levels of hydrophilicity/hydrophobicity. Marrink et al. (2007) described 9 different interaction levels with specific values of $\varepsilon$. Higher values of $\varepsilon$ can be used to model more attractive interactions among highly polar groups, whereas smaller values of $\varepsilon$ can be used to model various degrees of hydrophobic repulsions between polar and nonpolar phases.

Studies on atomistic level simulation have shown that C$_{60}$ fullerenes form spherical nanoparticles by closed packing large number individual entities to form spherical nanoparticles (Bedrov et al., 2005). At CG level, nanoparticles are modeled using clusters of small sized beads to form various shapes of nanoparticles (Knauert et al., 2007). We adopted a similar methodology to form spherical hydrophilic nanoparticles. The modeling of each nanoparticle inside the polymer matrix was performed using LJ beads of size 4.7 Å. A cluster of large number of beads connected to each other was created to form spherical nanoparticles. The number of beads needed to form these CG nanoparticle clusters were calculated using the volume of each individual bead and the total volume of each nanoparticle. The interactions between the beads were defined using only the LJ potential. For this work, the nanoparticle beads are named as “NP”. They are analogous to the bead type “P1” of MARTINI force field (hydrophilic beads with lower polarity). The strengths of interactions between nanoparticle beads and polymer beads are summarised in the table below.

<table>
<thead>
<tr>
<th>Bead types</th>
<th>$\sigma$ (Å)</th>
<th>$\varepsilon$ (kJ/mol)</th>
</tr>
</thead>
<tbody>
<tr>
<td>NP-NP</td>
<td>4.7</td>
<td>4.5</td>
</tr>
<tr>
<td>NP-A</td>
<td>4.7</td>
<td>2.7</td>
</tr>
<tr>
<td>NP-B</td>
<td>4.73</td>
<td>4.5</td>
</tr>
</tbody>
</table>
The simulation of nanocomposite coating was performed under NVT ensemble. These simulations were performed at longer time steps up to 25-30 ns to reach the equilibrium quicker. Under the melt condition at 500 K and atmospheric pressure, the polymer chains move freely and avoid getting trapped in one state of configuration. The sizes of nanoparticles and number of polymer beads decide the system size and thus the cutoff length for LJ potential. A long simulation time of 300 ns ensured reaching of the equilibration state and generation of sufficient trajectory length to extract data for measurement of structural properties. Generally, an average over a trajectory of last 50 ns is used for extracting data for analysis. The behavior of polymer beads was studied by keeping all the nanoparticles fixed at different locations inside the simulation box and the polymer was allowed to equilibrate.

The system was constructed using 15 chains of PMMA with 50 repeating units in each. The total number of polymer beads in the system were 1500. The size of the nanoparticle incorporated into the matrix was 3 nm. Note that the interaction among nanoparticle force sites is stronger than the interaction between polymer and nanoparticle beads. This stronger interaction prevents the polymer chains from entering inside the nanoparticle framework. It has been studied that the interface between the polymer and the nanoparticle plays a critical role in defining improvement in properties of nanocomposites (Starr et al., 2002; Jankar et al., 2010; Xiao et al., 2010). We studied the movement of polymer chains in the vicinity of nanoparticle surface. Figure 2.11 illustrates the behavior of polymer beads around nanoparticle surface at the end of equilibration process of the initially built system with one nanoparticle surrounding polymer chains. A “Visual Molecular Dynamics (VMD)” software package was used to visualize the 3D structure (Humphrey et al., 1996). The grey, green and blue colors represent
bead types NP, A and B respectively. As the simulation proceeds the polymer chains move towards the surface of nanoparticle and assemble into a structured pattern leading to a high density of beads at the interface. Figure 2.12 represents the number density distribution of polymer beads at different distances from nanoparticle center before (a) and after (b) the equilibration step. A peaks generated at the end of equilibration are due to the substantial increase in the number of polymer beads on the nanoparticle surface. It is the result of a formation of structured pattern of polymer beads at the interface. This phenomenon was also observed by Xiao et al. (2010).

Figure 2.11. Structures of nanocomposite: (a) the initial configuration, and (b) the final configuration after equilibration (where bead types NP, A and B are colored in grey, green and blue, respectively).
Figure 2.12. Number density distribution of polymer beads at different locations from the center of NP (3 nm size) (a) before and (b) after equilibration (cont’d).
2.5 CG Model Analysis and Evaluation of Properties

In order to be able to predict accurate properties and performance features of the coating system using this methodology, it is important to analyze the model for its accuracy in reproducing the data from atomistic simulations and also from experiments. The model being thermodynamically stable, it is also necessary to demonstrate its stability at different temperatures.

2.5.1 Radial Distribution Function

The analysis of the stability of the model and capacity to reproduce all the polymers structural features over a wide temperature range was demonstrated using radial distribution function $g(r)$ of polymer chains. When compared the curves representing the structure at different temperatures ranging from 200 K to 500 K, the CG model retained most of the structural features even at lower temperatures. The RDF plots of PMMA chains at different temperatures given in Figure 2.10 have all the peaks retained at same location. The height of the first peak is seen to be increasing with the temperature as a result of increasing stiffness of polymer chains at lower temperatures.

2.5.2 Microstructure Analysis

In this section we investigate various simulations performed on polymer-nanocomposite
samples by changing the structural parameters of both. We know that incorporation of nanoparticles in the polymer matrix causes polymer chains to move towards nanoparticle surface and orient in a highly dense pattern (Xiao et al., 2010). But there are several factors that resist the movement of bulk polymer and its microstructure on polymer-nanoparticle interface. By using MD simulation under NVT ensemble, we analyze the effect of nanoparticle size, polymer chainlength and polydispersity on the behavior of polymer beads in the vicinity of nanoparticle surface.

Effect of NP size. It has been proven that an interface between nanoparticle and polymer plays a key role in achieving superior properties of a nanocomposite coating. Xiao et al. (2010) demonstrated the formation of a structured pattern by polymer beads at the interface. The structured assembly of polymer beads causes an increase in the number density distribution. Consequently, it leads to improvement in the mechanical properties of the coating. The improvement in mechanical and rheological properties of nanocomposites is a function of surface area of nanoparticles exposed to polymer chains (Jancar et al., 2010). There are several different shapes of nanoparticles that are used to introduce different features into coatings. Nanoparticles could be rod like, spherical, sheet, platelet shaped, etc. Spherical TiO$_2$ and layered silicate nanoparticles of are amongst the most common ones in coating industries that give substantial improvement of mechanical, rheological or thermal properties and also sometimes brings in new functionalities like self-cleaning and anti-corrosion coatings (Zhang et al., 2005; Chow and Mohd Ishak, 2007; Chen et al., 2008; Radhakrishnan et al., 2009).

In order to model comparable size of nanoparticles, we first model spheres by close-packing multiple LJ beads into a spherical shape. As an initial case study, we model these type
of nanoparticles by assigning hydrophilic LJ parameters to individual beads which form spherical nanoparticles, as also explained in the “modeling of polymer nanocomposite” section.

Experimentally it has been determined that nanocoating samples with smaller sized NP show better performance of the final film than the ones with larger sized NP in it (Manoudis et al., 2007). In this work, the effect of size of spherical NP on the structured pattern formation of polymer around NP surface is investigated by simulating a single NP dispersed in PMMA. In each case, a total of 1,500 polymer beads is used. As shown in Figure 2.12, the highest density of polymer beads is obtained within 6 Å of the NP surface. Figure 2.13 shows that the density of polymer beads at the polymer-NP interface goes down as the size of the NP is increased.

![Graph showing polymer bead density at 6 Å distance away from NP surface for increasing NP size.](image)

Figure 2.13. Polymer beads density at 6 Å distance away from NP surface for increasing NP size.

The average radius of gyration of polymer chains is evaluated and used to analyze the structural properties. It is observed that incorporation of nanosize fillers can stretch polymer chains, thus increasing the radius of gyration. It is found that if the incorporated nanofiller is
larger in size, the average radius of gyration of polymer chains near the interface becomes larger. Figure 2.14 shows a linear increase of the ratio of radius of gyration in a nanocomposite relative to the pure polymer melt as the size of incorporated nanofiller is increased, which, indicates the stretching of polymer chains. This stretching creates more free volume at the interface, and thus reduces the polymer bead density.

![Graph](image_url)

**Figure 2.14.** The ratio of the radius of gyration of polymer in presence of NP to that in absence of NP as a function of the radius of NP.

**Effect of nanoparticle volume fraction.** In simulation, different concentrations of dispersed spherical NP are tested to evaluate the impact on nanocomposite performance. As shown in Table 2.6, the total surface area of NP is kept same (28.26 nm²) in all four cases. Then the average radius of gyration of polymer chains in each case is measured. The stretching of polymer chains due to interaction with NP’s gives rise to a large radius of gyration as compared
to a pure polymer case. Analysis of the radius of gyration suggests that the free volume available for polymer chains to stretch is less when the volume fraction of NP is higher.

Table 2.6. Radius of gyration of polymer chains at different NP concentrations.

<table>
<thead>
<tr>
<th>No. of NP’s</th>
<th>NP radius (nm)</th>
<th>Total NP surface area (nm$^2$)</th>
<th>$R_g$ (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1.5</td>
<td>28.26</td>
<td>1.65±0.15</td>
</tr>
<tr>
<td>2</td>
<td>1.06</td>
<td></td>
<td>1.612±0.13</td>
</tr>
<tr>
<td>3</td>
<td>0.866</td>
<td></td>
<td>1.633±0.16</td>
</tr>
<tr>
<td>4</td>
<td>0.75</td>
<td></td>
<td>1.59±0.19</td>
</tr>
</tbody>
</table>

Effect of polydispersity. It is a known fact that polydispersity of the polymer matrix under consideration plays a critical role in defining various performance properties of a nanocomposite. Gilman et al. (2000) studied the flame retardancy of a polymer nanocomposite having MMT clay platelets dispersed around polystyrene resin. It was determined that the organically modified PS-MMT grade does not show improvement in flame retardancy over immiscible PS-NaMMT sample. The organic treatment degrades the polymer thus increasing its PDI. According to Robello et al. (2004), the PS-clay nanocomposite grade with lower PDI of polystyrene shows an exfoliation of nanoparticles with very less agglomeration, unlike in the sample with higher PDI of PS. In this section we study the effect of polydispersity on the assembly of polymer beads around nanoparticle surface.

We carried out simulations of a nanocoating sample with varying number of polymer chains of different chain lengths. For the first case study, the chain length was varied from $n=40$ to $n=70$. The total number of polymer beads in the system were kept the same for each case. The size of the nanoparticle was fixed to 3 nm. The simulations were run at NPT ensemble for 300 ns with 30 fs time step. At the end of equilibration the total number of polymer beads at 6 Å
distance from nanoparticle surface were measured. The density of polymer beads give a measure of degree of assembly of polymer chains on the nanoparticle surface. Table 2.7 summarises the number of polymer beads gathering on the nanoparticle surface and average radius of gyration of polymer chains for all the samples. The density of beads increases consistently with the chain length for monodisperse samples. For bidisperse samples, the sample with more number of longer chains gives higher density. Although the samples of monodisperse and bidisperse polymer melts show higher polymer bead density for longer chains, these results are difficult to relate with tridisperse and tetradisperse samples. The loss of consistancy makes it difficult to correlate polydispersity with number density distribution of polymer beads. The calculation of average radius of gyration of polymer chains show reasonable trends with increase in polydispersity. The radius of gyration is higher for longer chains in monodisperse samples. Note that the radius of gyration of polymer chains is larger than pure polymer case due to the interaction with attractive nanoparticle surface. When compared with polydisperse samples, the radius of gyration keeps increasing with increase in polydispersity. This could be an effect of available free volume. At higher polydispersity, polymer chains have more volume available to stretch themselves. Shorter chains usually find more space and are more mobile compared to longer chains. The increase in the number of shorter chains assist in providing larger free volume for polymer chains that leads to increase in the radius of gyration.

Table 2.7. Density of polymer beads within 6 Å from the NP surface, and radius of gyration for monodisperse, bidisperse, tridisperse and tetradisperse polymer-nanocomposite samples.

<table>
<thead>
<tr>
<th>Polydispersity</th>
<th>Number of polymer chains</th>
<th>Total no. of beads</th>
<th>Number Density of beads in 6Å interface (no./ Å³)</th>
<th>Rg (Å)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Monodisperse</td>
<td>17 0 0 0</td>
<td>2380</td>
<td>0.0236</td>
<td>1.875±0.13</td>
</tr>
<tr>
<td></td>
<td>0 20 0 0</td>
<td>2400</td>
<td>0.0211</td>
<td>1.786±0.14</td>
</tr>
<tr>
<td></td>
<td>0 0 24 0</td>
<td>2400</td>
<td>0.0192</td>
<td>1.634±0.14</td>
</tr>
<tr>
<td></td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>39</td>
</tr>
<tr>
<td>----------</td>
<td>----</td>
<td>----</td>
<td>----</td>
<td>----</td>
</tr>
<tr>
<td>Bidisperse</td>
<td>0</td>
<td>15</td>
<td>6</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>0</td>
<td>10</td>
<td>12</td>
<td>0</td>
</tr>
<tr>
<td>Tridisperse</td>
<td>0</td>
<td>8</td>
<td>9</td>
<td>7</td>
</tr>
<tr>
<td>Tetradisperse</td>
<td>5</td>
<td>5</td>
<td>5</td>
<td>8</td>
</tr>
</tbody>
</table>

### 2.6 Summary

This work provided a detailed analysis of structural and thermodynamic behavior of polymer chains at the interface of polymer and NP. The modeling and simulation of PMMA chains surrounding hydrophilic NP cluster was performed using the MARTINI based coarse-graining methodology. Model verification was provided by reproduction of structural and thermodynamic properties of the polymer obtained through atomistic simulation that was performed using an optimized set of CHARMM force field parameters. The predicted densities, radii of gyration and distributions of bonds and angles were in good agreement with that from all-atom simulations over the temperature range of 200 K to 500 K.

The developed model of polymer-nanocomposite system was applied for the study of structural behavior of polymer chains around induced nanofillers with varying NP size, volume fraction and the polydispersity of the polymer PMMA. It is found that the radius of NP’s dispersed in nanocoating resin changes polymer bead density and the radius of gyration of polymer chains. For samples with large NP’s, the interfacial polymer bead density is smaller and the radius of gyration is larger than for samples with small NP’s as a result of availability of more free volume. It was also shown that increasing the polydispersity index of polymer matrix elongates polymer chains. These results favor a selection of polymer matrix with low polydispersity index with more number of longer chains and a higher volume fraction of NP’s with smaller size to obtain higher interfacial density of polymer beads.
The developed CG parameter set for PMMA and the simulation methodology for corresponding nanocomposite modeling should be promising as they can be valuable in the study of water-dispersible acrylic nanocoating systems. Additional nanocoating parameters, such as self-cleaning, and corrosion resistance, can also be investigated through the developed CG system.
CHAPTER 3
EXPERIMENTAL STUDY ON POLYMER NANOCOMPOSITE COATINGS

From past two decades, polymer nanocomposite coatings are being studied through various experiments to explore their potential to bring new and significantly improved functionalities into coating materials. This area of research is still ever increasing through millions of dollars being invested on it, as innumerable questions related to structure and material-property correlations of polymer nanocomposites are still unanswered.

The real challenge to this research comes from the limitation of scientific knowledge of nanocoating microstructure and required methodological-technical development. Needless to say, that the sustainable nanopaint development is achievable only with an integrated computational-experimental effort. Such study should be effective in analyzing all the aspects of the overall life cycle of nanocoating technology, which ranges from nanoparticles synthesis, through coating formulation, processing, and application to final life-cycle performance. The experimental analysis of various nanocoating materials and their characterization can not only provide the verification to the property predictions obtained from computational modeling, but also it can provide the key insights into the effects of various materials on microstructure-property correlations.
3.1 Objectives and Task Definition

The objective of the research is to develop a comprehensive multiscale methodology to assist in developing new nanocoating materials with multifunctional performance characteristics and improved sustainability than conventional coating materials. The outcome from computational findings would generate critical information about the microstructural behavior and the correlation between material and the bulk mechanical properties of nanocomposite resin. The computational study would also provide a variety of information for designing the experiments, which include the recommendations for selecting polymer matrix and nanoparticle type, size and shape. These findings would be useful in optimizing formulation parameters for the development of multifunctional nanocoating materials through experiments. In order to develop the correlation between paint material and bulk properties, computational work needs to be integrated with experiments. The verification of the modeling methodology through the case studies of real polymer nanocomposite coatings can support computational findings significantly.

In this work, the experimental efforts focus towards multiple aspects, such as providing verification to computational modeling and studying the bulk properties of nanosize polymer nanocomposite films. The steps followed to achieve these objectives are as follows:

(i) Polymer nanocomposite coating synthesis. TiO$_2$ nanoparticles are photocatalytic in nature. A nanocoating formulation using selected polymer resin and TiO$_2$ nanoparticles would be synthesized. The nanoparticles would be surface treated to aid the formation of stable dispersion without aggregation into polymer matrix. The selection of polymer resin type, solvents and size and shape of nanoparticles decide the performance of the final coating film.
(ii) **Analysis of nanocoating formulation.** The formulated nanocoating resin will be analyzed for nanoparticle aggregation or uniform dispersion using SEM (scanning electron microscopy), AFM (atomic force microscopy) or other high resolution electron microscopy.

(iii) **Application of coating material on panel surface and sample development.** Nanosize coating film would be generated using sophisticated spin coating applicator.

(iv) **Coating films’ analysis for multifunctional performance.** The cured nanocomposite films would be subjected to mechanical testing with nano-scratch indentation for the film stiffness analysis. Tapping mode AFM imaging would be employed to determine the nano-scratch depth and paint film morphologies. Self-cleaning functionality would be quantified through measurement of paint film surface properties. The measurement of contact angle change would give a degree of super-hydrophilicity achieved with the TiO$_2$ nanoparticles addition.

(v) **Verification of computational models.** The formulation recipes and coating films would be analyzed thoroughly to study the effects of nanosize film thickness and morphology on the mechanical properties. The film stiffness and mechanical properties will be compared with the computational modeling results and the prediction of same properties through molecular simulation will be verified.

**Project definition.** Nanoparticle induced coating compositions have significant number of benefits over their conventional counterparts. However, there are several factors related to the formulation, structure and application of nanocoatings which may affect their improved performance. These factors include types of polymer resin and nanoparticles, size, shape and chemistry of nanoparticles, distribution of nanoparticles within cured coating films, nanoparticle
aggregation, adhesion of the film to the substrates and with other coating films in multi-coat applications, rheological properties of nanocoating resins, application techniques, etc.

A thorough research is in progress to correlate nanoparticle morphological parameters with enhanced performance properties of the final polymer nanocomposite. In case of coatings, it is believed that the distribution of nanoparticles inside the coating film is a crucial factor which dictates the mechanical and barrier properties of the final film. It has been shown that the superhydrophilicity or self-cleaning feature of coating films can be controlled by applying layer-by-layer films of TiO$_2$ and SiO$_2$ nanoparticle dispersions of known thicknesses (Wu et al., 2007; Permpoon et al., 2008). This type of assembly of nanoparticle layers within a coating film can also change the mechanical properties of the film. A very little attention has been given towards this correlation between the mechanical properties and nanoparticles distribution inside the cured coating film. Our objective of the project is to analyze the change in mechanical performance as a function of location of nanoparticles layer inside the coating film. It is accomplished by sandwiching TiO$_2$ nanoparticles layer within two PMMA thin films, as shown in Figure 3.1.

![Figure 3.1. Coating film structure after sandwiching nanoparticle film between two polymer thin films.](image)

The current nanocoating films have random distribution of nanoparticles inside the coating film. We believe that if the distribution is controlled and the nanoparticles are concentrated at a location which would give maximum enhancement of the final properties, it
can lead to nanocomposite coating formulations with significantly improved performance. If this phenomenon is studied thoroughly and the research predictions are verified, the findings can also provide directions to the discovery of next generation nanocoatings which would require a single layer of paint to perform all the functions delivered by current technique of multiple layers of coatings.

### 3.2 Materials Selection and Sample Preparation

The polymeric material selected for this study was polymethyl methacrylate (PMMA) with average molecular weight $M_w = 120,000$ and glass transition temperature of $105^\circ$, purchased from Sigma-Aldrich. The nanoparticles were anatase-$\text{TiO}_2$ type with average particle size $< 25$nm, purchased from Sigma-Aldrich. The solvent chloroform was used to disperse PMMA. Silicon wafer substrates of 1 cm x 1 cm dimensions were used for application of nanosize films. The wafer surface was cleaned using methanol, acetone and water and by immersing the wafer substrates into freshly prepared pirhana solution for 30 minutes to remove all organic residues and make a clean hydrophilic surface. A typical pirhana solution is prepared by adding 30% hydrogen peroxide solution into concentrated sulfuric acid at a ratio of 1:3 (by volume). After removing from the pirhana, wafers were rinsed thoroughly with deionized water and dried using air jets.

The bare $\text{TiO}_2$ nanoparticles were tested for the solubility in organic solvents. The selection of appropriate solvent is critical to form a stable and uniform dispersion of nanoparticles which can be applied on the substrate. The solubility of the bare nanoparticles was found to be poor in most of the organic solvents. Nakayama and Hayashi (2008) suggested that
several efforts have been made to improve the solubility of nanoparticles by surface modifying these nanoparticles. However, these efforts produced nanoparticles with improved dispersibility in organic solvents but at a cost of high refractive index of TiO$_2$. The objective of their study was, (1) to prepare a good dispersion with stability of TiO$_2$ nanoparticles in organic solvents with surfactants of the minimal quantity, and (2) to disperse TiO$_2$ nanoparticles into the organic solvents, by exchanging the type of the organic surfactants, without changing the properties of TiO$_2$ nanoparticles as much as possible. TiO$_2$ has several Lewis acid sites (Ti$^+$) and Bronsted acid sites (Ti-OH) on the surface. If these sites are capped using both carboxylic acid and amine, the nanoparticles can form a stable dispersion in organic solvents. The scheme of surface binding at the TiO$_2$ surface is given in Figure 3.2 (Nakayama and Hayashi, 2008).

![Figure 3.2 Scheme of surface binding of TiO$_2$ nanoparticles surface.](image)

The procedure followed for the surface modification of these nanoparticles is as follows (Nakayama and Hayashi, 2008).
1. Bare TiO$_2$ nanopowder was added to excess of propionic acid and stirred for 24 hours.

2. The wet cake was obtained by centrifugation at 5000 rpm.

3. It was washed with ethyl acetate 5 times to remove excess of acid and added into methanol.

4. The solution of acid modified particles in methanol is mixed with excess amount of n-hexylamine and stirred for 1 hour.

5. The wet cake obtained by centrifugation at 5000 rpm was washed with ethyl acetate to obtain the surface modified TiO$_2$ nanopowder.

6. An overnight desiccation drying was done to obtain a dried nanopowder.

The FTIR spectrum of TiO$_2$ nanoparticles was then obtained to ensure the surface modification. The graph showed absorbance peaks at wavenumbers representing acid and amine functional groups (See Figure 3.3). These peaks ensured the successful surface modification of the TiO$_2$ surface. The solubility of the final surface modified TiO$_2$ was also tested in various organic solvents. The findings are given in Table 3.1.

<table>
<thead>
<tr>
<th>Solvent</th>
<th>IPA</th>
<th>Chloroform</th>
<th>Methanol</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bare TiO$_2$ Solubility</td>
<td>No</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>Surface modified TiO$_2$ Solubility</td>
<td>No</td>
<td>Yes</td>
<td>Yes</td>
</tr>
</tbody>
</table>
A polymer and nanoparticle dispersion films were applied by spin coating technique. The solutions of PMMA/chloroform were prepared at different concentrations of 0.5, 0.75, 1, 1.25, 2 and 2.5 weight %. All the samples were spun for 60 seconds at rotational speed of 3000 rpm. In order to reduce the surface roughness and to ensure removal of the solvent completely, the PMMA sample films were kept in an oven at 110º for 30 minutes. TiO₂ nanoparticles dispersion was prepared by dispersing the surface modified nanoparticles into solvent methanol at a concentration of 0.1 % (by weight). The dispersion was then applied onto the wafer substrate to form nanoparticle films by spin coating for 60 seconds at 3000 rpm.

For the analysis of elastic moduli of polymer nanocomposite films, three different samples were prepared using films of PMMA and nanoparticle dispersions as below:
(1) PMMA layer of thickness 48 nm using the solution with concentration of 0.75 % (by weight).
(2) PMMA layer of thickness 48 nm + TiO$_2$ nanoparticles layer using the dispersion of nanoparticles in methanol
(3) PMMA layer of thickness 48 nm + TiO$_2$ nanoparticles layer + PMMA layer of thickness 20 nm.

3.3 Film Thickness Analysis

The measurement of polymer film thicknesses was done using X-ray reflectivity (XRR) technique. Previous work by Walsh and Franses (2003) developed a correlation between PMMA film thicknesses and PMMA/chloroform solution concentrations at given conditions of spin coating (see Eq. 3.1).

$$d_1 = 4.30c^{0.33}\omega^{0.5}$$

where $d_1$ is the film thickness in mm, $c$ is the concentration of the solution in weight percent and $\omega$ is the rotation speed in rpm.

XRR measurement. In the past decade, X-ray reflectivity (XRR) technique has become an important tool to study the interfacial properties of thin films (Rao et al., 2004). XRR is a non-destructive, routine technique which is used for estimation of density, thickness and roughness of thin film structures. In general, it can be used with crystalline, amorphous and liquid samples. For the film thickness measurement XRR uses electron density fluctuations at the interface of two surfaces. The measurement of thickness is based on the total external X-ray reflection from the surface. This technique is especially suitable to study buried film surfaces as
X-rays can penetrate deep inside the coating material. This technique is used for layer thicknesses between 5Å to 400 nm. In this technique, the diffraction peaks are correlated with the thickness of polymer film.

For PMMA films applied using polymer dispersions of different concentrations, the samples were kept inside the XRR instrument and reflectivity data was recorded. The recorded data curve was analyzed and initial film thickness was predicted using FFT (fast Fourier transform).

The reflectivity of a single layer deposited on the substrate is expressed using an equation given as:

$$ R = \left| \frac{r_1 + r_2 e^{-2k_0 z t}}{1 + r_1 r_2 e^{-2k_0 z t}} \right|^2 $$

(3.2)

where $r_{1,2}$ are the Fresnel reflectivity coefficients of the free surface and the substrate interface respectively, $k_{0z}$ is the vertical component of the wave vector of the X-ray beam transmitted through the film and $t$ is the film thickness. The intensity maxima occurs at all points where $e^{-2k_{0z} t} = 1$.

During the determination of film thickness using XRR method, the roughness of the surface plays a key role. In reality, thin film surfaces are not perfectly smooth. The presence of high surface roughness decreases the specular intensity of the curve and gives rise to progressive damping of Kiessig fringes. Figure 3.4a,b represents the reflectivity data curves for polymer films applied onto a silicon wafer substrate using 1% and 2.5% (by weight) solutions of PMMA respectively. It can be observed that the fringes disappeared for the film of 2.5 % solution due to higher roughness of the surface. The roughness values are measured using tapping mode AFM and the values are reported in Table 3.2.
Table 3.2. Film thickness values for PMMA solutions at different concentrations

<table>
<thead>
<tr>
<th>Concentration (%)</th>
<th>Roughness (nm)</th>
<th>Measured thickness (nm)</th>
<th>Thickness from Eq. 3.5 (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.5</td>
<td>1.1</td>
<td>25.58</td>
<td>31.22</td>
</tr>
<tr>
<td>0.75</td>
<td>1.4</td>
<td>48.22</td>
<td>53.54</td>
</tr>
<tr>
<td>1</td>
<td>1.6</td>
<td>67.48</td>
<td>78.50</td>
</tr>
<tr>
<td>1.25</td>
<td>2.1</td>
<td>101.45</td>
<td>105.63</td>
</tr>
<tr>
<td>2</td>
<td>3.5</td>
<td>304.37</td>
<td>197.36</td>
</tr>
<tr>
<td>2.5</td>
<td>11.6</td>
<td>350.10</td>
<td>265.56</td>
</tr>
</tbody>
</table>

Figure 3.4. XRR plots for polymer film coated with (a) 1% solution of PMMA and (b) 2.5% solution of PMMA (cont’d).
The film thickness values are plotted against the concentration of the solutions of PMMA (see Figure 3.5). The values of the film thickness seem to match well with an error of less than 10% for low concentrations of the solution. At high concentration, the measurement technique does not produce reliable results because of increase in the roughness of the film surface.
3.4 Elastic Modulus Measurement

The elastic properties of the polymer film can be quantified using Young’s modulus measurement. Atomic force microscopy (AFM) is widely used to study the material properties and film morphology at nano-scale. The mechanical and topological properties of the polymer film can be calculated from force-distance curves which are obtained from AFM.

Figure 3.6a represents the operation scheme of AFM imaging by different modes. The AFM tip is generally able to probe into a very tiny area on the surface due to its extremely small radius around 5-50 nm. The tip is suspended with a very small spring through which all the highly sensitive forces of interaction from the tip are accessible for measurement. These cantilever beams are usually 100-200 microns long and can measure forces in the range of pico-Newton.
The AFM is used in the force calibration mode for the measurement of force applied by the cantilever on the tip. In this mode, the tip is brought together with the surface of the polymer film in a fluid cell at room temperature. The probe radius is 10 nm and the force constant (k) is 0.06 N/m. The force curves are obtained by bringing the tip in close contact with the surface in an aqueous medium and recording the force v/s deflection measurements after allowing the system to equilibrate for 10 min, until reproducible curves are observed. The data of deflection as a function of piezo position in the z-direction is obtained at different locations on the surface.

The data is analyzed at three different regions of the deflection curve. In the region with no contact between tip and the surface, the cantilever deflection “d” does not change with respect

Figure 3.6. Representation of (a) the outline of AFM imaging and (b) force-distance curve obtained from force calibration mode of AFM.
to the position of piezo in the $z$-direction. After the contact, the deflection begins to increase which represents the nonlinear region of the curve.

After the contact the tip indents inside the sample film to a distance $\delta$ resulting in small deflection $d = z - \delta$. After certain time, the tip reaches the infinite stiffness point where the deflection curve becomes linear with respect to piezo position and $d = z$. In generating the graph for force v/s displacement, the point at which the nonlinear region begins is considered as the starting point, $z_0$. The data of deflection against $z$-piezo position is converted into force against indentation by using Hooke’s law, where the force $F$ is proportional to the deflection through cantilever force constant, $k$.

$$F = kd = k(z - \delta)$$  \hspace{1cm} (3.3)

The final force against indentation curve is obtained by (a) calibrating the sensitivity, (b) shifting the deflection curve to (0,0) mark, (c) converting the $z$-piezo movement to indentation values, and (d) calculating $F$ values using deflection data through Hooke’s law.

The elastic deformation of the polymer film is represented by Hertzian mechanics (Mermut et al., 2003). The Young’s modulus then can be calculated using the equation,

$$F = \frac{4E\sqrt{\pi}}{3(1-\sigma^2)} \frac{3}{\delta^3}$$  \hspace{1cm} (3.4)

The Poisson’s ratio was selected as 0.35 for polymer PMMA film. The radius of the tip was 10 nm. From the linear plot of log $F$ as a function of log $\delta$, the intercept value was used to calculate the Young’s modulus by using the equation below.

$$\log F = \frac{3}{2} \log \delta + \log \left( \frac{4E\sqrt{\pi}}{3(1-\sigma^2)} \right)$$  \hspace{1cm} (3.5)

Three different samples were prepared for the analysis of elastic properties of the polymer and nanoparticle layers. The Young’s moduli were calculated for each of the samples to
study the effect of addition of nanoparticles into coating film. The plots of force to indentation for all the three samples are given in the following figure.

![Force vs indentation curves for (a) Sample 1, (b) Sample 2 and (c) Sample 3.](image)

The surface morphology was studied using AFM imaging under tapping mode. Figure 3.8 shows the AFM images for Sample 3 after coating of each layer followed by curing. It can be observed that a substantial number of nanoparticles are on the surface after application of the 2nd layer of nanoparticles. This layer disappears after applying the 3rd coat of PMMA of 20 nm thickness. A minimal amount of nanoparticles which are visible in the wet film also disappear after the sample is kept under heating for 30 min. After heating, all the solvent from the layer evaporates and the smoothness of the film improves.
Figure 3.8. AFM images of surfaces for sample 3 after application of nanoparticle layer (left), after application of top PMMA layer (middle) and heating of the top layer (right).

The graphs of force as a function of indentation for all three samples were converted into linear plots by taking a log on each side as shown in Eq. 3.5 and the elastic moduli were determined. The results are tabulated in Table 3.3.

Table 3.3. The elastic moduli of all three samples of polymer nanocomposites.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Description</th>
<th>Elastic Modulus (GPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>PMMA (48nm)</td>
<td>2.25 ± 1.1</td>
</tr>
<tr>
<td>2</td>
<td>PMMA (48nm)+TiO$_2$</td>
<td>94.15 ± 13.2</td>
</tr>
<tr>
<td>3</td>
<td>PMMA (48nm)+TiO$_2$+PMMA (20nm)</td>
<td>20.98 ± 10.6</td>
</tr>
</tbody>
</table>

3.5 Summary

In this work, the study of effect of distribution of nanoparticles inside a coating layer on the film properties was performed. For this study, an acrylic resin (PMMA) which one of the most common paint resins for water-borne formulations was used along with hydrophilic anatase
TiO$_2$ nanoparticles. The selection of water miscible ingredients was done by keeping in mind the emerging need for low VOC paint formulations.

The polymer and nanocomposite coating layers were analyzed and studied using AFM technique. The films were applied by spin coating method. The analysis of film thicknesses was carried out using a sophisticated XRR measurement method. The film thicknesses in the range of nanosize were obtained using spin coating under controlled operating conditions. The reliability of the values depends on the roughness of the coating film. The AFM under tapping mode was employed to determine the roughness values of polymer and nanocomposite coating films. The force calibration mode of AFM was used to determine the elastic properties of the films. The calculated elastic modulus for PMMA layer was in good agreement with the empirical value. It confirmed the applicability of the Hertzian mechanics for the measurement of elastic moduli of polymer/nanocomposite films. The samples were prepared by sandwiching a nanoparticles layer in between two PMMA coating layers. The coating layer thicknesses were controlled such that the final film thickness would remain within the range of 100-200 nm. A contact mode AFM was applied to perform the mechanical stiffness analysis of each of the coating layers. From the force to indentation curves, it was clearly seen that the elastic modulus of the coating film increased significantly after the addition of nanoparticles layer. The results were in good agreement with the prediction from the computational model which stated that the evenly distributed nanoparticles inside a coating layer improve the mechanical strength of the final film.

One of the limitations of this work was the accuracy of the Young’s moduli values. The values may not be reliable because of insufficient indentation of the AFM tip inside the coating film. The reliability of the results can be improved if the indentation at least reaches the bottom
layer (more than 50 nm). It can be achieved with a very strong material, such as diamond, for the AFM tip.

The future work should focus towards the measurement of accurate Young’s moduli of polymer nanocomposite films using hard diamond tips and a stronger cantilever beam of AFM. The values then can be compared directly with the elastic properties prediction obtained from computational modeling of the same material. The generation of more nanocomposite samples by varying the location of nanoparticles film from bottom of the layer to top, followed by determination of change in mechanical stiffness can generate critical information regarding the significance of nanoparticles distribution inside a coating layer. If this distribution can be controlled and optimized for maximum enhancement of the film properties, it can lead to breakthrough coating technology and a possible single coat application to serve a purpose of multiple coats (e.g. primer, basecoat, topcoat).
CHAPTER 4
MULTISCALE LIFE-CYCLE BASED SUSTAINABILITY ASSESSMENT (LCSA) OF NANOCOATING TECHNOLOGY

The concept of sustainability deals with the balance in the “triple bottom line”, which is to accomplish economic development, environmental cleanliness and societal satisfaction together. Collaborating sustainability approach with life cycle (LC) based study can support the economic growth of nanotechnology while improving the quality of life and without impairing public health or detrimental harm to the environment.

4.1 Objective and Significance

The nanocomposite coating material is synthesized by dispersing nanoparticles of different sizes and chemical nature into polymer resin. The application process of these coatings consists of spraying the paint on substrates and heating the substrate to allow the film to dry. The final paint film has numerous applications due to superior performance properties provided by incorporated nanoparticles. The life cycle of nanoparticle coatings can be divided into various stages. We define six stages of complete life cycle of nanocoating material, called as ‘cradle-to-grave’ continuum: (1) Raw material selection and preprocessing, (2) Paint Manufacturing, (3) Paint spray/flash, (4) Curing, (5) End use, (6) Disposal. First four stages account for nanopaint film development, and remaining two stages account for its use and
disposal. Figure 4.1 represents the pathway that connects all the stage of life cycle of a nanocoating.

![Life cycle of a nanopaint](image)

Figure 4.1. Life cycle of a nanopaint.

During these several stages, the nanoparticles are very prone release into the surrounding air and waste water. Assessing the release of nanoparticles and the possible exposure to environment and mankind is very crucial while framing critical stages of life cycle. These nanocoatings can be most susceptible to environmental air, water and soil at different stages of life cycle. TiO$_2$ and SiO$_2$ are most common types of nanoparticles that are added into coating formulations. Their use in nanocoating systems has been ever increasing due to all the benefits achievable with them.

The objective of this research is to identify and optimize ways to develop a sustainable nanocoating system with minimal environmental implications and improved societal safety and health care while preserving all the economic benefits of this novel technology. To meet this goal, all the key parameters related to the paint material and properties at each stage of the life cycle are identified and addressed. The correlations among sets of measurable parameters at each stage are established. The development of a sustainable nanocoating technology requires a comprehensive assessment through various appropriate sustainability indicators. The parameter sets at various stages will guide the development of sustainability matrix selection. The indicators are able to address the issues related to triple bottom line. The correlations among
indicator sets at different stages of life cycle are established. The overall life-cycle based sustainability assessment from all the stages should guide the development of nanocoating technology towards more sustainable future products and also highlight the areas of improvement at each stage of the life-cycle. Figure 4.2 represents a complete framework of the missions to be accomplished by this study. The developed assessment methodology is exemplified using case studies of conventional and nanotechnology based coating materials. The conclusions are drafted and recommendations are made to support the development of sustainable nanocoating material.

Figure 4.2. Life-cycle-based sustainability assessment structure.
4.2 Life Cycle Aspects

The term life-cycle assessment was originally coined and documented in early 1990 in ISA 14040/14044. This method comprises of four key phases. These key phases and their correlation is shown in Figure 4.3.

![Figure 4.3. LCA framework as defined in ISO 14040/14044.](image)

Life Cycle Assessment is a method for estimating and assessing the resource usage and environmental impacts attributable to the entire life cycle of a product, from raw material extraction and acquisition, through energy and material production and manufacturing, to use and end-of-life treatment and final disposal (ISO 14040, 2006). The impacts consist of ozone layer depletion, climate change, increase in toxicity of human health and ecosystem, water usage, etc. The details on the requirements and guidelines for the four phases shown in Figure 4.3 can be found in ISO 14044 (2006). The goal and scope definition phase provides the details of
product system boundaries and functional units. The inventory analysis phase is focused on the data collection and calculation procedures required to quantify all the measurable entities defined in phase 1. The life cycle impact assessment aim at aggregating the results from the inventory analysis and determine product’s potential for impact on environment and human health. This process involves connectivity between inventory data analysis phase with corresponding impact category indicators. The purpose of interpretation phase is to consider all the findings from inventory data and impact assessment and to provide appropriate conclusions and recommendations.

In this work, the LCA methodology described above is combined with sustainability assessment procedure. A comprehensive sustainability assessment of nanocoating technology considers following phases of the life cycle: (1) paint material selection and preprocessing, (2) paint manufacturing, (3) paint spray/flash application, (4) curing, (5) end use, and (6) disposal.

The examination of the stages for selection of paint material, manufacturing, processing and use and its final disposal stages requires careful assessment of energy, resources and impact on environment through different design and technology alternatives. Although most of the data required for the assessment is available only from the research papers and industrial practices, few parameters are very difficult to quantify due to unavailability of appropriate data. Thus, it is critical to highlight the parameters that can be measured and can be used for the assessment. These parameters are associated with different time and length scales of measurement. Figures 4.3a through 4.3f shows the framework of parameters selection and their correlations associated with nanocoating technology at different life-cycle stages. Note that the last stage of disposal has been ignored as it is least important and would make only a minor impact on the overall sustainability of the system.
For Stage 1 (material selection and preprocessing), seven parameters were identified which can be quantified either using the material data from industry or the data research papers. These parameters are as follows:

S1: Paint resin properties (e.g. molecular weight, PDI, functionality, etc.)
S2: Types of additives selected for the paint formulation
S3: Nanoparticles size, shape and orientation
S4: Particle size distribution for pigment
S5: VOC content
S6: Toxicity of each of the formulation ingredient
S7: Raw materials cost

The selection of these parameters was based on the availability of the data for analysis. More parameters related to energy, water and electricity consumption during each of the raw material extraction and synthesis could be added. Such parameters were neglected at this stage due to the limitation of the data.

Stage 2 (paint manufacturing) was assessed using nine different parameter sets which are given below. The selection of parameters was done such that all the key aspects related to environmental emissions, social satisfaction and cost for manufacturing and energy consumption would be considered.

M1: Paint system composition
M2: Concentration of nanoparticles released/exposed to the surrounding during manufacturing of the paint
M3: VOC emission during manufacturing
M4: Amount of solvent and water consumed
M5: CO₂ emission and water usage
M6: Quantity of wastes generated
M7: Toxicity index for all the raw materials and ingredients used during the manufacturing
M8: Amount of energy consumed for all the processes and electricity usage
M9: Recycle factor. It is a measure of quantity of raw materials which can be recycled

The data for these parameters can be easily obtained from a paint manufacturing industry. The analysis using such data can be very effective in improving the sustainability performance of the paint manufacturing stage.

For the assessment of Stage 3, the selected parameters can be quantified using the data from not only paint application industries but also computational modeling. These are as follows:

A1: Film surface topology parameters
A2: Paint film thickness data
A3: Emission of VOC’s and nanoparticles during paint application through spray technique
A4: Wet film defects (e.g. runs, sags, blisters, etc.)
A5: Paint transfer efficiency
A6: Ventilation system performance (e.g. air velocity, energy consumption)
A7: Energy efficiency of the paint-spray system

The data for some of these parameters is very difficult to obtain from the industrial practices and empirical measurements. Thus, computational modeling and simulation tools can
be employed to determine the performance of some of these parameters, especially in case of nanopaint application.

The paint spray and curing are one of the most energy consuming stages of the life cycle of coating technology. The quality of the final coating film is also dependent on these stages and optimization of corresponding operation settings. Thus, it is essential to perform a careful analysis of these stages in terms of energy, environment and quality. For the paint curing stage (Stage 4) eight measurable parameters were identified which are enlisted below:

- **C1**: Coating microstructure analysis
- **C2**: Surface roughness after curing
- **C3**: VOC and nanoparticles emission
- **C4**: Film thickness and uniformity (DFT measurement)
- **C5**: Inter-coat adhesion study (film quality parameter)
- **C6**: Over operation settings and parameters
- **C7**: Crosslinking reaction conversion
- **C8**: Net energy consumption by ovens

The 5th stage (Use/Reuse) is very crucial for social parameters and coating quality parameters. The performance at this stage is essentially decided by the sustainability performance of previous stages. The assessment of Stage 5 includes majority of the parameters related coating quality, performance and toxicity issues. Again, these sets of parameters can be modified depending upon the type of paint system to be studied. These parameters are enlisted below.

- **E1**: Gloss retention
- **E2**: Coating film functionalities
E3: Cost of the final product
E4: Toxicity and possible hazards on contact with the film
E5: Solvent and water resistance of the film
E6: Scratch resistance and mechanical stiffness
E7: Corrosion inhibition

All the parameter sets enlisted above are general and can be applied for most of the nanocoating systems. However, these parameter sets can be modified with new type of parameters if the data is available for a specific type of nanopaint to be studied. The correlations among these parameter sets over different time and length scales are shown below in Figure 4.4.
Figure 4.4. Life cycle of a paint material with measurable parameters and their correlations at: (a) all stages of life-cycle, (b) paint material selection stage (cont’d).
Figure 4.4. Life cycle of a paint material with measurable parameters and their correlations at (c) paint spray/flash and (d) coating curing stages (cont’d).
Figure 4.4. Life cycle of a paint material with measurable parameters and their correlations at (e) coating curing and (f) end use stages.
The quantities of these parameters can be obtained from past research work, computational models or industrial data. Based on the type of paint material and the available data, the final choice of parameters can be made which would define the sustainability assessment framework for the nanocoating system to be examined.

4.3 Sustainability Assessment Framework

In simplest words, “sustainability”, as defined by the 1987 World Commission on Environment and Development (WCED) as, “Meeting the needs of the present without compromising the ability of future generations to meet their own needs (Mebratu, 1998).” The sustainability assessment of nanotechnology deals with multiple dimensions of a design development and cause-and-effect relationships over various stages of life cycle.

The sustainability assessment is performed using appropriately selected environmental, social and economic indicator sets. Each indicator is quantified using the reference data for multiple cases and either absolute or normalized values are obtained. During the life cycle of nanopaint, each stage is associated with different sets of measurable parameters. Thus, the sustainability matrix is developed consisting of different indicator sets at each stage of life cycle based on the set of measurable parameters from that stage (see Table 4.1).
Table 4.1. Sustainability matrix over a life cycle of nanopaint.

<table>
<thead>
<tr>
<th>Stages</th>
<th>Economic</th>
<th>Environmental</th>
<th>Social</th>
</tr>
</thead>
<tbody>
<tr>
<td>Paint Material selection</td>
<td>Composition of paint (SI-2)</td>
<td>Amt. of toxic raw material (SI-1)</td>
<td>Human health impact (SI-4)</td>
</tr>
<tr>
<td></td>
<td>Material Cost (SI-6)</td>
<td>Amt. of solvent/water (SI-3)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Environmental impact factor (SI-5)</td>
<td></td>
</tr>
<tr>
<td>Paint Manufacturing</td>
<td>Raw material composition (MI-1)</td>
<td>Release/exposure of hazardous material (MI-2)</td>
<td>Waste generated (MI-4)</td>
</tr>
<tr>
<td></td>
<td>Recycle efficiency (MI-6)</td>
<td>Net solvent/water consumption (MI-3)</td>
<td>Toxicity assessment (MI-5)</td>
</tr>
<tr>
<td></td>
<td>Net manuf. Cost and capital investment (MI-7)</td>
<td>Total energy consumption (MI-8)</td>
<td></td>
</tr>
<tr>
<td>Paint spray/flash</td>
<td>Material transfer efficiency (AI-2)</td>
<td>Net VOC/NP emission (AI-1)</td>
<td>Film quality assessment (AI-3)</td>
</tr>
<tr>
<td></td>
<td>Net operational cost (AI-6)</td>
<td>Net energy usage (AI-5)</td>
<td>Human health burden (AI-4)</td>
</tr>
<tr>
<td>Curing</td>
<td>Energy efficiency (CI-5)</td>
<td>Curing process efficiency (CI-1)</td>
<td>Film quality assessment (CI-2)</td>
</tr>
<tr>
<td></td>
<td>Net operational cost (CI-6)</td>
<td>Toxicity assessment (CI-3)</td>
<td>Operational safety (CI-4)</td>
</tr>
<tr>
<td>End use</td>
<td>Product cost (EI-3)</td>
<td>Toxicity during use (EI-1)</td>
<td>Coating performance assessment (EI-2)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Customer satisfaction factor (EI-4)</td>
</tr>
</tbody>
</table>

Many of these indicators from different stages of the life cycle are correlated with each other. A change in the value of one indicator from one stage can affect the indicator value at another stage. Thus, while analyzing the sustainability performance of this technology, a holistic view is very important. The developed framework pinpoints some of the important sustainability indicators which are correlated with each other. In Figure 4.5, these correlations are represented and the locations of indicators are set at respective time and length scales of measurement.
Figure 4.5. Correlations among sustainability indicators at various stages of life cycle of a paint material.

4.4 Case Study

In this research, we selected an automotive paint system for a case study to perform a life cycle based sustainability assessment. The objective was to carry out the sustainability assessment using selected indicator sets for environmental, social and economic performance and determine the correlation among the indicators selected from at least 2 of the stages of life cycle.

The stages of life cycle we selected for the case study are, Stage 1: Material selection and Stage 3: Paint spray/flash.
4.4.1 Stage 1: Material Selection and Preprocessing

This stage of life cycle deals with selection of paint formulation and raw materials required to manufacture the paint. This is a critical stage which has direct correlation with the final performance of the coating film. The material selection is based on the required final properties of the coating film. Several factors related to the toxicity, energy efficiency, safety and economics of other stages of life-cycle can be improved by improving the sustainability of this critical stage. In this case we considered the material selection criteria for the determination of all the indicator parameter values. The preprocessing consists of the information from the extraction/synthesis of each of the raw materials selected for the formation of paint matrix. Due to the unavailability of such data for preprocessing, the indicators related to preprocessing stage were neglected.

For the case study, we considered two examples of an automotive coating system with 2 coats: basecoat and topcoat. The basecoat for both the examples was a waterborne paint system while the topcoat in first example was solventborne clearcoat and in second example it was powder clearcoat. The sustainability assessment of the system with basecoat and topcoat paint formulations used in the auto industry was carried out by selecting the paint formulations shown in Table 4.2. The quantities of raw materials given in this table are in weight percent of the total weight of each of the paint system.
Table 4.2. Automotive basecoat and topcoat formulations.

<table>
<thead>
<tr>
<th>Waterborne basecoat</th>
<th>Solventborne clearcoat</th>
<th>Powder clearcoat</th>
</tr>
</thead>
<tbody>
<tr>
<td>Material</td>
<td>Quantity (%)</td>
<td>Material</td>
</tr>
<tr>
<td>Water</td>
<td>40</td>
<td>Naptha</td>
</tr>
<tr>
<td>Titanium dioxide</td>
<td>21</td>
<td>Xylene</td>
</tr>
<tr>
<td>Naptha</td>
<td>4</td>
<td>Methanol</td>
</tr>
<tr>
<td>Polyurethane</td>
<td>3</td>
<td>Melamine formaldehyde</td>
</tr>
<tr>
<td>Polyester resin</td>
<td>7</td>
<td>Ethylbenzene</td>
</tr>
<tr>
<td>Melamine formaldehyde</td>
<td>4</td>
<td>N-butyl alcohol</td>
</tr>
<tr>
<td>EGME(^a)</td>
<td>7</td>
<td>Cumene</td>
</tr>
<tr>
<td>2-Hexyloxyethanol</td>
<td>2</td>
<td>MTS(^b)</td>
</tr>
<tr>
<td>PMMA</td>
<td>10</td>
<td>Butyl acetate</td>
</tr>
<tr>
<td></td>
<td></td>
<td>PMMA</td>
</tr>
</tbody>
</table>

The waterborne basecoat (WB) and solventborne clearcoat (SC) formulations are commercial products from DuPont and powder clearcoat (PC) is from Seibert which are used for automotive painting operations (Papasavva et al., 2001). The WB system contains a polyester and polyurethane resins. Both the resins together form a binder which contributes to 7-14% (by weight) of total formulation. The carrying medium is water which accounts for 40-60% (by weight). The pigment is white titanium dioxide based (21% by weight) powder. The average thickness of the basecoat for the application on automotive body is expected to be 1.0 mil.

The SC system is an acrylic solventborne system. The binder for this system accounts for 54% of the total weight of paint formulation. The expected film thickness to obtain optimum properties using this coating system is 1.8 mils. The resin for PC system is the glycidyl acrylic polymer. The resin and crosslinker together contribute to 94% (by weight) of the total formulation. The optimum film thickness for this system should be 2.2 mils.
In order to carry out the sustainability assessment we have selected a set of indicators which address environmental, social and economic aspects of Stage 1 of the life cycle of a paint. For the paint formulation to be environmentally sustainable it should have a minimum quantity of raw materials which contribute to VOC. The formulation with higher quantity of water should be more sustainable. The social indicators are selected to compare the toxicity for human health and the predicted performance of the final coating. The waterborne basecoat is expected to provide the color, adherence and hardness to the surface of the automobile, whereas the clearcoat systems should provide more properties such as appearance, gloss, scratch resistance and recoatability cost, etc. Economic indicators deal with the composition of each paint and the material cost. The indicator matrix with relative assessment of both paint formulations is given in the table below.

<table>
<thead>
<tr>
<th>Economic</th>
<th>Environmental</th>
<th>Social</th>
</tr>
</thead>
<tbody>
<tr>
<td>Material Cost</td>
<td>Quantity of toxic raw material (VOC/gal)</td>
<td>Predicted performance of the coating (scale of 10)</td>
</tr>
<tr>
<td>Composition of paint (number of ingredients)</td>
<td>Energy consumed per unit job</td>
<td></td>
</tr>
</tbody>
</table>

For environmental indicators assessment, it can be observed from the formulations that the system with powder clearcoat should perform better because of zero VOC. In each coating formulation, it is the polymer resin which consumes maximum amount of energy for the synthesis. Both the coating systems (WB-SC and WB-PC) are used for coating on an SUV vehicle. Figure 4.6 shows the amount of energy consumed for the material for each job of painting an SUV vehicle.
For social sustainability assessment, both the systems performed equally. The predicted properties of the final film with the selected basecoat and clearcoat systems were similar.

The economic assessment was done by analyzing the material cost and total number of ingredients required to formulate each type of coating system. The overall sustainability assessment and comparison between two examples is given in Table 4.4.

Table 4.4. Sustainability assessment matrix for Stage 1 of the life cycle of paint.

<table>
<thead>
<tr>
<th>Sustainability Aspect</th>
<th>Indicator</th>
<th>Example 1 (WB/SC)</th>
<th>Example 2 (WB/PC)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Basecoat</td>
<td>Clearcoat</td>
</tr>
<tr>
<td>Economic</td>
<td>Composition of paint (no. of ingredients)</td>
<td>9</td>
<td>10</td>
</tr>
<tr>
<td></td>
<td>Material Cost ($/job)</td>
<td>63</td>
<td>40</td>
</tr>
<tr>
<td>Environmental</td>
<td>Quantity of toxic raw material (VOC/gal)</td>
<td>1.5</td>
<td>4.0</td>
</tr>
<tr>
<td></td>
<td>Energy consumed per job (MJ)</td>
<td>467</td>
<td>514</td>
</tr>
<tr>
<td>Social</td>
<td>Predicted performance of the coating</td>
<td>100%</td>
<td>100%</td>
</tr>
</tbody>
</table>
### 4.4.2 Stage 3: Paint Spray/Flash

After the paint is manufactured, it is applied on to the substrate using different types of application techniques such as brush application, roller application, conventional spray, airless spray, electrostatic spray and high volume-low pressure spray. In an automotive industry, paint is applied using manual or robotic spray guns installed in a sophisticated spray booth. In general, the spray booth is equipped with spray guns, ventilation systems and various accessory machines, tools, appliances and equipment that are necessary for operators to apply paint on the object surfaces to be coated, for instance, pumps, compressors, and personal protective equipment.

During the paint spray process, paint species present in the paint formulation are continuously released into the air. Although most of the paint gets deposited onto the surface of auto part, a substantial amount remains which pollutes the air inside the booth. Volatile organic compounds (VOCs) are the most hazardous substances emitted during certain spray processes, which can cause acute symptoms to the involved workers, including headache and dizziness, and sometimes exposure to suspected carcinogens. In case of nanopaint application, released nanoparticles further add to the contamination of the air. This air is taken away by the exhaust and mixes with the running water through the grids at the bottom of the booth. It results in increased pollution of the environment and water and risk to health care.

The efficiency of the spray application depends on the final film topology and the amount of energy consumed. A comprehensive sustainability assessment of the process can help in quantitative evaluation of the overall performance of this stage of life cycle. Table 4.5 shows the indicator sets for initial assessment of the paint spray application stage.
Table 4.5. Sustainability indicator matrix for paint-spray application stage.

<table>
<thead>
<tr>
<th>Economic</th>
<th>Environmental</th>
<th>Social</th>
</tr>
</thead>
<tbody>
<tr>
<td>Net energy usage</td>
<td>Energy efficiency</td>
<td>Film quality assessment</td>
</tr>
<tr>
<td>Total amount of paint applied</td>
<td>Net VOC emission</td>
<td>Human health burden</td>
</tr>
</tbody>
</table>

**Paint spray booth system.** In an automotive industry, paint spray system consists of several zones through which automotive parts are carried on a conveyor belt. A complete layout of a typical paint spray system in an automotive plant is given in Figure 4.7. The paint is applied using compressed air spray guns operated by robotic bells at automatic spraying chambers and manually by the workers at manual interior cut-in chambers. After each type of paint film (basecoat and clearcoat) is applied, the parts are carried to the “heated flash off” zone where the parts are kept under warm ambience to allow uniform film formation and evaporation of solvents. After flash-off, the coated auto parts are carried to the curing ovens for final film build and hardening. The curing ovens are adjusted at a very high temperature above the boiling points of all the solvents and crosslinking reaction temperature in order to allow the film formation. The lengths of curing zones are adjusted such that the coated parts spend enough time under high temperatures to ensure high conversion and efficiency of crosslinking reactions.
There are 29 spray bells installed at different stations throughout the spray chambers as shown in Figure 4.7. The downdraft air is supplied by 15 exhaust fans of power of 25-hp each which are located at different places inside all the chambers. The electric draw of each motor of the exhaust fan is 18.64 kW. Conventionally in the paint spray operation, the energy is consumed by exhaust fans to provide ventilation air flows and by the spray facilities to generate shaping air flows. Industrial practice shows that maintaining ventilation air flows consumes much more power than shaping air flows. Thus in this work, the total energy consumption of a paint spray system is quantified solely using the energy consumed by the ventilation system. The power required for the exhaust fans \( E_d \) kW is proportional to the velocity of the exhaust air flow \( v_{dd} \) (Perry and Green, 1997).

\[
E_d = cv_{dd}
\]  

(4.1)
where \( c \) is a given coefficient (kW/m/s).

**Environmental assessment.** The environmental assessment was done using the data of two cases of industrial practices. Both cases assumed the settings for a sports utility SUV car getting painted in the spray booth. The input parameters and calculated values are given in Table 4.6. The basecoat was water-borne acrylic paint whereas the clearcoat was solvent-borne acrylic paint. In both the cases, the data related to application of waterborne basecoat onto the car body was used for the assessment. The VOC concentration of the basecoat paint was 0.4 kg/l. Thus, the VOC mass flow-rate was calculated as:

\[
\text{Mass flow rate of VOC} = \text{TE} \times \text{F. D.} \times \text{VOC concentration} \quad (4.2)
\]

where \( \text{TE} \) is the transfer efficiency (40\%) of the paint material and F. D. is a paint fluid delivery.

<table>
<thead>
<tr>
<th>Case No.</th>
<th>fluid delivery (cc/min)</th>
<th>Downdraft (FPM)</th>
<th>Downdraft (m/s)</th>
<th>VOC mass flow-rate*10^4 (kg/s)</th>
<th>c (energy coeff.) kW/m/s</th>
</tr>
</thead>
<tbody>
<tr>
<td>Case 1</td>
<td>21400</td>
<td>1159</td>
<td>5.88</td>
<td>570</td>
<td>47.48</td>
</tr>
<tr>
<td>Case 2</td>
<td>22100</td>
<td>1160</td>
<td>5.89</td>
<td>589</td>
<td>47.44</td>
</tr>
</tbody>
</table>

The energy efficiency was quantified using the energy coefficient. It gives a measure of amount of energy that is consumed by the ventilation system.

The net VOC emission was quantified using a relative parameter. The net VOC content inside the booth is proportional to the quantity of fluid delivered and the downdraft air. We define this relative term as “VOC factor”, which can be calculated as:

\[
\text{VOC factor} = \frac{\text{VOC mass flow rate (kg/min)}}{\text{total downdraft air in FPM}} \quad (4.3)
\]
Social assessment. The quality of the film was assessed by the percentage of paint film that was sprayed within the specified limit of film thickness. The standard specifications of film thicknesses for basecoat and clearcoat are tabulated in Table 4.7.

Table 4.7. Materials engineering paint film thickness specifications.

<table>
<thead>
<tr>
<th>for vertical and horizontal areas</th>
<th>Vertical (mils)</th>
<th>Horizontal (mils)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Basecoat</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Min Spec</td>
<td>1.0</td>
<td>1.0</td>
</tr>
<tr>
<td>Max Spec</td>
<td>1.2</td>
<td>1.2</td>
</tr>
<tr>
<td>Clearcoat</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Min Spec</td>
<td>1.8</td>
<td>1.8</td>
</tr>
<tr>
<td>Max Spec</td>
<td>2.0</td>
<td>2.0</td>
</tr>
</tbody>
</table>

The average film thicknesses at different locations of automotive parts are obtained by selecting 90 random data points distributed at car body’s hood, roof, doors and lift gate. The film thicknesses at each point were measured and percentage of data points within specified limit, below the limit and above the limit was determined. For two cases under consideration, the values obtained are given in Table 4.8.

Table 4.8. Percentage of data points measured within the paint specification and outside the specification.

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Basecoat</td>
<td>56.3%</td>
<td>19.7%</td>
<td>23.9%</td>
</tr>
<tr>
<td></td>
<td>Clearcoat</td>
<td>19.2%</td>
<td>67.6%</td>
<td>13.1%</td>
</tr>
<tr>
<td></td>
<td>Avg.</td>
<td>37.7%</td>
<td>43.7%</td>
<td>18.5%</td>
</tr>
<tr>
<td>2</td>
<td>Basecoat</td>
<td>78.9%</td>
<td>1.9%</td>
<td>19.2%</td>
</tr>
<tr>
<td></td>
<td>Clearcoat</td>
<td>25.8%</td>
<td>63.8%</td>
<td>10.3%</td>
</tr>
<tr>
<td></td>
<td>Avg.</td>
<td>52.3%</td>
<td>32.8%</td>
<td>14.8%</td>
</tr>
</tbody>
</table>
Economic assessment. The operational cost was determined by calculating the total energy required for running the spray operation. The energy is consumed by the system for ventilation airflows and shaping airflows. The values of ventilation airflows are recorded in FPM whereas the shaping airflows are recorded in liter/min. The surface area of the car is 131.8 ft². The energy (in kW) consumed by the shaping airflows was calculated by the following formula.

\[
\text{Energy consumed} = \frac{\text{Airflowrate} \cdot (\text{SP}-\text{TP}) \cdot 0.745}{6356 \cdot \eta}
\]  

(4.4)

where \((\text{SP}-\text{TP})\) is the static pressure with \(\text{SP}\) being outlet pressure (inches of water) and \(\text{TP}\) the inlet pressure (inches of water) from spray guns. \(\eta\) is the efficiency of the motor associated with each spray gun. The shaping air flow rate was calculated in CFM. The total energy consumed by ventilation and shaping airflows for 1 minute of operation in Case 2 (720 kW) is greater than in Case 1 (691 kW). The cost of each unit of electricity required to generate this energy was calculated by a two factor utility cost equation which is given as:

\[
C_{S,u} = a \cdot (\text{CE PCI}) + b \cdot (C_{S,f})
\]  

(4.5)

where \(C_{S,u}\) is the price, \(a\) and \(b\) are the utility coefficients, \(C_{S,f}\) is the fuel price in $/gal and CE PCI (Chemical engineering plant cost index) is the inflation parameters for all the projects in U.S. (Ulrich and Vasudevan, 2006). The value of \(a\) is \(1.3 \times 10^{-4}\) and \(b\) is 0.010. The inflation parameter is selected to be in the range of 550 to 600. The fuel price was assumed to be $3.5/gal. Thus, the unit value of electricity required to generate the energy for case 1 was,

\[
C_{S,u} = 1.3 \times 10^{-4} \times 575 + 0.010 \times 3.5 = $0.1097/\text{kWh}
\]  

(4.6)

The human health burden was measured as a function of mass flow rate of VOC. The overall assessment and comparison between two cases is given in Table 4.9. Figure 4.8
represents the assessment results in bar-chart, where the “green” colored bars indicate the more sustainable case and “red” colored bars indicate less sustainable case. It can be observed that Case 1 is more sustainable compared to Case 2 for four indicators out of selected six selected indicators.

Table 4.9. Overall sustainability assessment of Stage 3.

<table>
<thead>
<tr>
<th>Sust. Factor</th>
<th>Indicator</th>
<th>Case 1</th>
<th>Case 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Environmental</td>
<td>(1) Net VOC emission factor (\times 10^3)</td>
<td>2.954</td>
<td>3.048</td>
</tr>
<tr>
<td></td>
<td>(2) Net energy consumption (kW/m/s)</td>
<td>47.48</td>
<td>47.44</td>
</tr>
<tr>
<td>Social</td>
<td>(3) Film quality assessment (% In Spec.)</td>
<td>37.7%</td>
<td>52.3%</td>
</tr>
<tr>
<td></td>
<td>(4) Human health burden*10^4 (kg/s)</td>
<td>570</td>
<td>589</td>
</tr>
<tr>
<td>Economic</td>
<td>(5) Net operational cost ($)</td>
<td>4548</td>
<td>4739</td>
</tr>
<tr>
<td></td>
<td>(6) Quantity of paint applied (cc/min)</td>
<td>21400</td>
<td>22100</td>
</tr>
</tbody>
</table>
Figure 4.8. Overall comparison of the two cases for the assessment of Stage 3.
CHAPTER 5
CFD MODELING OF NANOPAINT APPLICATION SYSTEM – METHODOLOGY AND SYSTEM DESCRIPTION

In an automotive industry, the topcoat layers are applied on the car bodies by manual or robotic spray-guns installed inside a spray booth. The paint is applied under optimized conditions of temperature and humidity. For painting of irregularly shaped objects, manual spray application is preferred. A typical paint spray technique uses a compressed gas to atomize the paint material and direct it at a high velocity to the receiving panel through spray guns (Colbert and Cairncross 2006). Although most of the sprayed paint lands on the target, a certain amount of paint is carried by the ventilation air and enters the water in the drain through grids on the booth floor. The effectiveness of paint spray and final coating film quality depends largely on the operational settings, such as paint flow rate, shaping air velocity and electrostatic voltage (Li et al. 2007). During paint spray operation, the workers inside the spray booth may become prone to multiple types of hazards as a result of exposure to VOC, heat overload, electric shock or electrocution, noise and other types of body injuries. It has been recognized that VOC’s affect human health and productivity the most in the manual paint application process due to possible acute symptoms to workers, including headache, dizziness and exposure to suspected carcinogens (Feldman et al. 1999; Kim et al. 2000).

For nanopaint, the overspray paint mist is laden with VOC and toxic nanoparticles. It may severely pollute the booth air as well as the water flowing through the washout. Dunnett
(1994) studied how the presence of contaminants and the air flow pattern affect the workers’ health. If exposed to severe concentrations of toxic nanoparticles such as nanosilica or nano-titanium dioxide, it can cause fatal health disorders (Reijnders, 2009). Thus, it is important to develop an optimized paint-spray operation which would be more sustainable in terms of environmental emission, coating film quality and high transfer efficiency.

5.1 Benefits of CFD Modeling for Paint-Spray System

The study of paint-spray process requires detailed analysis of fluid flow, emission, atomization parameters and film topology changes. Such analysis is only possible through computational modeling where individual paint droplets can be tracked and the concentration at various locations in the spray-booth can be precisely measured. The computational model which simulates the paint spray application method should also consider the evaporation of solvent from individual paint droplets during the course of spray. A thorough investigation of the air quality and energy consumption in the paint spray booth requires an understanding of the complex fluid flow phenomena in the occupied space of the ventilation air, for which finite difference-based and Computational Fluid Dynamics (CFD)-based modeling and simulation are suitable techniques (Tabor et al. 2012).

Li et al. (2013) demonstrated the potential of CFD-based modeling methodology for the comprehensive analysis of a manual paint-spray system. It was shown that the spray booth geometry and operation parameters could be adjusted to control the VOC emission below threshold limit value and to improve the energy efficiency. But, it was assumed that the VOC was evaporated instantaneously after the paint was injected from the gun. A CFD modeling was
also used to characterize the mass transfer and flow of VOC in the booth environment (Flynn and Sills 2000; Fogliati et al. 2006). It was applied to simulate the breathing-zone concentration for a paint spray method in a simple spray booth unit with cross-flow ventilation [Flynn, 2000]. Kim et al. (2002) analyzed a small-scale painting process with various exits of air leaving at different suction velocities. Lu and Howarth (1996) presented numerical results of the prediction of air velocities, aerosol particle deposition and migration in two interconnected ventilated zones. Dunnett (1994) presented a study on the effect of the presence of workers on the air flow pattern and the factors affecting the exposure to contaminants. However, more systematic and quantitative VOC emission characterization and energy-efficient VOC reduction analysis for surface coating systems have not been thoroughly studied yet.

5.2 Paint Spray System Design

Since last decade, nanopaints are seen to be replacing conventional paints at several application areas including automotive, industrial, architectural, military, etc. The induced nanoparticles of different size, shape and chemical nature provide substantial improvement in existing product performance and also bring in newer functionalities such as self-cleaning, super-scratch, anticorrosion, etc. Despite of all the benefits from nano-induced coating materials, it poses serious concern to environmental safety and health care by unintended release of nanoparticles into the surrounding medium during manufacturing, application and use phases. For automotive paint application, the material is applied using conventional, airless or electrostatic spray techniques. During spraying, the nanoparticles are prone to release in the surrounding atmosphere and also in the drainage waste water causing serious hazards to
atmosphere. Thus, continuous efforts are being made to optimize these spray techniques and reduce the risk factors associated with the use of nanocoatings.

A typical paint spray technique uses a compressed gas to atomize the paint material and direct it with high velocity to the receiving panel through spray guns (Colbert and Cairncross, 2006). In automotive sector, the paint spray operation is performed by robotic manipulators. The paint particles are ejected towards the vehicle body inside the spray booth at a high speed. Although most of the sprayed paint lands on the target, a certain concentration of paint is carried by the downdraft air and enters the water in the drain through grids on the booth floor. The efficiency of paint-spray and the final coating film quality depends on several factors such as paint flow-rate, paint injection velocity, atomization method, ventilation air velocity, spray angle, the distance between the gun and the substrate, etc. (Li et al., 2007) The overspray released during the painting operation is removed by the downdraft air that flows through the booth geometry and is absorbed by the water flowing underneath the exhaust grid. The removal of the overspray can be more efficient with higher downdraft velocity of ventilation air. But, it leads to increased energy consumption. High transfer efficiency can also minimize the overspray concentration. To improve the process economy and coating film quality, high transfer efficiency is critical. While spraying nanopaint, the exposure to nanoparticles via different pathways such inhalation or dermal can cause severe health hazards (Klaine et al., 2008). TiO₂, SiO₂, MMT clay, Al, etc. are among the most commonly used nanoparticles in coating formulations. Research has shown that exposure to high concentrations of such material may result into possible acute symptoms to workers, including headache, dizziness and exposure to suspected carcinogens and sometimes can also affect central nervous system (Feldman et al., 1999; Kim et al., 2000).
A paint spray unit includes a spray booth, spray guns/bells, ventilation system, tools, appliances and equipment such as pump, compressor, conveyor belt and personal protective gears which are necessary for an operator to apply paint on the object surface to be coated. Paint material is sprayed using spray bells rotating at high speed. A typical automotive paint spray application unit is shown in Figure 5.1.

The air inside the bells atomizes the paint material and breaks it into tiny paint particles. For nanopaint, the particles are filled with individual or agglomerated nanoparticles inside it. The paint particles having nanoparticles are then ejected from the bells at high speed to form a film on automotive panels. The transfer efficiency of water-borne paint material is generally around 30-40 % while that of solvent-borne is 60-80%. The remaining paint material is emitted into the surrounding atmosphere resulting in contamination of the air inside the spray booth.
This contamination includes high concentrations of nanoparticles and VOCs. A sophisticated ventilation system is used to remove the air contaminated with these paint particles.

As described before, the painting of irregularly shaped objects is performed inside a manual paint-spray booth. As described by Li et al. (2013), a manual paint-spray facility consists of an air ventilation unit, spray-gun and a placement of substrate to be coated. Figure 5.2 shows a typical manual spray-booth design.

![Figure 5.2. Sketch of manual paint-spray booth.](image)

The design consists of a paint spray booth with operational parameters adopted from a real industrial practice. According to Lin et al. (2005), the spray booth is applicable to chemical agent resistant coatings, production paints and camouflage and powder paints application.
The size of the booth is 19 ft by 39 ft, which is housed in a room of dimensions approximately equal to 350 ft long and 60 ft wide. Inside the booth, a conveyor belt is located for moving parts through the working area. Paint is applied by a compressed air-forced paint gun operated by a worker positioned next to the substrate panel. Generally, an epoxy primer coat is first applied by spraying, followed by a topcoat films. After painting, the parts are carried out of the booth into dryers for flash-off and curing. At room temperature, the primer coat dries to touch in 15~45 minutes and the topcoat dries to touch in 15 minutes. The appropriate paint thickness of 1.0~1.5 mils for the primer and 1.8 mils for the topcoat is achieved. A successful paint application job should produce a smooth, continuous, adherent paint film free of runs, sags, blisters, orange peel, streaks, craters, blotches, fish eyes and pinholes.

The roof of the paint booth is open and air is drawn through a 13 ft wide opening down to both exhaust air intakes with 4 ft wide slots located on each side of the booth floor. Four 25-horsepower motors power the exhaust fans to discharge paint fumes outdoor, which give the exhaust air flows at 80,000 CFM (Cubic Feet per Minute). During the painting operation, fresh air enters the booth from the open roof and flows down to the floor continuously (see Figure 5.2). Both ends of the booth are kept open. The heating and ventilation facility operates continuously at all times.

5.3 Integrated Modeling Methodology

The transport of ventilation air, shaping air, injected paint material, VOC and nanoparticles can be thoroughly examined through CFD-based simulation. Figure 5.3 represents a general scheme of the modeling methodology and different types of models that are used for
the calculation of energy efficiency, VOC emission and nanoparticles emission during the spray process. This methodology consists of five different models which are described below. Four of the models were used by Li et al. (2013) for the analysis of VOC emission and energy efficiency of a paint booth for conventional paint spray. In this work, the discrete phase model was combined with the previous model to make it suitable for the analysis of nanoparticles-based paint systems.

5.3.1 Air Flow Model

The air flow pattern within the spray booth is determined by the geometry of the booth and ventilation system’s design and setting. It can be characterized by the following dynamic models, based on the mass conservation principle (see Eq. (5.1)), the momentum conservation...
principle (see Eq’s. (5.2)-(5.4)), as well as the realizable k-ε turbulence principle (see Eq’s. (5.5) and (5.6)) (Colbert and Cairncross 2006; Ellwood and Braslaw 1998; Shah et al. 2006). The air is a continuous phase in the simulation. The prediction of flow of continuous phase is obtained through a realization k-ε model. It is an improved turbulence model as compared to the previous standard k-ε model (Boulet et al., 2010). The governing equations for these models are given below.

\[
\frac{\partial}{\partial t} \rho_a = -\nabla \cdot (\rho_a \mathbf{v}) + S_m
\]

(5.1)

\[
\frac{\partial}{\partial t} (\rho_a \mathbf{v}) = -\nabla \cdot (\rho_a \mathbf{v} \mathbf{v}) - \nabla p + \nabla \cdot \mathbf{\tau} + \rho_ag + \sum_i \left\langle \dot{S}_i \right\rangle
\]

(5.2)

where

\[
\tau = \left( \mu_a + \mu_i \right) \left( \nabla \mathbf{v} + (\nabla \mathbf{v})^T \right) - \frac{2}{3} \left( \mu_a + \mu_i \right) \left( \nabla \cdot \mathbf{v} \right) I
\]

(5.3)

\[
\mu_i = \rho_a C_{\mu} \frac{(k)^2}{\epsilon}
\]

(5.4)

and

\[
\frac{\partial}{\partial t} (\rho_a k) = -\nabla \cdot (\rho_a \mathbf{v} k) + \nabla \cdot \left[ \left( \mu_a + \mu_i \right) \nabla \epsilon \right] + G_k + G_b - \rho_a \epsilon - Y_M
\]

(5.5)

\[
\frac{\partial}{\partial t} (\rho_a \epsilon) = -\nabla \cdot (\rho_a \mathbf{v} \epsilon) + \nabla \cdot \left[ \left( \mu_a + \mu_i \right) \nabla \epsilon \right] + \left( C_{1a} \frac{\epsilon}{k} C_{3b} G_b + C_{4a} S_c - C_{2a} \rho_a \frac{\epsilon^2}{k \sqrt{\epsilon}} \right)
\]

(5.6)

where \( \rho_a \) is the air density (kg/m\(^3\)), \( \mathbf{v} \) is the velocity of air (m/s), \( S_m \) is the rate of mass addition into the gas phase per unit volume from the dispersed second phase (kg/m\(^3\)s), \( p \) is the pressure (Pa), \( \tau \) is the stress tensor of the air (Pa), \( \mathbf{g} \) is the acceleration due to gravity (m/s\(^2\)), \( \dot{S}_i \) is the external force on gas phase per unit volume from the \( i \)-th species particle (N/m\(^3\)), \( \mu_a \) and \( \mu_i \) are, respectively, the shear viscosity (Ns/m\(^2\)) and turbulent viscosity (Ns/m\(^2\)), \( I \) is the unit
tensor, $C_\mu$, $C_{1\varepsilon}$, $C_{2\varepsilon}$, $C_{3\varepsilon}$, and $\sigma_k$ are the model constants, $k$ is the kinetic energy of turbulence ($m^2/s^2$), $\varepsilon$ is the dissipation rate of turbulence ($m^2/s^3$), $G_k$ is the generation of turbulent kinetic energy due to the mean velocity gradient ($kg/ms^3$), $G_b$ is the generation of turbulent kinetic energy due to buoyancy ($kg/ms^3$), and $Y_M$ is the contribution of the fluctuating dilatation in compressible turbulence to the overall dissipation rate ($kg/ms^3$). More details about the realizable $k-\varepsilon$ model can be found in the ANSYS FLUENT theory guide (2009).

The air within the spray booth is initially static. The fresh air then begins to blow from booth intake to the exhaust. It is interfered by the shaping air flow near the spray guns. The air flows out of the booth to the environment through exhaust grids located on the floor. These operating conditions inside the spray booth are defined by the following initial and boundary settings of the air flow model.

$$v(x,t) = \begin{cases} 
  v_{SG} & \text{at the outlet of spray guns, } t > 0 \\
  v_{DD} & \text{at booth intakes, } t > 0 \\
  0 & \text{at other physical boundaries except the exhausts, } t > 0 \\
  0 & \text{in the spray booth, } t = 0
\end{cases} \quad (5.7)$$

where $v_{SG}$ is the air velocity from the spray gun (m/s), and $v_{DD}$ is the air velocity of the downdraft (m/s).

### 5.3.2 Species Transport Model

Paint species are ejected from the spray guns with a momentum that is governed by two types of forces: the drag force from the surrounding turbulent air flow and the gravitational force.
The velocity of paint species \((u)\) can be modeled by the Newton’s second law of motion as shown in the following equations (Colbert and Cairncross 2006; Im et al. 2004).

\[
\frac{du(t)}{dt} = \frac{3\pi\mu_aD_p}{m}(\nu(t) - u(t))f + g
\]  

(5.8)

where

\[
m = \frac{1}{6}\pi(D_p)^3\rho_p
\]  

(5.9)

\[
f = \begin{cases} 
1 + \frac{(Re_r)^{2/3}}{6}, & Re_r < 1,000 \\
0.0183 Re_r, & 1,000 \leq Re_r \leq 3 \times 10^5 
\end{cases}
\]  

(5.10)

\[
Re_r = \frac{D_p[v(t) - u(t)]\rho_a}{\mu_a}
\]  

(5.11)

where \(u\) (m/s) is the velocity of the species having particle diameter \(D_p\) (m) and the mass \(m\) (kg), \(\nu\) (m/s) is the velocity of air surrounding the species, \(f\) is the drag factor, \(\rho_p\) is the paint density (kg/m\(^3\)) and \(Re_r\) is the Reynolds number.

The initial species velocity from the spray guns for this model is \(u_0\):

\[
u(0) = u_0
\]  

(5.12)

The species trajectory \(x_{tr}(t)\) is determined from the species velocity that is determined by Eq’s. (5.8) – (5.12). The initial position of all paint species is assumed to be at the outlet of the spray gun. The dynamic model becomes:

\[
\frac{dx_{tr}(t)}{dt} = u(t)
\]  

(5.13)

\[
x_{tr}(0) = x_0
\]  

(5.14)
The models for the air flow and the species transport can be solved numerically. The solution can be obtained at various locations within the booth and at different times. The results from the CFD simulation can characterize the emission and diffusion of VOC’s in the system as well as the effect of ventilation during the paint spray operation.

The VOC emission and energy consumption of the paint spray booth can be quantified by the following models.

### 5.3.3 VOC Concentration Model

The air quality inside the spray booth is quantified by measuring the mean VOC concentration \( C_m, \text{ kg/m}^3 \) in the system; which is the ratio of total mass of VOC’s to the total space of the booth \( V_{bh}, \text{ m}^3 \).

\[
C_m = \frac{\int V_{bh} \rho_s \, dV}{V_{bh}}
\]  

(5.15)

where \( \rho_s \) is the species flow density (kg/m³) obtained from the species transport model, and \( Y \) is the mass fraction of VOC in the paint species flow. This model accounts for the VOC content released in the air from the paint being applied onto the substrate and also the overspray. The VOC evaporation from the deposited wet film is neglected as the substrate is immediately moved from the paint booth to the next production stage.
5.3.4 Energy Consumption Model

In paint spray operation, energy is consumed by exhaust fans that provide ventilation air and by the spray facilities that generate the shaping air. Industrial practice shows that maintaining ventilation air flows consumes much more power than keeping shaping air flows. Thus, in this work, the total energy consumption of a paint spray system is quantified solely using the energy consumed by the ventilation system. The power required for the exhaust fans ($E_d$, kW) is proportional to the velocity of the exhaust air flow ($v_{dd}$) (Perry and Green 1997) i.e.,

$$E_d = cv_{dd}$$

(5.16)

where $c$ is a given coefficient (kW/m/s).

5.3.5 Discrete Phase Model

In addition to solving the transport equations of the continuous phase, FLUENT also provides a tool for simulating the discrete second phase. The paint acts as a discrete second phase, which is sprayed in a continuous air domain. The dispersed phase can exchange mass, momentum and energy with the continuous phase. It is treated by the Lagrangian discrete phase model (DPM) which follows the Euler-Lagrange approach. The DPM is combined with species transport model to calculate the paint flow trajectories and nanoparticles emission (See Figure 5.3). This discrete phase is sprayed in the form of large number of tiny particles representing real paint droplets with the same properties. By this representation, one can control the size distribution of paint droplets that are being injected from the gun. The droplets are traced through the booth environment at each time interval of iteration. The trajectory of individual
droplet is determined by solving the equation of motion for each droplet. This equation of motion is obtained from the particle force balance written in a Lagrangian frame of reference (See Eq. 5.17). This equation considers the forces in the $i^{th}$ direction of cartesian coordinates due to drag, gravity and pressure gradient. The thermophoretic and Brownian motion forces are neglected.

$$\frac{du_i}{dt} = F - \frac{6}{D_i} \rho \left( \rho_p - \rho \right) + \frac{1}{2} \frac{d}{\rho_p dt} \left( u_i - u_i^p \right)$$ (5.17)

$$F_{D_i} = \frac{18 \mu \rho_p C_D R_e |u_i^p - u_i|}{d_p^2}$$ (5.18)

$$R_e = \frac{\rho d_p |u_i^p - u_i|}{\mu}$$ (5.19)

where $F_{D_i}$ is the drag force per unit particle mass, $u_i$ and $u_i^p$ are the velocities of fluid phase and particles respectively, $\rho$ and $\rho_p$ are the densities of the fluid phase and particles respectively, $C_D$ is the drag coefficient, $d_p$ is the particle/droplet diameter, $\mu$ is the viscosity of the discrete phase, $g_i$ is the acceleration due to gravity, and $R_e$ is the Reynolds number.

During the spray of stable nanopaint material, the nanoparticles are not released in the natural form, but they are surrounded with resin material. Essentially, the overspray paint mist carries the nanoparticles in the booth environment. The emission of nanoparticles thus depends upon the paint droplet size and the concentration of the overspray. The simulation of such mixture of species is accomplished by activating the FLUENT multicomponent particle law (law 7). It allows FLUENT to model each droplet consisting of multiple components with different masses and densities. The injected particles are allowed to collide or break on interaction with each other in the air and on the deposited film. The mass of each paint droplet changes depending upon the evaporation of solvent and collision and break-up phenomena. Droplet spreading on the substrate film depends on multiple factors, such as, droplet size, injection speed,
angle of impact, paint surface tension, viscosity and surface properties such as surface energy, surface roughness, porosity, etc. The amount of deposition and spreading is determined by the driving and resisting forces. The driving force is provided by the kinetic energy of the droplet, whereas resistive force is provided by viscosity and surface tension of the paint material. When the droplet impacts the surface, its initial spherical shape may change into oval-like form. In FLUENT, the variation in the droplet shape is accounted by the “dynamic drag model”. The distortion of shape significantly affects the drag coefficient of the droplet and the value changes significantly from the drag coefficient of the spherical droplet. The dynamic drag model calculates the drag coefficient depending on the extent of distortion as given by Eq. 5.20.

\[ C_D = C_{D, sphere}(1 + 2.632y) \]  

where \( y \) is the extent of droplet distortion from spherical to the maximum of disc like shape. In the case of zero distortion \( (y = 0) \), the drag coefficient of a sphere is used, while the maximum distortion \( (y = 1) \) produces the drag coefficient of a disc shape.

### 5.4 Theory for Paint Spray Atomization and Modeling

A spray is a system of droplets immersed in a continuous gaseous phase. Sprays can be produced in a variety of ways. Most of the spray application devices achieve atomization of bulk liquid by creating a high velocity between the liquid and the injected gas required to atomize the liquid. In order to atomize the bulk liquid into tiny droplets, energy needs to be supplied to the system. This energy can be provided to the liquid material by employing either high pressure, or kinetic acceleration. Flowing air bubbles bursting at the liquid surface lead to its conversion into small droplets which are then sprayed out of the nozzle of the spray applicator.
The important properties for the process of atomization of any liquid are density, surface tension and viscosity. The atomization of the liquid forming tiny droplets occurs when the stable liquid with certain surface tension and viscosity is influenced by the disruptions from many internal and external forces. When the disruptive force barely exceeds the surface tension of the liquid, it leads to atomization and formation of droplets. Most of the bigger droplets are unstable during the initial breakup of the liquid sheet and form smaller droplets by undergoing further disintegration. Thus, the characteristics of a liquid spray are governed by the droplet sizes produced during primary atomization as well as by the extent to which these droplets are later disintegrated into smaller ones during secondary atomization.

The diameter of droplets which are formed from the flowing liquid through a thin circular outlet of the spray gun nozzle can be calculated from the following equation (Lefebvre, 1989).

\[ d = \left( \frac{6\sigma d_0}{\rho_L g} \right)^{1/3} \]  

where \( d_0 \) is the outlet diameter, \( \sigma \) is the liquid surface tension, \( \rho_L \) is the liquid density and \( g \) is the gravity constant. The effect of the aerodynamic drag force on break-up of droplets can be summarized using a Weber number as shown in following equation.

\[ \text{We} = \frac{\rho_{\text{air}} U^2 d}{\sigma} \]  

where \( \rho_{\text{air}} \) is the gas density, \( U \) the relative velocity and \( d \) the droplet diameter.

In spray coating system, the liquid paint is atomized at the nozzle by pressure or ultrasounds and then it is directed towards the panel by a flowing gas. The tiny droplets that are formed after atomization land on the surface producing either a full wet layer, in the case of large paint-flow rate, or sparse depositions and arrangements of dots on the surface, in the case of smaller flow rates. This droplet spreading in spray coating application is influenced by various
factors such as, droplet sizes, impact velocity, impact angle, liquid surface tension, liquid rheology and substrate surface properties such as surface energy, surface roughness and porosity.

When a droplet hits the surface, its initial spherical shape is forcefully converted into a pancake like shape. The spreading of paint droplets is decided by the balance of driving and resisting forces between the droplets and the surface. The driving force is provided by the kinetic energy of the droplet, whereas resisting force is provided by the surface tension and viscosity of the paint material. The relative values of the surface and kinetic energies can be represented with Weber number. For large values of the Weber number the kinetic energy overcomes the surface tension of the paint and spreading occurs. Whereas, for low values of Weber number, (e.g. low impact velocity,) the droplets tend to stay spherical and minimal wetting of the surface takes place.

Droplets are generated by high pressure of the gas during an atomization process. The injection velocity from the nozzle can be of the order of 100 m/s. However, a reduced velocity is used for spray nozzles with larger cross sectional area. But, such adjustment is not recommended as the surrounding air from ventilation may affect the jet velocity and slow down the droplets flowing towards the surface. Typical droplet sizes for paint material injection through spray guns are given in the literature which fall between 20-80μm (Toivakka, 2003).

5.4.1 Concept of Particles and Parcels

During DPM simulation, the spray module tracks the discrete phase movement throughout the system domain. In our case, the discrete phase is paint material. This discrete phase is allowed to exchange energy, mass (evaporation) and momentum (drag) with the air
(continuous phase). In case of spray application, the injected paint material is converted into millions of tiny droplets which flow through the simulation domain. During unsteady DPM modeling, the tracking of each of the millions of individual droplets is extremely expensive in terms of computational efficiency. Thus, in order to improve computational efficiency, FLUENT code in fact tracks “parcels” and not particles/droplets.

Each parcel is a representative entity of the behavior of many identical droplets after they are sprayed from the gun. The behavior of the parcels is determined by the behavior of individual particles. The total numbers of parcels that are generated are determined by various factors, such as, total mass flow rate, number of particles, particle size distribution, number of holes on the nozzle tip, and other spray injection inputs.

In general, the number of parcels is calculated by the following equation (Eq. 5.23).

\[
N_{\text{parcels}} = N_{\text{tracks}} \times N_{\text{sizes}}
\]  

(5.23)

where \(N_{\text{tracks}}\) is the number of random locations on the gun through which the paint material is injected and \(N_{\text{sizes}}\) is the number of diameters defined in Rosin-Rammler option for particle size distribution.

The number of particles in each of the parcels is calculated by Eq. 5.24.

\[
N_{\text{particles}} = \frac{\dot{M} \times \Delta t}{N_{\text{parcels}} \times \rho_p \times V_{\text{p}i}}
\]  

(5.24)

where \(\dot{M}\) is the mass flow rate of the paint, \(\Delta t\) is the time step of injection, \(\rho_p\) is the particle/droplet material density, \(V_{\text{p}i}\) is the particle/droplet volume. FLUENT tracks these parcels carrying different number of particles at all the time during simulation. After each instant of time iteration, the number of parcels is recalculated. The particles can exchange among different parcels during the course of simulation.
5.4.2 Boundary Conditions

The droplet-substrate interaction is an important aspect responsible for formation of the paint film on the panel surface. FLUENT provides a “wall-film” boundary condition which can cause the deposition of the paint on the panel. It allows the fluid droplets to impinge on the surface and form a thin film. The wall-film considers four key phenomena of interaction between droplets and sample based on the impact energy and wall temperature: spreading, sticking, rebound and splashing (Stanton et al. 1998 and O’Rourke 2000). Eq. 5.25 represents the impact energy ($E$) as a function of material parameters and the film thickness.

$$E^2 = \frac{\rho_p u_p^2 d_p}{\sigma_p} \left\{ \frac{1}{\min\left(\frac{h_0}{d_p}, 1\right)} + \frac{\delta_{bl}}{d_p} \right\}$$  \hspace{2cm} (5.25)

where $\rho_p$ is the paint density, $u_p$ is the droplet velocity, $d_p$ is the diameter of the droplet, $\sigma_p$ is the surface tension of the paint material and $\delta_{bl}$ is the boundary layer thickness. If the wall temperature is below the boiling temperature of the liquid, sticking, spreading and splashing occur. The number of splashed droplets is set to the default value of 4.
CHAPTER 6

CFD MODELING OF NANOPAINT APPLICATION SYSTEM – ANALYSIS OF ENVIRONMENTAL EMISSIONS AND COATING QUALITY

The CFD-based integrated modeling and simulation methodology which was described in Chapter 5 was used to investigate an industrial case of a paint-spray booth. Various geometries and operational settings were simulated and analyzed for determination of an optimum set of working parameters for minimal VOC emission and energy consumption. The test cases of various spray-booth systems and detail analysis can be found in Li et al. (2013). The VOC emission analysis was performed using species transport model and VOC concentration model. The analyses of nanoparticles emission, coating film topology, transfer efficiency were performed by integrating a DPM theory with the models for air flow, species transport and VOC concentration.

6.1 Spray-booth and Paint Material Details

The paint booth geometry for the base case was adopted from the industrial example (Li et al., 2013). In automotive industries, nanopaints have major applications in clearcoat formulations. In this work, the paint material selected for modeling was an epoxy-based clearcoat resin, product no. D.E.R. 538 (www.dow.com/scripts/litorder.asp?filepath=/296-00312.pdf). For nanopaint formulation, silicon dioxide nanoparticles of uniform size distribution
of 100 nm were dispersed into the coating matrix (Huynh, 2006). The solvent used for all paint formulations was acetone. Based on the industrial data, the paint flow rate was selected as 150 ml/min (Im et al. 2004). The shaping air flow rate from the spray gun is set to 5 m/s. Such a high value of shaping air flow avoids the coating spray quality from getting affected by the downdraft ventilation air. The paint droplets were injected at a high velocity of 20 m/s. For the simulation and analysis of the paint-spray through DPM, three paint systems were selected as: 1) nanopaint I (with 10% (by weight) nanoparticles); 2) nanopaint II (with 5% (by weight) nanoparticles); and 3) conventional paint. The solvent concentration and material parameters of all three systems are enlisted in Table 6.1.

Table 6.1. Material properties of various paint systems

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Nanopaint I</th>
<th>Nanopaint II</th>
<th>Conventional Paint</th>
</tr>
</thead>
<tbody>
<tr>
<td>Paint Density</td>
<td>1.286 g/cc</td>
<td>1.286 g/cc</td>
<td>1.23 g/cc</td>
</tr>
<tr>
<td>NP type</td>
<td>SiO₂ with 100 nm particle-size</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>NP mass fraction</td>
<td>10%</td>
<td>5%</td>
<td>0%</td>
</tr>
<tr>
<td>Viscosity</td>
<td>46 cP</td>
<td>43 cP</td>
<td>40 cP</td>
</tr>
<tr>
<td>Paint flow-rate</td>
<td>150 ml/min</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Size of Paint droplets</td>
<td>10-100 μm</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Solvent (Acetone) concentration (lb/gal)</td>
<td>4</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

For the ventilation air unit, the booth was provided with four 25-hp fan motors. The flow rate of the exhaust air generated by the fans was 80,000 CFM. Thus, the exhaust air velocity was calculated as,

\[ v = \frac{80,000 \text{ CFM}}{(2 \times 4 \text{ ft} \times 39 \text{ ft})} = 256.4 \text{ FPM (Feet Per Minute)} = 1.30 \text{ m/s} \quad (6.1) \]

where \((2 \times 4 \text{ ft} \times 39 \text{ ft})\) is the intake area for the exhaust air. The specified system information was used to construct the CFD-based models by employing Eq’s. (5.1) through (5.16).
spray booth consists of four 25-hp fan motors, and the electrical draw of each motor is 18.64 kW. Thus, the energy coefficient in Eq. 5.16 is calculated as:

\[
c = 4 \times 18.64 \text{ kW} / 1.30 \text{ m/s} = 57.35 \text{ kW/m/s}
\]  

(6.2)

The performance of the paint spray system, in terms of air purification by ventilation, quality of the deposited paint film and transfer efficiency depends on multiple factors such as ventilation air velocity, shaping air velocity, spray pattern, injection tracks, injection angle, surface roughness, gun-to-substrate distance of separation, etc. For nanopaint spray, toxic nanoparticles are released in the atmosphere which must be removed to keep the booth environment safe for workers. The quality of the film may alter due to the addition of nanoparticles in the paint matrix. The prediction of the coating film topology provides key information about the role of nanoparticles. For the process to be economical, transfer efficiency of the paint is a crucial factor. All the parameters are a function of the method of spraying and operational settings. The optimum performance can be obtained by adjusting the operational parameters appropriately for each type of coating material to be used. This works studies these crucial operational parameters which can improve the sustainability performance of the paint spray system for different types of coating materials.

6.2 Case Study Description

In this work, the paint material was applied on the panel surface with three different spray patterns. The spray patterns are described below as Cases 1 to 3. In each case, the separation between the gun and the substrate was fixed at 8”. The performance of each of the three paint systems was analyzed through CFD-based modeling and simulation to compare the film quality,
topological properties and emission of VOCs’ and NPs’. The spray patterns and resulting coverage of the sample by the paint material are shown in Figure 6.1.

Case 1: The paint was sprayed in a cone shape at the center of the sample at a spray angle of 66.02º. The spray angle was adjusted such that no paint could be released directly in the surrounding (See Figure 6.1 (a)).

Case 2: The paint was sprayed in a cone shape from the center at a spray angle of 71.67º. This spray angle ensured a complete coverage of the panel surface. However, a certain portion of paint was allowed to release directly in the surrounding as overspray (See Figure 6.1 (b)).

Case 3: The paint was sprayed at 5 different locations on the panel for 2 seconds at each at a fixed spray angle of 58º. The location of spray gun was changed to four corners and the center of the panel (See Figure 6.1 (c)) in order to improve the surface topology of the deposited film.
Figure 6.1. Spray patterns for (a) Case 1, (b) Case 2 and (c) Case 3.
In each case, the paint was sprayed for 10 sec. The fluid was injected along fixed 500 droplet streams from the outlet of the gun. It ensured even distribution of the paint material on the panel surface in each case. The streams injected droplets of different diameters ranging from 10μm to 100μm. The intermediate droplet size distribution was calculated by applying Rosin-Rammler distribution function with 50μm as the average size (Alderliesten, 2013). The spread parameter was set equal to 2. The Rosin-Rammler distribution function calculates the droplet sizes based on Eq. 6.2.

\[ Y_d = e^{(\frac{d}{\bar{d}})^n} \]  \hspace{1cm} (6.2)

where \( Y_d \) is the mass fraction of the paint material injected in the form of droplets with diameter \( d \), \( \bar{d} \) is the average diameter and \( n \) is the spread parameter. The plot of initial size distribution based on Rosin-Rammler equation is given in Figure 6.2.

![Initial droplet size distribution based on Rosin-Rammler function.](image)

The CFD model of the paint spray system considered certain assumptions for the analysis of each case. These assumptions are,
1. The nanoparticles were emitted in the atmosphere in the enclosed form inside a paint resin.

2. All the nanoparticles were spherical with identical size of 100 nm.

3. On collision of paint droplets on the substrate, the droplets showed one of the four behaviors: splash, spread, stick and rebound. In the case of splashing, each droplet would break to form 4 tiny fractions of same size and equal mass fractions.

4. The effect of gravitational force which may cause sagging of wet film was neglected.

5. The collision on walls of paint-booth lead to sticking of the paint material.

6.3 Simulation Details

The models described above were implemented using the ANSYS CFD software FLUENT, Version 14. A fine mesh was generated with 356746 total elements. The mesh resolution near the gun outlet was higher to accurately predict the flow dynamics and the interaction between spray particles and the air. The droplets were injected in a conical fashion from the spray gun. Unsteady particle tracking was applied for DPM particle treatment. The particles were sprayed at each particle time step of 0.1 second. Droplet collision and droplet break-up (WAVE) models were activated to simulate the paint flow behavior during the spray. Stochastic tracking was applied by discrete random walk model. Coupled scheme was implemented for pressure-velocity coupling. The pressure discretization was performed under the second order scheme.
The air flow model predicted the air flow profile inside the booth, which was generated as a result of interaction of downdraft air and shaping air. The contour of air flow velocity is shown in Figure 6.3.

![Air Velocity Contour](image)

Figure 6.3. Air velocity contour (a) front view, (b) side view.
6.4 Spray Trajectories Analysis

The paint droplets were injected from the gun nozzle at a high speed of 20 m/s. Due to such high speed, the continuous phase had minimal effects on the flow and transfer efficiency of the paint. For the DPM simulations, the boundary condition for the walls of the booth was selected such that the droplets would stick after the contact. The downdraft air was allowed to flow from the opening on the roof to the grids mounted at the floor of the booth. The panel surface was set as “wall-film”. The trajectories were calculated at every 0.1 sec of iteration time step. In all the cases, the droplets ejected from the gun were either deposited on the panel or released in the air.

The flow of paint droplets along the spray booth is represented in Figure 6.4. It can be seen from Figure 6.4a that the particles with highest residence time (red color) are lying on the wall of the booth. However, the particles with smallest residence time (blue color) are observed near the panel. Figure 6.4b represents the paint particles in the air at the end of 10 sec of spray and as a function of VOC content present in them. The coloring shows that the particles which are further away from the substrate and have spent longer time in the booth have lesser amount of VOC in them. This is expected due to higher evaporation of solvents at longer residence time.
Figure 6.4. DPM droplet tracks at the end of 10 sec of paint spray as a function of (a) residence time, and (b) VOC mass fraction.
6.5 VOC Emission Analysis

During the spray application, the solvent was continuously evaporating from the paint material. This VOC vapor contributes to the increasing air pollution inside the booth which could be hazardous to the worker standing inside. Using this CFD simulation for all the cases, the vaporized VOC concentration distribution was analyzed. Figure 6.5 represents this concentration of VOC. It can be seen that the vapor concentration is highest near the sample. This VOC vapor is taken away with the downdraft air through the grids at the bottom. The spray booth and ventilation system efficiency is determined by its ability to remove this VOC from the booth ambience and keep the environment clean for working.

The VOC emission was also analyzed more precisely by calculating the solvent concentration remaining inside 20 randomly selected droplet streams injected initially at a time zero. Figure 6.6 represents the trend of solvent concentration change in each of the 20 droplet streams of Case 1 for all three paint samples. It can be observed that the solvent concentration was exponentially decreasing along the time. This decrease was a result of continuous solvent evaporation contributing to increasing VOC in the booth. The occasional increase in the solvent concentration of few droplets could be caused due to coalescence of several droplets together after depositing on the panel film. Majority of the droplet streams did not reach 10 sec. It could also be a cause of coalescence of the droplets with others and spreading on the panel film. A close similarity among all three graphs (Figures 6.6a through 6.6c) suggests that addition of nanoparticles in the paint matrix did not affect the solvent concentration trends of all the droplet streams. Thus, nanoparticles had negligible effect on VOC evaporation during the paint spray.
Figure 6.5. Concentration of VOC vapor inside the spray booth after 10 sec of spray.

Figure 6.6. Solvent concentration change during the spray operation of Case 1 for (a) Nanopaint I, (b) Nanopaint II, and (c) Conventional paint (cont’d).
Figure 6.6. Solvent concentration change during the spray operation of Case 1 for (a) Nanopaint I, (b) Nanopaint II, and (c) Conventional paint.
The deposition of the solvent on the panel surface was also analyzed and compared to study the effect of nanoparticles addition into coating matrix. Figure 6.7 represents the contours of solvent concentration that was deposited on the panel surface after the spraying of Nanopaint I and conventional paint material through the spray pattern used in Case 1. The close proximity of the two contours suggests that nanoparticles have negligible effect on the solvent deposition on the spray panel surface.

![Figure 6.7. Contours of solvent deposition on panel surface by spray pattern of Case 1 for the spraying of (a) Nanopaint I, and (b) Conventional Paint.](image)

6.6 Paint Transfer Efficiency

The paint-spray process can be economical and the quality of deposited paint film can be improved with increasing the transfer efficiency. Although spray painting provides a good control over the quality of the finish, lower transfer efficiency may cause higher emission of
paint mists. The transfer efficiency of the spray process primarily depends upon the injection speed and the gun-to-substrate separation distance. The non-transferred paint contributes to the overspray which causes worker’s exposure to higher concentrations of paint mist.

The separation distance between gun and sample was fixed to 8” for all the cases (Beuerle, 1996). The transfer efficiencies of all the cases and types of paints are reported in the Table 6.2. As expected, Case 1 provided the highest transfer efficiency followed by Case 3 and Case 2. It can be observed that the nanopaint samples with increasing nanoparticle concentration have better transfer efficiencies. This could be due to the presence of nanoparticles and increase in the paint viscosity. The spray of more viscous paint deposited more content on the film and formed lesser paint mist.

Table 6.2. Analysis of performance parameters for all three cases of paint spray

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Cases</th>
<th>Geometry and spray pattern</th>
<th>Nanopaint I</th>
<th>Nanopaint II</th>
<th>Conventional Paint</th>
</tr>
</thead>
<tbody>
<tr>
<td>Transfer Efficiency (%)</td>
<td>1</td>
<td>Spray at the center (partial coverage of panel)</td>
<td>85.09</td>
<td>81.73</td>
<td>74.93</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>Spray at the center (full coverage of panel)</td>
<td>74.85</td>
<td>73.86</td>
<td>72.05</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>Spray at 5 locations (full coverage of panel)</td>
<td>80.91</td>
<td>76.46</td>
<td>72.26</td>
</tr>
<tr>
<td>Paint removed by the ventilation system (%)</td>
<td>1</td>
<td>Spray at the center (partial coverage of panel)</td>
<td>31.48</td>
<td>31.09</td>
<td>32.02</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>Spray at the center (full coverage of panel)</td>
<td>30.61</td>
<td>30.71</td>
<td>31.07</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>Spray at 5 locations (full coverage of panel)</td>
<td>31.12</td>
<td>31.08</td>
<td>32.45</td>
</tr>
<tr>
<td>NP’s concentration in booth air (no./m³)</td>
<td>1</td>
<td>Spray at the center (partial coverage of panel)</td>
<td>1.351x10¹²</td>
<td>8.323x10¹¹</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>Spray at the center (full coverage of panel)</td>
<td>2.307x10¹²</td>
<td>1.197x10¹²</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>Spray at 5 locations (full coverage of panel)</td>
<td>1.742x10¹²</td>
<td>1.074x10¹²</td>
<td>0</td>
</tr>
</tbody>
</table>
The overspray, consisting of paint solids, nanoparticles and VOC, results in the worker's exposure to toxic substances. This overspray must be removed efficiently by the downdraft air in order to keep the booth environment safe for working. For all the simulated cases, the mass flow and velocity of downdraft air was kept unchanged. The efficiency of the ventilation system for different cases of paint systems was studied by calculating the proportion of paint removed by the downdraft air out of total overspray. The corresponding values are reported in the Table 2. It confers that the efficiency of the ventilation system, in case of conventional paint, is slightly better than that of nanopaint. However, it is preferred to have a high efficiency of ventilation system, especially in the presence of toxic nanoparticles. Thus, there is a trade-off between the transfer efficiency and environmental emissions.

6.7 Nanoparticles Emission

The nanoparticles emission during each case was analyzed by calculating the total concentration of nanoparticles in the booth environment at the end of paint-spray operation. The values obtained from the analysis are reported in Table 6.2. It can be seen that the higher nanoparticles content in the paint system invites further risk of nanoparticles emission and exposure to higher concentrations. Highest concentration of nanoparticles was emitted during the spray of nanopaint samples through Case 2. With the environmental regulations on NP’s emission becoming more strict day-by-day, it is crucial to have an optimized ventilation system for nanopaint spray to minimize this exposure which can cause several health concerns.
6.8 Film Topology Analysis

The key benefit of paint-spray application method is the ability to control and improve surface morphology and coating film topology. Spreading of the paint material on the film and the prediction of velocity profiles of paint droplets are the fundamental factors required for the accurate determination of surface morphology. Fogliati et al. (2006) determined that the Realizable $k$-$\varepsilon$ model gives most accurate prediction of these characteristics. Thus, the same model was adopted for the simulation of all the cases.

In order to determine the film thickness profile of the deposited paint layer, the sample surface was modeled as a “wall-film” which predicts four different phenomena of spread, stick, rebound and splash of the impacting paint droplets depending on the impact angle and velocity. The paint mass and deposited film thickness values at various locations on the panel were recorded. Using these values the plots of distribution of paint thicknesses over the panel surface were reconstructed and analyzed. The plots obtained for the spray of Nanopaint I using all three cases of spray patterns are represented in Figure 6.8. The roughness of the film surface was calculated as the “arithmetic mean roughness ($R_{\text{mean}}$)” by using the Eq. 10.

$$R_{\text{mean}} = \frac{1}{n} \sum_{i=0}^{n} |y_i|$$

(6.3)

where $y_i$ is the vertical distance from the mean thickness line to the $i^{th}$ data point and $n$ is the number of selected ordered and equally spaced data points along the surface of the panel. The values of the average thickness and roughness of the deposited film from all the cases are reported in Table 6.3. In order to ensure the reliability of the simulations and the accuracy of the thickness data, these average thickness values were compared with the theoretically calculated thickness values. The theoretical thickness is the ratio of quantity of paint that is transferred on
the substrate over the cross sectional area of the substrate. The final values were calculated for each case and the percent error is reported in Table 6.4. For all the cases, the error was less than 8% which verified the reliability of the simulation and the final results.

Table 6.3. Average film thickness and roughness values for all the cases of all the paint samples

<table>
<thead>
<tr>
<th>Cases</th>
<th>Average film thickness (μm)</th>
<th>Arithmetic mean Roughness (μm)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Nanopaint I</td>
<td>Nanopaint II</td>
</tr>
<tr>
<td>1</td>
<td>29.70</td>
<td>28.54</td>
</tr>
<tr>
<td>2</td>
<td>25.55</td>
<td>24.55</td>
</tr>
<tr>
<td>3</td>
<td>27.31</td>
<td>26.17</td>
</tr>
</tbody>
</table>

Table 6.4. Theoretical thickness values and percent error in the calculations for all the simulation cases

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Cases</th>
<th>Geometry and spray pattern</th>
<th>Nanopaint I</th>
<th>Nanopaint II</th>
<th>Conventional Paint</th>
</tr>
</thead>
<tbody>
<tr>
<td>Transfer Efficiency (%)</td>
<td>1</td>
<td>Spray at the center (partial coverage of panel)</td>
<td>85.09</td>
<td>81.73</td>
<td>74.93</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>Spray at the center (full coverage of panel)</td>
<td>74.85</td>
<td>73.86</td>
<td>72.05</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>Spray at 5 locations (full coverage of panel)</td>
<td>80.91</td>
<td>76.46</td>
<td>72.26</td>
</tr>
<tr>
<td>Theoretical Thickness (μm)</td>
<td>1</td>
<td>Spray at the center (partial coverage of panel)</td>
<td>29.68</td>
<td>28.50</td>
<td>27.32</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>Spray at the center (full coverage of panel)</td>
<td>26.11</td>
<td>25.76</td>
<td>26.27</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>Spray at 5 locations (full coverage of panel)</td>
<td>28.22</td>
<td>26.67</td>
<td>26.35</td>
</tr>
<tr>
<td>% error</td>
<td>1</td>
<td>Spray at the center (partial coverage of panel)</td>
<td>0.08</td>
<td>0.14</td>
<td>3.24</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>Spray at the center (full coverage of panel)</td>
<td>2.11</td>
<td>4.71</td>
<td>7.58</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>Spray at 5 locations (full coverage of panel)</td>
<td>4.64</td>
<td>3.36</td>
<td>7.78</td>
</tr>
</tbody>
</table>
Figure 6.8. Film thickness profiles of Nanopaint 1 from (a) Case 1, (b) Case 2 and (c) Case 3 (cont’d).
Figure 6.8. Film thickness profiles of Nanopaint 1 from (a) Case 1, (b) Case 2 and (c) Case 3.

From Figure 6.8, it can be observed that the film topology from Case 3 was substantially improved as compared to that from Case 1 and Case 2. The thickness values from Table 6.3 suggest that the average thicknesses of nanopaint samples were greater than the conventional paint. Case 3 showed about 7% improvement of the average film thickness than the worst case. The roughness was substantially reduced in Case 3 as compared to Case 1. Thus, the spraying of paint uniformly over the cross-section of the panel showed significant improvement in the film topology.
CHAPTER 7
CONCLUSIONS AND FUTURE WORK

This chapter concludes the dissertation report with the summary of all the research findings in the areas of multiscale molecular modeling, life-cycle based sustainability assessment of nanocoating technology and CFD modeling and simulation of paint application process for the prediction of nanopaint performance and environmental emissions as compared to existing conventional paint systems. The summary is followed by recommendations for possible extension of this work in the future.

7.1 Conclusions

A comprehensive methodology and framework for the multiscale life-cycle-based sustainability assessment of nanocoating technology has been successfully developed. This novel technology was studied using computational modeling and experimental analysis on various levels of time and length ranging from nano- to macro- scales. These modeling methodologies and corresponding results generated key information required for the quantification of several sustainability matrixes representing different stages of the life cycle of nanocoating technology.

The key challenge to the research on sustainability assessment of an emerging technology such as “nanocoating technology” comes from the limited scientific knowledge and the reliable
data availability. Since last decade, nanoparticle-based coating materials are being studied extensively to determine all the attractive properties achievable with it. It has been proven that the addition of nanosize fillers of different types and shapes can enhance the polymer composite as well as coating resin’s properties tremendously. The final product can have significantly improved mechanical stiffness, scratch resistance, super-corrosion resistance, antifouling properties, mar resistance and also certain smart functionalities such as self-cleaning, self-healing, thermochromic, heat resistance, significantly improved solvent and chemical resistance, etc. However, it is still unknown how these nanoparticles are responsible for such property enhancements. The experiments solely are not capable of answering questions and studying the microstructure-property correlations among nanoparticles and polymer resins. Through this research, an integrated computation-experimental approach was provided to study these microstructure-property-performance correlations.

The computational modeling was performed using molecular simulation techniques such as Monte-Carlo and Molecular Dynamics (MD). In the past, modeling methodology was developed on macro- to meso- scales of material. This work adopted the bottom-up, deductive systems engineering approach of materials development. The nanocoating system was simulated on nano-, micro- and meso- levels and the methodology was connected with the previously developed system to complete the multiscale modeling methodology of nanocoating system development.

The nanocoating material was first simulated through MD simulation technique on a microscale. The microstructural parameters such as bond lengths, bond angles, atomic charges, dihedrals, etc. were taken from a well-established atomistic force field called “CHARMM”. The atomistic system consisted of a large number of entities which made the simulation of a
nanocoating system with huge design space and complexity computationally very expensive. Thus, the atomistic design was mapped onto a coarse grained system of “beads”. This was accomplished by following MARTINI force field development protocols. The bonded and non-bonded interaction parameters for the system of PMMA-based resin material and TiO$_2$ type nanoparticles were developed and the system was studied extensively to analyze the structure-property correlations which cause the enhancement of properties in case of nanoparticles dispersion. The factors such as, nanoparticles size, volume fraction, polydispersity index, etc. were studied to determine their effects on final nanocomposite structure. This work was later coupled with experimental analysis in order to accomplish three key objectives: (a) to provide validation to the computational modeling work, (b) to study the effect of nanoparticles distribution inside a coating film on the final mechanical properties, and (c) to lead the computational-experimental approach towards a new knowledge discovery and optimization of a “self-cleaning” smart functionality.

It was realized that the optimization of nanocoating formulations solely cannot bring these emerging products into commercialization due to various types of environmental hazards, health concerns and economic issues associated with it. Thus, a comprehensive sustainability assessment framework was developed to look into all the aspects of nanocoating technology, not only at the manufacturing stage, but over its entire life-cycle which includes all the phases from material selection and preprocessing, paint manufacturing, paint application and film formation to its end use and disposal. The developed methodology demonstrated how the sustainability indicators at various stages of life are correlated with each other. This technology can gain significant value and the nanoproducts can be commercialized only if its sustainability
performance is assessed, compared with existing technologies and further improved to make it a safer alternative.

However, the challenge to such detailed assessment emerged from the availability of data for the nanopaint. Thus, Computational Fluid Dynamics (CFD) approach was adopted to simulate these nanopaints and study the coating quality parameters and environmental emission factors during application phase. In automotive industries, paint-spray is the most common application method used for coating the auto bodies. In this research, an industrial paint booth was designed and the spray of various paint systems was studied. The components which may primarily affect the environment during paint spray are VOC (Volatile Organic Contents) and nanoparticles. Through CFD simulation tool, ANSYS FLUENT, the paint spray process was simulated and the emission of VOC’s and nanoparticles was thoroughly assessed. This simulation generated crucial information related to the concentrations of overspray paint and VOC’s that was released in the atmosphere and could cause health hazards to the workers inside the spray booth and pollute the drainage water passing through these paint booths. The final coating quality parameters were also studied to ensure the superior performance of nanopaints over conventional paints. The quality parameters such as, film topology, roughness, paint spray efficiency, etc. were determined and the results for different types of conventional and nanopaints were compared.

The sustainability assessment framework and multiscale computational modeling methodologies generated in this research can provide a key pathway towards optimizing an emerging nanocoating technology and justify the inclusion of nanopaint products into future commercial market. This research can also assist experimentalists in learning the structure-
property correlations of nanocomposite coatings and discover novel formulations of these materials for the future.

### 7.2 Future Work

The findings presented in this dissertation provide a solid foundation for the research related to the development of sustainable nanocoating technology. It can be extended in three major directions: multiscale molecular modeling of nanocoatings, life-cycle based sustainability assessment and modeling and simulation of nanopaint application process.

**Multiscale molecular modeling of nanocoatings.** This dissertation focused on the development of the multiscale molecular modeling and simulation of nanopaints for the study of structure-property correlations. The force field parameters and the CG methodology developed in this research was applied for the study of polymer nanocomposite microstructure. This sophisticated methodology and the MARTINI force field for PMMA/TiO$_2$ system can be applied for the study of several other key properties. However, it is important to validate all the model parameters and the property predictions presented in this dissertation through experiments. The experimental efforts initiated in this research for the study of polymer nanocomposite films and determination of properties related to mechanical stiffness and scratch resistance can be further explored and the validation to the multiscale framework (Figure 1.3) can be provided. The accomplished work and some of the directions for future work using multiscale MD simulation method is shown in Figure 7.1.
With the accomplished work on the multiscale modeling of thermoplastic nanocomposite coating, the properties such as mechanical stiffness, scratch resistance, barrier properties and self-cleaning can be predicted. However, a majority of the automotive paint are 2-component, thermoset systems. The existing methodology can be extended to simulate such thermoset system by developing appropriate force field parameters for the crosslinker molecules and verifying them through experiments and empirical data. If succeeded, this model can also provide validation to the past work which used Monte-Carlo based simulation method to model generic thermoset nanocomposite coating material.

The CG model for nanocoatings with TiO₂ type of nanoparticles can be applied for the study of “smart” coating materials with self-cleaning functionality. The hydrophilicity of the nanoparticles play a key role in determining the performance of the eventual nanopaint. This
MARTINI force field based CG model can predict the interactions among paint resin and surface of nanoparticles with different hydrophilicity. This phenomenon can assist in the development of optimum formulations of nanopaints which can deliver self-cleaning functionality.

The thermoset paint system is a 2K (2 component) system where the 2\text{nd} component is a cross-linker/curing agent. The curing agent is mixed with the paint vehicle before the mixture is applied onto car bodies. The cross linking reaction helps in the formation of very hard films which have significantly better mechanical properties of the film as compared to thermoplastic paints. The thermosets have been well rooted into automotive paint systems and most of the existing basecoat and topcoat formulations are thermoset paints. However, the development of nanoparticles-induced thermoset paints remains a mountain top area of research. The optimization of thermoset nanopaint formulations is a challenge due to the limited availability of knowledge about its microstructure. The CG model developed in this research can be further extended for the study of such systems and the microstructure-property-performance correlations can be established for thermoset nanopaint materials. Through the MD tool NAMD, it can be accomplished by developing an optimized CG force field parameters for crosslinker molecule and then simulate the cured system (as shown in Figure 7.1). However, NAMD does not support the formation of new chemical bonds and thus the curing reaction to happen during the course of simulation. This shortcoming of NAMD can be avoided by the use of the MD tool which supports such reactions. LAMMPS and GROMACS are the best alternate routes which can perform such reactions during simulation and thus the thermoset nanocoatings can be modeled.

The performance of nanopaints for automotive bodies heavily depends on the size distribution and shapes of dispersed nanoparticles. The nanoparticles such as TiO$_2$, SiO$_2$, alumina, MMT clay, etc. have different shapes (e.g. spherical, oval, platelet, crystalline, etc.) and
thus they exhibit different characteristics in the final coating material. The multiscale modeling methodology developed in this dissertation can be applied to study the effects of such shapes of nanoparticles on the final performance of the coating film.

One of the challenges that the nanopaints are facing for becoming commercialized is from the issues related to durability. This factor is also related to destabilization of paint due to aggregation/agglomeration of nanoparticles. This crucial phenomenon can be studied by modifying the developed PNC system with a system where each nanoparticle is represented by a single Van der Waal bead. The interaction parameters can be determined by following the same MARTINI methodology which was applied in this research. Such system can efficiently address the concerns related to aggregation/agglomeration of different types of nanoparticles in the paint formulation.

**Life-cycle-based sustainability assessment.** It has been evidenced by several studies that the nanoparticles pose serious threat to environment and human health. Needless to say, that the nanocoating, after commercialization in the future, can also pose such threats to human life and environment. Thus, this dissertation was focused on the development of a comprehensive life-cycle based sustainability assessment framework (Figure 4.2) to analyze the performance of existing nanocoating technology and determine the areas to improve in terms of environmental, economic and social performance of this technology for making it sustainable. This framework can be further refined with much extensive research on each of the stages of life cycle by obtaining more reliable and accurate data for all the sustainability matrix parameters. The accuracy of the data is the most important aspect which can justify the credibility of this methodology for the prediction of sustainability performance of different types of nanopaint systems. The indicator matrixes developed in this work are also generic. More indicators can be
explored and this framework can be further improved to make it applicable for a specific
category of nanopaint. For example, the assessment of a nanopaint system which has application
in automotive industry can be more effective if the indicators are more focused towards the
factors such as, durability of the film, energy consumption during application of the films in
paint shops, and final performance of cured coating surfaces.

Another aspect which can be introduced is the “decision-making”. The current LCSA
framework does not include the details of decision-making based on the assessment results. The
decision making methodology can be combined with the assessment to provide
recommendations for improving the sustainability performance of the nanocoating technology.

**CFD modeling and simulation.** The part of the research was focused on the study of
automotive paint application process through CFD-based modeling to assess the environmental
emissions and coating quality parameters during paint-spray. The energy efficiency and VOC
emission patterns were studied through CFD modeling for the conventional paint spray (Li et al.,
2013). In case of nanopaint spray, there are added concerns related to emission of nanoparticles
along with VOC’s during the spray application. The presence of nanoparticles can also affect the
performance of spray technique and thus the quality of the final film. These factors were studied
through CFD based modeling of a manual paint-spray booth. In future, this approach can be
extended to study automatic paint spray booths which consist of several robotic spray bells to
ccoat the car bodies passing through the booth zones.

The energy consumption and heating of wet films in curing ovens are other key factors
which can influence the overall sustainability performance of the nanocoating technology. The
CFD modeling can be used for the assessment of curing zones to determine the emissions and
effects of the operational settings on the quality of the final film. Such models can greatly help
in obtaining the data for the sustainability assessment of application stage of the life-cycle of nanocoating technology.
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ABSTRACT

MULTISCALE DESIGN AND LIFE-CYCLE BASED SUSTAINABILITY ASSESSMENT OF POLYMER NANOCOMPOSITE COATINGS

by

ROHAN G. UTTARWAR

December 2013

Advisor: Dr. Yinlun Huang
Major: Chemical Engineering
Degree: Doctor of Philosophy

Nanopaint with significantly improved performance properties could have numerous applications in the automotive, aerospace, ship-making, chemical, electronics, steel, construction, and other industries. It is predicted that nanopaint, especially that providing multiple new functionalities to nanocoatings will be dominant surface coating material in the near future. However, how to ensure that such a nanocoating material be sustainable is a very sophisticated scientific problem. The challenges in research arrive from the insufficient knowledge about formulation-structure-property correlations of nanocoating materials and their design complexity, and more seriously the prediction of their life-cycle sustainability performance, especially environmental impact and health risk.

This research focuses on the study of multiscale sustainable nanocoating design, which could have applications from novel function envisioning and idea refinement, to knowledge discovery and design solution derivation, and further to sustainability performance evaluation. The nanocoating design is studied through developing multiscale models and conducting simulations using advanced computational techniques. The nanocoating’s sustainability assessment was performed throughout its life cycle. Computational simulation aims at
integrating top-down, goals/means, inductive systems engineering and bottom-up, cause and effect, deductive systems engineering approaches for material development. The *in silico* paint resin system is a water-dispersible acrylic polymer with hydrophilic nanoparticles incorporated into it. The nano-scale atomistic and micro-scale coarse-grained (CG) level simulations are performed using molecular dynamics methodology to study structural and morphological features, such as effect of polymer molecular weight, polydispersity, rheology, nanoparticle volume fraction, size, shape and chemical nature on the bulk mechanical and self-cleaning properties of the coating film. At the macro-scale, a paint spray system commonly used for automotive coating application is studied using CFD simulation technique to generate crucial information about the effects of nanocoating formulation on environmental emission and coating film quality. The life-cycle based sustainability assessment study addresses all the critical issues related to economic benefit, environmental implication and societal effect of nanocoating technology through case studies on automotive coating systems. It is accomplished by identifying crucial correlations among measurable parameters at different stages and developing sustainability matrices for analysis of each stage of life-cycle. The findings from the research can have great potential to derive useful conclusions in favor of future development of coating systems with novel functionalities and improved sustainability.
AUTOBIOGRAPHICAL STATEMENT

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• NSF Travel Award for ICOSSE ’13 - 3rd International Congress on Sustainability Science & Engineering Conference, Cincinnati, OH, 2013
• Membership, American Institute of Chemical Engineers (AIChE), 2010 - present.
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• Co-Chair, Ninth Annual Graduate Research Symposium, Wayne State University, Detroit, MI, October 27, 2009

PUBLICATIONS AND TECHNICAL PRESENTATIONS