Recycled Carpet Materials for Infrastructure Applications

Ranji Vaidyanathan, Ph.D., P.E.
Raman P. Singh, Ph.D.
Tyler Ley, Ph.D., P.E.
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Recycled carpet materials for infrastructure applications

The objective of this project was to develop novel composite materials for infrastructure applications by recycling nylon based waste carpets. These novel composites have been proven to possess improved mechanical and sound barrier properties to meet the needs of infrastructure in the transportation industry as sound barrier walls at the highway turnpike. This was achieved by scaling up the Vacuum Assisted Resin Transfer Molding (VARTM) technique of manufacturing carpet composites. Additionally, these composites were also tested and characterized for long-term stable mechanical properties through the introduction of nano-sized fillers for improved resistance to moisture and UV.

This project has proven the concept of innovative-engineered composite materials with tailored mechanical and physical properties from discarded waste carpet reducing its impact on the environment. In particular, this project will take advantage of the 3-D fiber architecture, aspect ratio of the carpet fibers and the layered nature of the discarded carpet to provide a unique combination of impact, mechanical and acoustic insulation properties that could be advantageous to the infrastructure and transportation industries.
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ACKNOWLEDGMENTS

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Recycled Carpet Materials for Infrastructure Applications

Ranji Vaidyanathan, Ph. D, P.E.
School of Materials Science and Engineering

Raman P. Singh, Ph. D
School of Mechanical and Aerospace Engineering

Tyler Ley, Ph. D, P.E.
School of Civil Engineering

Oklahoma State University
Stillwater, OK 74078

FINAL REPORT

Project OTCREOS9.1-49

June 2013

Oklahoma Transportation Center
2601 Liberty Parkway Suite 110
Midwest City, OK 73110
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EXECUTIVE SUMMARY

The objective of this project was to develop novel composite materials for infrastructure applications by recycling nylon based waste carpets. These novel composites have been proven to possess improved mechanical and sound barrier properties to meet the needs of infrastructure in the transportation industry as sound barrier walls at the highway turnpike. This was achieved by scaling up the Vacuum Assisted Resin Transfer Molding (VARTM) technique of manufacturing carpet composites. Additionally, these composites were also tested and characterized for long-term stable mechanical properties through the introduction of nano-sized fillers for improved resistance to moisture and UV.

This project has proven the concept of innovative-engineered composite materials with tailored mechanical and physical properties from discarded waste carpet reducing its impact on the environment. In particular, this project will take advantage of the 3-D fiber architecture, aspect ratio of the carpet fibers and the layered nature of the discarded carpet to provide a unique combination of impact, mechanical and acoustic insulation properties that could be advantageous to the infrastructure and transportation industries.

Composites were fabricated from recycled carpet materials and exposed to a variety of environmental conditions including moisture, ultra violet radiation and a combination of the two. Significant reduction in flammability in the carpet composites was observed due to use of clay. The linear burning rate as per ASTM D 635-10 was reduced by 20 % for composite samples coated with the epoxy-clay coating, while it was reduced by 34 % for composite samples infused with 4 % nanoclay.

Moisture exposure for these samples indicated that both clay coating and clay infusion reduced the deleterious effect of moisture on the composites. Significant reduction in diffusivity was found in clay coated and clay infused composites. All composites showed a reduction in flexural strength and modulus, where severity of degradation was less in clay coated and clay infused carpet composites. For alternating moisture and UV exposure cycles, the clay-coated samples exhibited the lowest change in mass, while the neat epoxy recycled carpet composites showed the highest mass gain. However, It was also observed that the combined moisture/UV exposure resulted in an increase in strength in the case of all composite types. Finally, wind load tests proved that the composite is strong and withstands extreme wind conditions similar to tornadoes.
Three graduate students have completed their Masters work on this funded project. A patent based on this work was applied for during the project and has recently been issued – A. K. Singh, G. Pandey, R. Vaidyanathan, and R. P. Singh, “Fabrication Method of Multi-functional Composites from Pre/Post-Customer Carpet,” U.S. Patent Application No. 12/695730, August 5, 2010. An Oklahoma manufacturer (K. T. Plastics, Calera, OK) has signed an option to license for this patent. It is expected that KT Plastics will be providing the technology to another Oklahoma manufacturer of highway sound barriers (Apache Precast, Jenks, OK).

Several papers have been published and presented, while also giving the opportunity to intern for undergraduate students at Oklahoma State University.
1. INTRODUCTION

This project has proven the concept of innovative-engineered composite materials with tailored mechanical and physical properties from discarded waste carpet reducing its impact on the environment. In particular, this project will take advantage of the 3-D fiber architecture, aspect ratio of the carpet fibers and the layered nature of the discarded carpet to provide a unique combination of impact, mechanical and acoustic insulation properties that could be advantageous to the infrastructure and transportation industries.

The world’s primary energy needs have increased by 66 percent in the last 30 years and today’s energy path would mean rapidly increasing dependence on fossil fuels with alarming consequences for climate change and energy security [1]. Experts accurately predicted large and permanent increases in oil prices after the year 2000 due to increasing scarcity. A series of episodes including the oil reaching a record price of $120 a barrel in March 2008 has made this evident. The president of OPEC (Organization of Petroleum Exporting Countries) has warned of oil reaching $200 a barrel clearly indicating huge demand for oil. The rapid economic development of developing countries around the world (including India and China) has produced insatiable demand for oil and other energy sources [2]. However, apart from its demand-supply problem, there are even larger challenges in the energy industry, such as consumption of energy yielding to increase in greenhouse gas emissions that in turn results in increase of global temperature. Hence energy is crucial for sustainable development and World Energy Outlook 2009 report suggests that there is an urgent need to limit the global temperature rise to $2^\circ$C [1].

For sustainable development, the materials recovery and recycling industry has to deliver marketable products which are based on cost effective technologies and have a positive life cycle environmental impact [3]. This served as a motivation to address a solution to the problem of recycling of waste carpet lying in the landfills. Land-filling is not an environmentally friendly solution because carpet fibers are not biodegradable [4]. In the United States, environmental concerns and governmental regulations have started putting efforts in the direction of recycling all synthetic polymers, of which carpet and carpet fibers constitute a significant percentage [4]. Carpet is a petroleum-based product and it is estimated that carpet recycling can save more than 700,000 barrels of oil per year, conserving 4.4 trillion BTUs of energy [5].

In the United States alone, about 200 million tons of carpet waste is generated every year, and only 2% of this amount is recycled [6]. Out of the 5 billion pounds of carpet sent to landfill
only 1% of old carpet has been recovered as per Carpet America Recovery Effort (CARE), an organization committed to carpet recycling in USA [6]. In every municipal solid-waste stream of USA, post-consumer carpets account for 1 wt% of the stream thereby indicating the high usage of carpet. Carpet offers compelling benefits like underfoot comfort, sound absorption, improved slip resistance and thermal comfort and has therefore been the reason for covering 60 percent of all floors in the United States [7]. The Carpet and Rug Institute sustainability report says that “carpet industry in USA produces more than 19 billion square feet of carpet each year” [7].

Carpet waste is both an environmental and economical problem because of the cost involved in disposal of waste carpet. The cost also involves discarding of valuable raw material in the form of high engineering value fibers like Nylon 6.6, polyester, polyvinyl chloride (PVC), polypropylene (PP), other olefins etc. going to landfills apart from the cost of transporting the waste to landfills. Thus post-consumer carpet represents a significant environment burden on the landfills, tax burden due to state sponsored subsidies, and a lost economic opportunity. Hence it is imperative to recycle post-consumer carpets for a sustainable society.

The chart in Fig. 1.1 has been adapted from the data provided by the Environment protection agency on Municipal Solid Waste (MSW). Waste carpet is an important constituent of the MSW and the chart below gives an idea about the steady trend in recovery of carpets over the decade. The rapid economic development throughout the world has increased the generation of municipal solid waste. The imbalance between the amount of discarded waste and its minimal recovery indicates a deplorable outlook for the recycling industry. However the steady increase in awareness for recovery of municipal solid waste gives a hope for the waste recycling industry.

In 2009, CARE reported that 311 million pounds of post-consumer carpet were diverted from landfills, of which 246 million pounds were diverted due to a direct result of reuse and recycling [9]. This indicates that there is an increasing trend in awareness of carpet recycling and the products that fall under the topic of recycled carpets would be well received.
The 2010 Annual report by CARE pointed out that Nylon 6 and Nylon 6,6 represent 80 percent of all carpet diverted from landfills in 2009 [9]. The two predominant forces driving the waste nylon based carpet’s reclamation are – 1) the opportunities for usage of recycled nylon fiber in various end markets and 2) the huge market share of nylon carpet sold in United States [9]. The carpets made of other face fiber materials like polyester, PVC and PP are therefore not economical to recycle with current technology because these face fiber materials are cheaper virgin materials and to recycle them is uneconomical [11]. Among the carpets sold in the United States, about 65 percent of them are nylon based because nylon is a more durable fiber and has excellent performance characteristics [12]. There has been extensive research on recycle of nylon-based carpets due to the huge market for nylon fibers. Nylon 6 fibers are derived from caprolactum, which has a significant market value and are thereby recycled by chemical recycling methods. However the chemical recycling techniques for carpet fail to deliver in sustainable product development due to their characteristic requirement of high-energy inputs.
Literature suggests that the energy consumed in manufacture of nylon fibers for polymerization, spinning and finishing is between 369 and 432 MJ/kg [13]. The calorific value of crude oil is between 38 and 46 MJ/kg and therefore it can be concluded that for 1 kg of fiber about 11 kg of crude oil is necessary [13]. Though thermal utilization is a lucrative option, the process of thermal utilization uses only the calorific value which is about the same or slightly lower compared to crude oil [13]. Hence it is imperative to evaluate a process to save the 11 kg of crude oil getting wasted for every 1 kg of nylon wasted in the carpet waste. Therefore carpet recycling becomes critical and to ensure sustainable development.

**TABLE 1.1. NATIONAL DISCARD ESTIMATES PROVIDED BY THE CARPET AND RUG INSTITUTE [10]**

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<tr>
<td>Landfill Diversion Rate</td>
<td>3.8%</td>
<td>10%</td>
<td>19%</td>
<td>23%</td>
<td>27-34%</td>
</tr>
</tbody>
</table>

*Note* - (1) Cement kilns use incineration of carpets as a fuel supplement in kilns (2) NA - Not available.

At Oklahoma State University (OSU), an innovative carpet recycling technology has been developed to recycle waste carpets from landfills into structural composites using VARTM. The main advantage of VARTM is that the carpets are recycled without any physical or chemical
treatment which therefore results in process optimization and cost reduction. Fig. 1.2 includes a schematic of the VARTM process used to recycle carpet in this study.

**Figure 1.2.** An illustrative diagram of VARTM process to make carpet composites.

The vitality of a globalized economy is directly linked to the productivity and efficiency of its surface transportation system among other factors. For instance, the development of Interstate Highway System has stimulated the economy of the United States, but it also brings in increasing demands for safe, efficient and environmentally sound efficient transportation system with lower noise decibel levels. This indicates the feasibility for novel materials with better acoustic properties as highway noise barriers. In this study it has been proposed to use novel composite materials like carpet composites as sound barrier walls at turnpike.
2. BACKGROUND AND LITERATURE REVIEW

2.1 CARPET – A BRIEF REVIEW

2.1.1 Carpet Construction

A typical carpet has four main layers – face fibers, primary backing, binder and secondary backing. The arrangement of these four layers is as shown in Fig. 2.1.

![Figure 2.1 Carpet (cut-pile type) and its constituents – fibers, backing and binder [4].](image)

The top layer consists of face fibers that are generally tufted through the primary backing, usually made of polypropylene (PP). Other fibers like polyesters, polyethylene and rayon are also used as backing materials [4]. Latex adhesive, usually made of a styrene butadiene rubber co-polymer (SBR) adhesive is applied under the primary backing to secure face fiber [4]. The SBR adhesive combined with inorganic filler materials like calcium carbonate or barium sulfate is then used as a binder to bond the secondary backing to the primary backing [4]. The secondary backing is usually made up of same material as material as primary backing [4]. The face fibers contain dyes, soil repellents, fire retardants, and other additives to improve the carpet quality [4].

The following table, Table 2.1 gives a detail account of the constituents of a typical carpet with PP/SBR construction, which accounts for 95 percent of all the residential and commercial carpets in the United States. Carpet recycling is a complicated issue mainly due to the multi-component construction, presence of backing material and presence of dyes and coatings.
TABLE 2.1. COMPOSITION OF A TYPICAL CARPET WITH PP/SBR CONSTRUCTION [4]

<table>
<thead>
<tr>
<th>Sr. No.</th>
<th>Constituent</th>
<th>Composition (in oz. avg. wt.)</th>
<th>Composition (in percentage wt.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Face Fiber (Nylon 6 / Nylon 6,6)</td>
<td>30</td>
<td>45.80</td>
</tr>
<tr>
<td>2</td>
<td>PP (Primary backing)</td>
<td>4</td>
<td>6.11</td>
</tr>
<tr>
<td>3</td>
<td>Latex Adhesive</td>
<td>29 (SBR – 6 oz., CaCO₃ – 23 oz.)</td>
<td>44.27</td>
</tr>
<tr>
<td>4</td>
<td>PP (Secondary backing)</td>
<td>2.5</td>
<td>3.82</td>
</tr>
<tr>
<td>5</td>
<td>TOTAL</td>
<td>65.5</td>
<td>100</td>
</tr>
</tbody>
</table>

Figure 2.2 Illustration of the 3-D architecture of carpet [14].

Carpet padding is a cushioned material placed underneath carpet for longevity, comfort and noise absorption. It is mostly made with felt rubber and urethane. The cushion backed carpet incorporates the padding as an integral part of backing while incorporating coal fly ash into polyurethane [14].

2.1.2 Carpet Classification

Carpets can be classified in three ways – (1) based on its consumption, (2) based on its construction and (3) based on the type of face fiber used.

Based on its consumption, carpets can be classified as virgin carpets, pre - consumer carpet waste and post - consumer carpet waste. A virgin carpet is an unused carpet and it is
typically a carpet without any history of foot traffic. Virgin carpets are free of dirt, but they mainly consist of fibers, backing and a variety of chemical adhesives, fire retardants, dirt resistant, stain resistant dyes, color dyes and other materials. Pre-consumer carpet waste consists of the waste generated at the landfills largely due to the manufacturing process of carpet and the carpet fitting process. In the carpet manufacturing process, the edges of carpet need to be trimmed. These edge trims and off-cuts are disposed as pre-consumer carpet waste. The amount of carpet waste in the form of trimmings and waste is about 12% of the total carpet production [4]. In the automotive industry, a process called fitting process is used, where the carpets are cut into various irregular shapes and sizes. These cut edge trims and the waste carpet contribute to pre-consumer carpet waste in the landfill. Post-consumer carpet waste is defined as, “A used carpet generated at the landfills due to disposal of old carpets from residential and industrial areas.” The average age of a carpet is 8 to 10 years. The weight of post-consumer carpet waste is 20 percent more than virgin carpet due to presence of dirt, dog hair, nails, metal fragments, food, bacteria and other microorganisms etc.

In this study the focus has been on the recycle of post consumer carpet. The recycling process suggested could also be applied to pre-consumer carpet but it would need further improvement in process modifications of the established scale-up process which is not desired. Moreover the amount of pre-consumer carpet waste is low in comparison to post consumer carpet waste. Virgin carpets are not a part of the research to develop carpet composites in this study because this study aims at recycling the waste carpets in the landfills into highway noise barriers and it would defeat the objective of the study if virgin carpets are included in recycling of carpet composites.

Based on its construction, carpets can be classified as (1) cut-pile, (2) loop pile, (3) patterned loop and (4) cut & loop styles. These types of carpets can be manufactured by either weaving, tufting or by needle felt type of manufacturing process. Woven produces the highest grade of carpet in a loom by a process that is similar to woven cloth (cut pile type) [15]. Tufting is a widely followed industrialized process which involves sewing strands of yarn into the backing material and thereby creating thousands of yarn loops. In this study cut pile carpets were used for fabrication of carpet composites.
Based on the type of face-fiber material used, carpets can be classified into - Nylon (made of Nylon 6 or Nylon 6,6), Polyester, PP, Polyethylene Terephthalate (PET) and poly (vinyl chloride) (PVC), other olefins, jute, rayon, wool and cotton based carpets [4, 17]. Almost 97 percent of carpet manufactured in the United States is made from synthetic fibers because synthetic fibers are more resistant to stains while being less expensive [17]. However, as
mentioned earlier nylon based carpets are used more significantly in the consumer world. About 90 percent of all residential carpets and 65 percent of all carpets are made of nylon fiber [17]. Nylon 6 and Nylon 6,6 account for 45 and 55 percent of the nylon carpet market [11]. In this study Nylon 6,6 based cut pile carpet was used for fabrication of carpet composites.

2.1.3 Nylon fibers

In the field of material science, there is a wealth of information about the invention of synthetic fibers for advancement in quality and availability of textiles. The discovery of Nylon 6,6 in 1931, by the chemists at E. I. DuPont de Nemours and Company pioneered the revolution of textile industry and a huge impact of scientific world in the society [4]. Nylon 6,6 is a polyamide made from two monomers of 6 carbon atoms each, therefore the designation 6,6 [4]. Nylon 6,6 is formed by a polymerization reaction between its monomers hexamethylene diamine and adipic acid [4]. Currently, Nylon 6,6 is made by DuPont, Solutia and other manufacturers and is sold as fibers [11]. By around 1940, Paul Schlack of I.G. Farben Company in Germany obtained a different form of polyamide called Nylon 6 by using caprolactum as a monomer [4]. Currently, Nylon 6 is made by Honeywell, BASF and several other manufacturers and is extruded into fibers by these companies or by the carpet manufacturers. The manufacturers of carpet procure the nylon material from these companies or have their own stand-alone nylon production.

2.2 CARPET RECYCLING

Carpet recycling poses significant challenges due to the inhomogeneous and chemically diverse nature of the materials that constitute a carpet. However literature suggests that there is an increasing trend in research and development of carpet recycling methods. The carpet recycling methods can be broadly classified into four categories:

1) **Primary recycling or depolymerization:** It involves methods to breakdown the long polymer chains into their original monomers that can be repolymerized [4].

2) **Secondary Recycling:** It involves recovery of individual components of a polymeric mixture without breaking them into monomeric forms but by extraction and separation methods [4].
3) **Tertiary Recycling**: It consists of preparing a thermoplastic mixture by melt blending of carpet waste by reactive extrusion and compatibilization. Injection molding on this blend would yield products of lower quality [4].

4) **Quaternary Recycling**: It involves energy recovery by incineration of carpet waste [4].

Research and development of carpet recycling has mostly been a part of one of the above four recycling techniques. However, there are companies like Polyamid 2000 that have processes to integrate all the above four recycling techniques into one recycling process. Polyamid 2000 has a plant at Premnitz (near Berlin, Germany) at operation since 2000 that uses a recycling process similar to the one illustrated in Fig. 2.7.

**Figure 2.7** *An adapted block diagram of Nylon recycling process used by Polyamid 2000 [18]*

The following sections cover a detailed technical literature review of the carpet recycling methods, covering patent and non-patent literature review.

2.2.1 *Primary Recycling*

Primary Recycling or Depolymerization is currently the most preferred route for the carpet recycling industry, since it allows recovery of monomers from carpet fibers (nylon 6 and/or nylon 6,6) which can be repolymerized into new nylon products of high quality [4]. In the non patent literature review, the paper by Mihut C et al. has provided valuable background information. Mihut C et al. mentioned the following depolymerization techniques of carpet in their paper [4] –

(a) Depolymerization with Ammonia (Ammonolysis)
Acid/Base Catalyzed Depolymerization

Depolymerization with water

Pyrolysis

The depolymerization reaction of nylon 6 is a thermodynamically favored at low pressures and high temperatures, above the boiling point of caprolactum (around 267°C). At low pressures it becomes easier to separate the monomers from the melt. High temperature using superheated steam is maintained because the process is endothermic. For depolymerization of nylon 6,6 the key focus is the recovery of hexamethylene diamine and adipic acid. Some depolymerization processes include addition of catalyst for easier processing of polymeric melt by decreasing its viscosity [4]. The following Fig. 2.8 illustrates the companies involved in depolymerization process of nylon based carpet.

Figure 2.8 An illustrative chart listing major companies in depolymerization of nylon carpets

Depolymerization of nylon 6 with superheated steam is a highly researched depolymerization technique. Patent literature review suggests that Sifniades et al. of Allied Signal Inc have several patents on the recovery of caprolactum from nylon 6. In their patent they developed a depolymerization technique using superheated steam [19]. In multicomponent mixtures or composites like waste carpets that contain nylon 6, the recovery of caprolactum is complicated due to presence of other components [20]. Sifniades et al. August 1999, mention in their patent that, “In the absence of added catalyst, contacting the multi component waste material with superheated steam at a temperature of about 250°C to about 400°C and at a pressure within the range of about 1 atm to about 100 atm and substantially less than the
saturated vapor pressure of water at the temperature wherein a caprolactum-containing vapor stream is formed.

The formed caprolactum may then be used in the production of engineering resins and fibers” [20]. Mayer et al. of Allied Signal Inc. modified the depolymerization process by stating that “if the crude caprolactum from depolymerization of polyamide-containing carpet is subjected to a simple flashing operation, the condensed flashed material may be crystallized from the aqueous solution and good quality, washable crystals are obtained. The resulting caprolactum has purity greater than 99.9 weight percent (excluding water) and a permanganate number less than 3 and a color number less than 2 and thus, is world class caprolactum.” [21]. Hendrix et al. (1997), mention that depolymerization can take place at a pressure of between 0.5 and 5 atm in the presence of a nitrogen containing compound [22]. By this process, the carpet waste would undergo a depolymerization reaction to yield monomeric components and by-products.

These by-products can be removed with a nitrogen containing compound via gas phase [22]. Allied Signal Inc. patented a process to remove caprolactum from hydrolyzable polymer like nylon based carpet waste by depolymerization with water [23]. The process by Jenczewski et al. (1997), uses “whole carpet,” that is carpet that has not been subject to any mechanical separation, including cleansing of carpet [23]. The whole carpet is then fed to a continuous stirred tank reactor (CSTR) or a series of CSTR’s via a gear pump, wherein it is mixed with water to form a multicomponent polymeric waste material [23]. This mixture is then subjected to heat and pressure to form a liquid aqueous solution and therefore dissolution of nylon 6 and partial depolymerization of nylon 6 is achieved [23].

Caprolactum is obtained by flash distillation while the water insoluble waste is sent to centrifuging to obtain solids which are combusted in a separate step to provide process heating and evaporation [23]. Frentzen et al. suggest their patent process for depolymerizing nylon 6 and recovering caprolactum from the depolymerized products by extraction with alkyl phenolic compounds [24]. Corbin et al. 1999, focused on the process improvement in depolymerization of nylon-6 carpet with water [25]. Corbin et al. invented the process in which the carpet is fed in a separator to prepare a scrap containing nylon -6 and auxiliary materials, from which, the scrap is fed to a depolymerizer reactor to produce a distillate rich in epsilon – caprolactum [25]. Here in the distillate epsilon - caprolactum is separated from other volatiles and the auxiliary materials are recovered and reused [25]. While most of the patents focused on recycling of nylon -6 to
caprolactum by depolymerization with water, Dos Santos et al. 1999, focused on treatment of material comprising nylon 6,6 to recover adipic acid and other diacids [26].

A patent literature review revealed that depolymerization of nylon fibers into its monomers by pyrolysis is not as studied in comparison to other depolymerization techniques. However a few like Moens et al. (1999), invented a process to purify caprolactum from recycled nylon by pyrolysis [27]. Moens et al. (1999), used carbon dioxide to form a precipitate in a solvent dissolved pyrozylate containing caprolactum mixture [27]. The precipitate is removed from the solution and pure caprolactum is recovered from the solution [27].

The non-patent literature suggests that depolymerization of carpet fibers by acid/ base catalysis has been a highly reviewed and widely researched topic. Recovery of epsilon - caprolactum from waste polyamide 6 is competitive with traditional synthesis process and has a significant environmental impact [28]. The Zimmer AG process, which is applicable only for pure polyamide 6 materials, performs depolymerization using steam and a liquid catalyst like phosphoric acid [28]. The disadvantage of Zimmer process is the high yield of salts and traces of phosphoric acid in the recovered ε-caprolactum [28]. Generally, depolymerization of carpet using acid catalysis requires separation of fibers (beneficiation) from other components because CaCO3 present in fillers can consume an equivalent amount of acid catalyst used [4]. Moran and McKinney have worked extensively on the subject of depolymerization using catalysts using acid catalysts using aliphatic carboxylic acid like acetic acid and propionic acid to obtain caprolactum and adipic acid. According to the patent by Moran et al. for DuPont the following process for depolymerization of waste carpet was developed -

“Waste carpets can be entirely dissolved in the aliphatic acids used, at temperatures of about 110°C and atmospheric pressures. Carpet components, such as the primary and secondary backings and adhesive binder, which are insoluble in the acid solution, can be separated by hot filtration. The acid filtrate, containing the nylon fibers, can then be subjected to the depolymerization process” [4].

Bockhorn et al. investigated the use of two types of catalysts – basic catalysis by NaOH/KOH and acidic catalysis by H3PO4 (by Zimmer procedure), which remain liquid under degradation conditions [28]. Bockhorn et al. concluded that “The basic catalyzed degradation of polyamide 6 proceeds via an anionic chain mechanism, which explains high reaction rates in contrast to the non-catalyzed and acid catalyzed reaction. For these reasons, basic catalyzed
Degradation of polyamide 6 is the best suited for an application in a procedure for epsilon-caprolactum recovery from polyamide 6 wastes” [28]. Depolymerization with ammonia involves heating of a mixture of polyamide 6 and polyamide 6,6 in the presence of ammonia at high temperatures and pressures [4]. Lewis acid catalyst precursors makes the process highly efficient [4].

**Figure 2.9** An adapted block diagram of Dupont’s Nylon recycling process [18]

### 2.2.2 Secondary Recycling

In the process of secondary recycling, the entire waste carpet is dissolved in a solvent at high temperature and nylon can be extracted from its fibers in its polymeric forms by extraction process. Although the final product can be used in various applications by further processing using injection molding, the problem associated with this process is the selection of a suitable solvent that selectively dissolves the nylon fibers and does not interact with or dissolve with any of the other carpet components [4].

A patent literature review of secondary recycling of carpet reveals that Roberts et al. in 1997 patented a process for recovering and recycling carpet waste by contacting the carpet waste with a solvent (like formic acid or phenol) which selectively dissolves nylon to form a polymer solution. The solvent and polymer solution is then contacted with an anti-solvent (like carbon dioxide) which causes the polymer to nucleate and precipitate from the solution [29]. The process is efficient in terms of safety and cost because it can be operated using low temperature solvents and anti-solvent like carbon dioxide for re-crystallization of nylon [29]. Costello et al. devised a method to remove the adhesive material from the waste carpet material feedstock to
obtain a mixture containing face fiber material which is passed with a liquid medium to a hydrocyclone for separation of solids on the basis of specific gravity for recovery [30].

2.2.3 *Tertiary Recycling*

Tertiary recycling of carpet materials consists of melting or extruding the waste carpet to form a blended mixture which is then used in production of injection molding and thermoplastics [4]. Although melt blending is cost effective compared to the expensive separation and depolymerization procedures it produces low quality melt blended plastics due to poor chemical compatibility of carpets [4].

Kotlair and Fountain developed a simple process to convert carpet waste to individual shreds, which on coating with structural adhesives gives fibrous composites like “synthetic wood” [4]. Young et al. focused on recycling of automotive carpet scrap by extruding the shredded and granulated carpet scraps [4]. The melt produced is then pelletized and used in injection molding applications [4]. This process for Lear Corporation included applications like flexible floor mats, automotive carpet backing with superior sound-insulating properties and rigid materials like door panels and truck liners [4]. David et al. developed an alternative method of melt blending for Monsanto in which, the carpet waste without excess dirt is fed into a twin screw extruder at a shear rate of 200 to 400 sec\(^{-1}\), a temperature between 250°C and 350°C and a pressure of 350 to 450 psi [4]. Intense mixing converts the carpet into a flowing homogenous mixture containing all the components of carpet - face fiber, backing and SBR adhesive, which on cooling yields a resulting material exhibiting thermoplastic characteristics in the form of pellets or chips [4].

2.2.4 *Quaternary Recycling*

Incineration or quaternary recycling helps in partial recovery of the energy content of the waste materials like carpet waste. Two types of ash are produced by the incineration process: fly ash (fine particles entrained in the incinerator exhaust gases) and bottom ash (the large and heavy particles removed from the bed of the incinerator) [31]. Land filling these ash residues is not always acceptable because of the potential for groundwater and soil pollution due to leachate carrying heavy metals like lead and cadmium [31]. Even though there has been public resistance for incineration due to its emission of toxic fumes, there has been significant improvement in technology which makes incineration a robust recycling option [31]. Most of the municipal solid waste (MSW) combustion currently practiced in the United States involves recovery of an
energy product (generally steam or electricity). The resulting energy reduces the amount needed from other sources, and the sale of the energy helps to offset the cost of operating the facility [8]. The USEPA report of 2009, in its chapter on Management of Municipal Solid Waste states that “Total U.S. MSW combustion with energy recovery, referred to as waste-to-energy (WTE) combustion, had a 2009 design capacity of 94,721 tons per day. There were 87 WTE facilities in 2009, down from 102 in 2000” [8].

There is also a trend in facilities seeking combustion of source separated MSW for use of rubber tires and used paper / wood products in boilers, cement kilns, pulp and paper mills, industrial boilers and dedicated scrap tire to energy facilities [8]. The use of tires have decreased the air emissions by 30 % for a plant at Atlanta, Georgia and has therefore helped the company to meet the stringent nitrogen oxides (NO\textsubscript{x}) guidelines [32]. It is estimated that about 3.1 million tons of MSW were combusted in this manner in 2009, with tires being the majority in this contribution [8]. However, the volume of these MSW is reduced by about 75% after incineration and the post-combustion ash needs to be treated separately and then land filled thereby owing to general public concerns [32]. The use of carpet waste in cement kilns is also an attractive option since the relatively high fuel value of carpet polymers can reduce the need for fuels, and the calcium carbonate in carpet could become a raw material for cement [32].

2.3 WASTE CARPET RECYCLING

Fig. 2.10 shows the products obtained by recycling of carpet using the traditional methods of recycling. The traditional methods of carpet recycling have many shortcomings of which the predominant drawbacks are - 1) being highly uneconomical and 2) excessive process time. This defies the aim of a sustainable development process for product recycling. Recycling is defined as a process to make products better in terms of properties and value than its current value. But the recycling process should be cheap and viable to make it lucrative. This helped us in choosing better products like - polymer composites, which are cheaper in value and their manufacturing process for recycling waste carpets. Hence VARTM was used to recycle waste carpets into structural composites, which can be used as highway noise barriers. The next section talks in detail about the need for highway noise barriers and the requirements for a material to qualify as a highway noise barrier.
2.4 HIGHWAY NOISE BARRIERS

For vehicles in motion in highways, friction between the vehicle’s body and the air touching the vehicle causes a change in the gradient of the air pressure field induced by the friction [33]. This friction renders aerodynamic effect called noise which can be heard at significant distances [33]. Noise is also created due to engine operations, exhaust systems and the contact of grooved tires on pavement surfaces [33]. This noise with a spectrum of multiple frequencies is called traffic noise [33]. A majority of this spectrum falls within the range of 250 Hz and 4000 Hz [33]. Human ear can easily hear a noise of this frequency and therefore traffic noise causes great discomfort to mankind. Noise barriers are built along highways to control the noise pollution because the barriers contain and absorb the noise within them and do not allow any propagation of noise to any significant distance [33]. As shown in the illustration below in Fig. 2.11 traffic noise is caused mainly due to traffic volume, speed of the traffic and volume of trucks.

Figure 2.10 A diagram of products obtained by traditional carpet recycling processes
A formal definition is needed to define highway noise barriers as it one of the potential applications for our material - carpet composite. A highway noise barrier is defined as a barrier wall that attenuates noise by preventing the direct propagation of noise between the noise source and the listener by absorbing the sound energy incident upon the surface on the barrier [35]. Hence measurement of absorption of sound energy incident upon the surface of the barrier material is the most critical parameter for evaluation of the performance of the material as a barrier. This measurement can be achieved using acoustic characterization test based on the American Standard for Testing Materials, ASTM standard C 384 - 04. This characterization would enable a correct assessment of the structural material as a highway noise barrier. Barriers are generally applicable to the design of new highways and are an attractive means of providing noise relief to communities along existing highways [35]. There are four primary requirements for an acceptable highway noise barrier [35]:

1) It must function effectively as a noise reduction structure

2) The barrier must be a safe roadside structure

3) The total installation costs must be acceptable, and

4) The barrier must have an acceptable appearance.

Currently most highway noise barriers are built with precast concrete or concrete slabs, which have high acoustic reflectivity (95% and above) and low noise absorption for the frequency band of highway noise between 250 Hz and 4000 Hz [33]. There is a possibility for alternative highway noise barriers made up of materials with higher noise absorption coefficient.
or even the use of vegetation as a noise barrier. Vegetation can decrease highway traffic noise if it is high, wide and dense enough. A 200 ft width of dense vegetation can reduce noise by 10 dB, which cuts the traffic loudness into half, but is impractical to plant such scale of vegetation – at this era of rapid development [34].

![Concrete Noise Barrier, Wooden Noise Barrier, Earthberm Noise Barrier](image)

**Figure 2.12 Types of Highway Noise Barriers [34]**

Due to exponential economic growth of the world economy in the past few decades, there has been a drastic increment in highway traffic and a need to develop new and better noise reduction barriers for highways and airports. This trend in increase of highway traffic is likely to continue in the near future and so has been an increase in dedicated research efforts to develop new noise barriers. Polycarbonate noise reduction panels made by a California based company, “Quitite International”, have been used to build polycarbonate noise wall near Culver City park in Los Angeles, California in 1996 [33]. Also a jet engine testing center was installed at Albany airport, Albany, New York in 1997 using polycarbonate noise barrier [33]. In both these cases the polycarbonate was made from Lexan plastic produced by GE Plastics [33]. Carsonite International of Early Branch, South Carolina developed hollow panels made planks of reinforced composite material crumbed rubber tires as nano-fillers [33].

The literature review indicates that Wolfe et al. investigated the usage of cement wood composite as highway noise barriers while Watts et al. studied the effects on roadside noise levels by applying sound absorptive materials to the traffic face of noise barriers [36, 37]. Watts et al. also used the boundary element method to study the combined effects of porous asphalt surfacing and noise barriers on traffic noise [37]. Yamaguchi et al. studied the sound absorption mechanisms, void ratio and acoustic properties of porous asphalt surface by comparison with other porous materials like mineral wool and synthetic wool [38]. Meiarashi et al. and Oshino et al. studied the sound absorption mechanism of porous asphalt pavement and proposed a model
The factors that influence the structural design of barriers are both acoustic and non-acoustic. Non-acoustic factors include wind loads, fire-retardancy, and the weather and aging effects (which cause warping and corrosion) [42].

### 2.5 ENVIRONMENTAL DEGRADATION OF CARPET COMPOSITES

The performance of the carpet composite demonstrated high potential as a new material for highway sound barrier and other structural application in terms of acoustic and mechanical properties. However, during long-term outdoor type of applications, the composite panels would be exposed to UV, moisture, heat, wind, dust and other pollutants that could potentially deteriorate the properties of the composite panels. [43] Among all the environmental elements, moisture and UV play important roles in physical and chemical change or degradation process of polymers. [43]

The following section is about the possible degradation that the carpet composites may undergo due to the interaction with moisture and Ultra-violet (UV) radiation.

#### 2.5.1 Effect of water on polymer

Chemical and physical changes are observed in polymers due to interaction with moisture. Chemical changes involve hydrolysis and leaching of additives. As a result, plasticization effect can be seen in polymers that lead to changes in mechanical properties. [44]

The water absorption kinetics in polymers is generally divided into two types: 1) Fickian and 2) non- Fickian. For the first kind, absorption follows the Fickian law of diffusion, which in one-dimensional case is governed by the following equation:

\[
\frac{\partial c}{\partial t} = D \frac{\partial^2 c}{\partial x^2} \quad \text{.................................(1)}
\]

This equation can be simplified in the following form relating mass gain to diffusivity and thickness h with time[14] :

\[
\frac{M_t}{M_\infty} = 4 \left( \frac{Dt}{\pi h^2} \right)^{\frac{1}{2}} \quad \text{.................................(2)}
\]
Taking initial mass $M_\infty$, thickness $h$, and diffusivity $D$ as constant, mass gain becomes proportional to square root of time. As a result Fickian diffusion can easily be identified from linear relationship of mass change and square root of time. Fickian diffusion of polymers involves absorption through nano or micropores of the polymer, where non-Fickian indicates boundary separation in different polymer phase and causes swelling and residual stresses in polymers. [45]

Epoxy has strong affinity towards water. Absorption and transportation mechanism of water has been of interest for many researchers. Different techniques have been utilized to figure out whether the absorbed water molecules are in motion or static. Most of the findings came to the conclusion that water is stable inside the epoxy and water molecules are attached to specific locations of the molecular chain of epoxy. It is assumed that nano voids open spaces for the water and polarity of the epoxy facilitates attachment of water molecules with polymer chain. NMR study by Fuller et al. indicated absorbed water molecules by epoxy is not in motion. [46] They also found hydrogen exchange in between water and epoxy. [47] Similar findings was reported by Ellis and Karasz and they suggested that one water molecule can interact with six OH sites of epoxy polymer. [48, 49] Along with the polarity, free void of epoxy facilitates water motion. During the formation of covalent bond in curing process, nanopores of 5-6 Å size are created which is big enough to occupy water molecule. [50] As the absorption process goes on, water absorption is slowed down not only because of the decrease of free void, but also for the hindrance created by the other water molecules strongly attached to the polar sites of the epoxy chain. [51]

However, epoxy can be saturated by 1 to 7% of water of its weight depending on the nature of epoxy. [52] Due to interaction with water, there can be change in physical and mechanical properties. One of the after effects of water absorption is depression of glass transition temperature ($T_g$) that is attributed to plasticization of epoxy. [53] Moy and Karasz reported that depression of $T_g$ increases with the increase in amount of absorbed water. [54] Absorption of water leads to formation of microvoid and microcracking, as a result of which loss of mechanical properties is observed. [55]

Similar effect of moisture is observed in nylon. Nylon 66 can absorb up to 9% of moisture of its weight. [43] Reduction in tensile strength and modulus has been reported due to degradation by moisture. Reduction in elastic modulus may occur in nylon, and high temperature
hygrothermal ageing can increase crystallinity. [56] Reduction in strength has been found as much as 82% after exposure to 100% relative Humidity in 100 °C up to saturation moisture absorption. [57]

In fiber matrix composites, interfacial bond between fiber and matrix is highly affected due to moisture absorption process. Debonding and weakening of the fiber-matrix bond is observed due to interaction with water. [57-59]

2.5.2 Effect of UV on polymer

Ultra violet (UV) radiation is responsible for chain scission, cross linking reaction, and photo-oxidation of polymer. UV exposure of epoxy leads to chain scission and surface cracking. [60] Due to irradiance of UV, surface damage and initiation of crack have been found in glass epoxy composites. [61]

Chemical, mechanical and physical changes in nylon are found when exposed to UV. Many authors have studied chain scission of nylon after UV exposure. Figure 2.13 shows chain scission reaction in the most possible order [62]. FT-IR analysis showed that chain scission reaction of nylon 6,6 can be accompanied with emission of gases such as H₂, CO, NH₃, carbonyl compound. [62, 63]. Longer exposure of UV can lead to increased cross-linking reaction. [64] Along with the chain scission and cross-linking reaction, photo oxidation is another possible way of degradation of nylon in presence of UV. [65] Chain scission makes the polymer weaker and cross-linking leads to brittleness and cracking.

Synergistic effect of UV and moisture is seen on polymer and polymer matrix composites. Kumer et al reported results of combined effect of UV and condensation on carbon fiber reinforced epoxy composites. Degradation process is accelerated and erosion can happen in polymer matrix by alternative exposure of UV and condensation. [66] It is assumed that the role of moisture in the combined effect of UV and moisture is to increases the mobility of the catalyst and thus enhance the degradation process.
2.6 CLAY-POLYMER NANOCOMPOSITES

Clay is a montmorillonite mineral which has 2:1 sheet-layered smectite platey structure. The elementary structure consists of 2 silica tetrahedron and 1 aluminum octahedral sheets. [67] These three sheets together make a clay layer whose thickness is 0.96nm. [68] High aspect ratio and high specific surface area have made nanoclay as an attractive nanofiller for polymers. The aspect ratio (thickness to diameter ratio) of nanoclay ranges 200 to 500, and specific surface area is more than 750 m²/g. [67] Because of this high surface area, addition of small amount of nanoclay can contribute to significant improvement in mechanical properties as well as barrier properties against liquid and gaseous substants. To obtain the maximum performance of nanoclay, proper dispersion or exfoliation of nanoclay layers is extremely required. Based on the exfoliation or orientation of clay layers in polymers, nanoclay-polymer composites are classified into three groups (Figure 2.14) : 1) Conventional composite, 2) Intercalated Nanocomposite, and
3) Exfoliated Nanocomposite. [69] In conventional composites, there is no inclusion of polymer molecule in between the clay layers. In the second type, molecules of polymers enter in between clay layers and increase the distance of the layers. For exfoliated nanocomposites, individual clay layers are separated from each other by polymer molecules.

![Diagram of different clay-polymer composites]

*Figure 2.14: Schematic illustration of different clay-polymer composites. Modified from [69]*

2.6.1 Effect of clay on properties of polymer and composite

Clay has been used with different polymers as a reinforcing agent. Proper mixing of clay restrains the movement of polymer chain, and enhancement in modulus is generally observed. Kojima et al. introduced use of clay polymer and reported enhancement of tensile strength, tensile modulus, flexural strength and modulus. [70] Lan and Pinnavaia found increasing trend of tensile strength and modulus in DGEBA epoxy with increasing amount of clay loading. [71] Chan et al. used different amount of clay and measured the strength and modulus. They found increasing trend of clay modulus and strength of epoxy up to 5% wt of clay loading and reduction in properties with higher amount of clay. According to their observations, clay platelets can hinder crack propagation in polymer matrix and thus help improving strength. On the other hand, higher loading creates agglomeration and works as discontinuity in matrix. [72] Zerda et al. has found similar effects of agglomeration. They reported improvement of properties up to 3.5% of clay loading and decrease with higher amount of clay. [73] Zaarei et al. found enhancement of microhardness, toughness and wear resistance for use of clay with epoxy as coating. [74]
2.6.2 Effect on barrier properties

Reduction in absorption and enhancement in barrier properties for liquid and gas have been observed by addition of clay. These effects have been attributed to the tortuosity effect due to higher aspect ratio of clay platelets. Based on the tortuosity effect, Neilson's model has been developed which predicts the permeability properties of clay composites against liquid and gaseous molecules by the following equation [69], given in Figure 2.15:

\[
P_{\text{nanocomposite}} = \frac{(1 - \phi)P_{\text{matrix}}}{1 + \alpha \frac{\phi}{2}}
\]

Here, \( \alpha \) is aspect ratio, \( \Phi \) is the volume fraction of nanofiller and \( P \) represents permeability. According to the model, because of the high aspect ratio of nanoclay, small volume fraction of nanoclay can provide significant improvement in barrier properties.

![Illustration of Neilson's tortuous path model for reduction in permeability. Image modified from [69]](image)

Experimental results show that the tortuosity effect significantly reduces the permeability of moisture and gas in nanoclay-polymer composites than the polymer itself. [45, 75, 76]

2.6.3 Impact of clay on degradation mechanism

Clay can bridge nanovoids and hinder degradation mechanism. This is the way it reduces the moisture absorption rate in epoxy. Not only that, by preventing formation of micro-crack due
to environmental effect, it can reduce the effect of degradation on mechanical properties. Kim et al. used different types of clay and found reduction in moisture absorption while compared to neat epoxy. [45] Studies of effect of NO\textsubscript{x} on the mechanical properties have also suggested that clay-polymer composites can retain tensile strength better than neat polymer after degradation. [77] Fractographic analysis by woo et al. indicated that initial degradation process by UV cannot be delayed by the use of clay, but effect of longer UV exposure is prevented. Clay can hinder diffusion of oxygen and other free radicals and thus prevents crack propagation along the thickness of the material. [60] Bagherzadeh et al. applied 60±5μ coating of epoxy-clay on metal and found better anti-corrosive behavior. [78] Singh et al. reported the effect of clay on retention of mechanical properties of epoxy against alternating exposure of UV and condensation. According to them, clay decreases the formation of cracks and micro-channels, and provides higher flexural strength and modulus after combined degradation. [79]

2.6.4 Effect of clay on fire retardancy of polymer

Polymers are more vulnerable than metal, ceramic, or concrete as construction materials because of their flammability. Polymer can start a flame once the surface temperature reaches ignition point. For most polymers, ignition temperature ranges from 275°C to 475°C. [80] Norman Grassie proposed a simple model of fire propagation mechanism through polymer (Figure 2.16). [81] According to the model, once the fire is ignited, because of the heat, polymer undergoes thermal decomposition and produces liquid and gaseous components that take part in the burning process.

**Figure 2.16: Flame propagation model proposed by Norman Grassie.** [51] (Reused with permission from Elsevier Limited)

Different approaches to improve fire retardancy are targeted to discontinue the fire propagation cycle by breaking it on the point A, B, or C (Figure 2.16).

Use of nanoclay or any other nanomaterial can create barriers against propagation of combustible product produced by heat and delay the flame propagation. Nanoclay hinders the
physical movements of flammable product by two methods: 1) Barrier effect and, 2) formation of char. [82]

The network structure formed by nano particles creates a thermal shield for heat, oxygen and other combustible product and reduces supply of fuel for flame. On the other hand, in many cases, use of nano components helps formation of robust char that prevents flow of combustible product produced by heat during burning process. [82] Kashiwagi used nanosilica, nanoclay, and carbon nanotube with PMMA, polyamide, and polypropylene, respectively, and observed fire properties in a cone calorimeter. It was reported that addition of nanoparticles can delay ignition time, reduces peak, and average heat release rate compared to neat polymers. The aspect ratio of the nanoparticle was attributed as the controlling parameter on the effect of flame retardancy. Analysis of the final burning product showed higher aspect ratio of the particle is favorable for formation of char and reducing crack during burning process. [83] Clay has been used in polymer with other flame retardant chemicals and reduced flammability was reported. Nazare et al used nanoclay in polyester with different flame-retardants and found delay in ignition time and decrease in total and peak heat release. [84]
3. COMPOSITE FABRICATION

The VARTM process for recycling carpet has several advantages compared to traditional carpet recycling techniques like depolymerization and melt blending that require high energy. Singh et al. (2009) [6] made a detailed analysis of recycling waste carpet into carpet composites by VARTM with various thermoset resins. Singh et al. (2009) [6] used hot pressing for high temperature post curing step to achieve carpet composites as a final product. The use of hot press limits the fabrication of composites of different shapes and sizes primarily because hot press is designed for flat composite panels of a fixed size. Jain (2010) [85] extended the work performed on feasibility of carpet composites by studying the effect of carpet fiber treatment with Lewis acid / base chemicals on resulting carpet composites by VARTM. Jain (2010) [85] makes use of hot pressing as a post curing step to make carpet composites.

However the challenge remains to scale-up carpet composites of larger dimensions and different shapes. Hence a scaled-up process of fabricating large-scale carpet composites was required. The following process of scaling up of VARTM was followed -

Samples were fabricated using a simple, vacuum assisted resin transfer molding fabrication technique with different combinations of layers. In the scaled up VARTM fabrication process caul plates and green mesh cloth (similar to SCRIMP process) are used. Initially, the glass plate is cleaned with acetone. Three layers of mold release wax and one layer of PVA release film coat are applied over the surface to ensure an easy removal of composite after completion of resin curing period.

Nylon bagging material (procured from Airtech International Inc.), tacky tape and connecting tubes (procured from Fiberlay Inc.) are cut to the required size. The tacky tape is laid to a size slightly greater than the size of effective material (e.g., the infusion area over the glass plate) and the carpet is placed between the tapes. The scaled-up VARTM fabrication process involves use of a thin release sheet placed over the carpet for minimum resin use. Thin pores on the sheet allow excess resin to flow out of the pores, which is then carried to the next level of green mesh (SCRIMP model) which adds local stress points inside the bag so that resin runs through carpet sample to reach the vacuum end of the set-up. A polyester release cloth is placed over the green mesh cloth to ensure that the resin does not stick to the vacuum bag and the bag can be attempted to be made reusable. Breather cloth is placed over the release cloth to ensure distribution of air (if any) in the vacuum bag. The breather cloth distributes the air evenly in the
setup if in case of air pocket. In the conventional VARTM nylon bag is used to enclose the entire set up from end to end over the tacky tape with attention paid to avoiding air pockets at any end. In the scaled up VARTM process, the same procedure is used but over an aluminum sheet placed over the breather cloth layer (thereby sandwiching the entire carpet sample assembly). This sheet of aluminum is a caul plate that saves in the cost and simplifies VARTM by avoiding the hot press. The caul plates used are of the same dimensions as the sample of carpet cut.

3.1 MATERIALS AND COMPOSITE FABRICATION

Fig. 3.1 (a) and Fig. 3.1 (b) show a schematic of the setup of scaled up VARTM process used for fabrication of carpet composites.

Figure 3.1 (a) An illustrative schematic of the scaled up VARTM process (Plan View).
Figure 3.1 (b) An illustrative schematic of the scaled up VARTM process (Elevation View).

The scaled up VARTM set-up constitutes laying the preform fabric (nylon based carpet) on a smooth surface tool or glass plate or a flat table. A release material like wax is applied to the mold so that after the cure of the part it is easy to demold the composite from the mold or glass plate. The resin inlet and the vacuum outlet piping are placed at both ends of the laid fabric. A release film, peel ply, low permeability resin distribution media, and breather cloth, in that order, are placed on the top of the fabric. Finally a vacuum bag is placed on top of this set up and is sealed with the help of a sealing tape. Finally, vacuum is drawn using a vacuum pump. After the setup of this process, the resin is infused (pulled by the vacuuming force of the pump) through the carpet. Thereafter the inlet manifold is clamped so that the resin feed can be stopped. The impregnated carpet fabric is allowed to cure. The type of resin and hardener combination used for resin impregnation decides the curing time.

Mold/Glass Plate

The mold is defined as a rigid material that does not change its shape during the VARTM process and acts as a stable base. In this study a stiff glass plate with fine finish was chosen as a mold. The glass plate gave a smooth finish for one side of a flat panel. However in case of VARTM for larger samples, of the size 0.61 m x 1.22 m (2 ft. x 4 ft.) the surface of the work-bench was used as a mold and sometimes, the floor was used as the flat end mold surface. Although the resulting panels had a smooth finish, the surface of workbench and floor as a mold
gave other operational problems like cleaning the spillage of excessive resin and change in flow pattern of resin.

In case of curved shaped panels, a thin aluminum plate was curved to the shape of an arch and used as a mold.

**Reinforcing Fabric - Carpet**

Reinforcing fibers are the backbone of any composite material. In this study Nylon, cut loop type of carpet of the commercial name ESSENTL TWTGO, color Sandstone by Shaw Industries was chosen.

These carpets of size 3.66 m x 1.22 m (12 ft. x 4 ft.) were bought from M/s Georgia Carpet Industries Inc., Atlanta, Georgia, and then pieces of sizes 0.15 m x 0.15 m (0.5 ft. x 0.5 ft.), 0.15 m x 0.24 m (0.5 ft. x 0.8 ft.), 0.24 m x 0.24 m (0.8 ft. x 0.8 ft.), 0.30 m x 0.30 m (1 ft. x 1 ft.), 0.61 m x 0.61 m (2 ft. x 2 ft.) and 1.22 m x 0.61 m (4 ft. x 2 ft.) were cut in the sample preparation process for fabrication of composites.

The carpet was laid up in three different configurations - TBBT, BTTB and TBBTTBBT (4 layer carpet), shown in Figure 3.2.

*Figure 3.2 Images of carpet configurations used for VARTM*
**Resin**

The sole purpose of the use of thermoset resin is to impregnate the carpet layers and act as a matrix for the resulting carpet composite. The characteristics of the resin that directly affect the VARTM process are viscosity, gel time and the type of resin. Less viscous resins might infuse quickly, resulting in deformed parts. This deformation is due to uneven wetting of the fabric making some portions resin rich and others full of voids with dry-spots, while faster curing resins, with high viscosity gel too quickly and cannot sometimes impregnate the entire part [53].

In this study, the resin systems used were vinyl ester, polyester and epoxy resins. Even though vinyl ester and polyester resins are available cheaply in the market, styrene emissions is a problem, causing shrinkage in the parts resulting in stresses between the layers [86]. There were attempts to do VARTM with epoxidized soybean oil (ESO) but it proved to be unsuccessful because it could not crosslink.

In this study, the resins - Epoxy 2000 (hardener Epoxy Hardener 2120) and polyester and vinyl ester (with hardener: methyl-ethyl-ketone-peroxide) (MEKP) were procured from Fiberglass Corporation. The Epoxy 2000 resin was prepared by mixing with hardener with a weight ratio of 27/100. For example for every 1000 grams of epoxy resin, 270 grams of hardener was added and then stirred for effective cross linking. The pot life of the resin was 120 minutes due to the addition of Epoxy 2120 hardener used for mixing and therefore proved advantageous for larger samples. Polyester resin was prepared by adding hardener MEKP in the weight ratio of 1/100 while on the other hand for vinyl ester resin MEKP was added in the weight ratio of 1.25/100. Thus for every 1000 grams of polyester or for every 1000 grams of vinyl ester resin, 10 gm of MEKP and 12.5 grams of MEKP are added respectively. Since the curing temperature of both polyester and vinyl ester resins takes place at room temperature itself, the pot life for these resins is as low as 20 minutes. A curing period of 24 hours was maintained to ensure complete curing for all resin types.

A Mettler Toledo weigh balance was used to measure the accurate amount of resin and accurate amount of hardener mixture before infusion. The weigh balance needs to be accurate as improper ratios can cause uneven curing, change the properties of resin flow inside the preform.
and cause gel times to change [86]. A balance with an error range of 0.01 gm was used because the proportion of hardener to the resin should match exactly the manufacturer specifications.

The resin distribution media is used to accentuate the distribution of resin from the inlet piping to the preform and makes the resin moves faster due to the pressure gradient created. The resin resin distribution media used was “Greenflow 75” bought from Airtech International Inc. The resin resin distribution media efficiently distributes resin with little waste due to its low profile and tight construction.

Generally, the resin distribution media is placed from the inlet-piping end to the outlet, without touching the outlet helix tubing. Generally, the gap between the outlet helix tubing and the resin distribution media is 31.75 mm (1.25 in) [87]. However in the scale-up of VARTM process, the resin resin distribution media covers the entire area including the outlet tubing. This would create micro voids in the composite panels fabricated. This is because there is always a possibility that some air gets trapped in the helix tubing and addition of the resin resin distribution media over the outlet tubing would aggravate the problem by actually trapping air in the micro - gradients created.

One of the main aims for scale-up of VARTM is to produce composites with excellent acoustic properties. Creation of micro-voids ensures that there is an increase in porosity, which in turn results in improvement of acoustic performance of composites.

3.1.2 Nanoclay

The nano-filler used in this study was Nanomer I.28 (Nanocor, Inc., IL, USA). This clay had been selected because of its compatibility with epoxy. This clay has particle size of 15 -20 micron. The clay surface was modified with trimethyl octadecyl ammonium chloride (Figure 3.3) and modifier concentration is 25-30%. In order to allow the dispersion of clay in an organic medium, a surface modification was required to make it organophilic. I.28 clay has been chosen for the longer chain of modifier salt. Longer chain of the modifier helps better dispersion and higher mechanical strength of the nanocomposites than the clay modified with shorter chain. [88]
3.2. SCALING UP OF VARTM PROCESS

The following section discusses in detail the changes made in order to achieve scale up of VARTM process.

*Vacuum Degassing*

Mixing of the resin and hardener cause air bubbles while mixing that get trapped in the resin. These air bubbles are the main reason for formation of micro-voids because these air bubbles stay in the resin, even after catalysis of resin and travel through the fabric during infusion and contribute to an increase in void volume fraction.

Vacuum degassing set-up can address the arrest of these pre-existing air bubbles in the resin pot after mixing of resin and hardener. The vacuum degassing set-up consists of a vacuum chamber in which the pre-mixed resin pot is placed. Vacuum is drawn into the chamber and therefore the air bubbles rise up to the top of the resin pot and get collected there. After a while there is reduction in the air bubbles. Literature suggests that the process of vacuum degassing creates micro bubbles, which are not seen by naked eye and there is always scope for improvement in degassing, such as addition of nucleation agent [89-90].

However in this study, for scaled up VARTM process, it was intended to retain (or rather incorporate, if possible) micro-voids in the resultant composite. This is because of the unique research objective in the study to develop carpet composites with better acoustic properties. Literature suggests that in energy absorption testing, composite made out of higher void content out-performed a composite made of the same materials and a void content of less than 1% [91].
This indicates that the design of the composite structure should be made keeping in mind the product application intended for the composite. In this study, it was advantageous to have composite samples with higher void content for their use as highway noise barriers in terms of their superior acoustic characteristics.

*Large and Complex structures*

In the conventional VARTM setup for fabrication of large-scale composites, there is structural instability in the final part for composites of the size dimensions 1.22 m x 0.61 m (4 ft. x 2 ft.) and greater sizes. This is because the vacuum bag side is not smooth, which leads to structural instability due to thickness variation throughout the part [86]. Therefore it is quite often a problem to fabricate large scale composite by the conventional VARTM process. This problem could also occur due to inability of the vacuum end in achieving greater compaction of stacked up carpet layers.

This problem was arrested in the scaled up VARTM process where a caul plate could ensure smoothness of surface on the vacuum bag side and thereby possibly minimizes the thickness variation throughout the part. Hence it was observed that it is advantageous to use scaled up VARTM process for fabrication of large-scale composite panels.

For instance multiple inlet ports can be used for fabrication of large-scale composites of the order of size 0.61 m x 1.22 m (2 ft. x 4 ft.) and greater sizes. This has been shown in the following Fig. 3.4 where multiple inlet ports were used for fabrication of large composites because the resin covers a larger area quickly.

*Figure 3.4 A picture of VARTM process using multiple resin inlet ports*

It was envisaged to demonstrate the feasibility of fabrication of complex structures using scaled up VARTM process. Therefore fabrication of complex shapes like arch was planned as an
experiment. The mold prepared for this sample was an empty round shaped canister with an external flat base. One of the caul plates was stuck to the canister to form an arch shaped base of the mold. Carpet layers were stacked in the BTTB configuration. The scaled up VARTM process was rigorously followed and the mold was sealed with a vacuum bag on the top and a caul plate below the bag. Essentially there are two caul plates - in the top of the mold to give a shape to the resulting composite and in the bottom of the mold for aiding in providing an arch shaped base for mold. The BTTB carpet layer configuration was placed between the caul plates after waxing and the standard procedure for fabrication of carpet composites by scaled up VARTM process was followed. During the resin infusion it was observed that the flow of resin is only because of the vacuum inside the bag and was not influenced by other forces like gravity. Post-infusion, a curved shape composite was observed as shown in Fig. 3.5 with an arch with inside diameter of 18 cm.

**Figure 3.5** Image of a curved composite fabricated by scaled up VARTM process

### 3.3 COST ECONOMICS

A preliminary cost model has been developed by monitoring the VARTM process, which has been regarded as a cost effective manufacturing method. VARTM provides significant savings in tooling cost because it requires only a one-piece mold and a vacuum bag is required to close the mold [63]. The total manufacturing cost (TC) consists of material cost (MC), direct labor cost (LC), equipment cost (EC) and the overhead cost (OC) as shown in the following equation 3.2.

\[ TC = MC + LC + EC + OC \]  

(3.1)
Material cost is the major factor in the total manufacturing cost for this particular operation, mainly because the resin system and its curing agents used are expensive. The cost of resin required to fabricate a 0.0935 m² (1 ft. x 1 ft. panel) is around $11. This cost also involves the cost of the curing agent. There could be a maximum wastage of $3 of resin for this size of panel and the cost of curing agent therefore the total materials cost (MC) could be equal to $14 as per the current manufacturing practice of scaled-up VARTM process. However the cost of wastage could be optimized and therefore the total approximate cost of carpet composites prepared by the current manufacturing method could be $12.1.

\[
\text{Resin cost, 11} \quad \text{Total Cost, 12.1} \quad \text{ROM Cost of concrete sound barrier}
\]

\[
\begin{align*}
\text{Resin cost, 5} & \quad \text{Wastage, 3} \\
\text{Total cost, 5.5} & \quad \text{Wastage, 1}
\end{align*}
\]

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{cost_chart.png}
\caption{Preliminary cost model}
\end{figure}

The future material cost could be low because of a regular bulk use of resin. The future cost is estimated to be around $5.5 for a carpet composite panel of the size 0.0935 m². The costs have been estimated on the basis of the material cost data obtained from the material supplier Fiberglast Inc., based in Ohio, USA.

However it can be seen that the cost of concrete is low, about $7 for a 0.0935 m² sample because concrete is a cheap material. However the cost associated with concrete is mainly the
cost of labor and its transportation. The cost of $7 for a concrete panel is a rough order of magnitude cost for the concrete panel of size 0.0935 m².

The cost analysis in this study has been non-conclusive and therefore a judgment cannot be made on the basis of the data presented above. The material cost data suggested in this section is preliminary in nature and has been developed on the basis of lab scale experience. It is an attempt to give a precursor to the detailed cost estimation recommended for this study. A rough understanding of labor involved could be made due to experience in fabrication of composites in this study. VARTM method of manufacturing could be cheap but it is highly labor intensive. The primary labor cost would be the labor involved for 1 worker with 100% presence for making a carpet composite panel of size 0.3045 m x 0.3045 m (0.0935 m²). However as the size of the panel increases to 0.371 m² (0.609 m x 0.609 m) there will be a need for the presence of another worker at a 50% presence factor. This understanding of the requirement of labor at various presence factors was developed during the fabrication of the carpet composites of different sizes in the laboratory.

The following table, Table 3.1 enlists the time taken for various process steps in the scaled up VARTM process for fabrication of carpet composites for a square panel with an area of 0.0935 m² (0.3045 m x 0.3045 m).

3.4 FABRICATION OF COMPOSITES WITH NANOFILLERS

Two types of composites were fabricated for evaluating the effect of nanoclay fillers in the recycled carpet composites. a) clay coated, and b) clay infused composites.

To fabricate clay coated composite, epoxy resin and clay in the weight ratio of 100:4 clay were mixed by magnetic stirring for 24 hours at 120°C, and hardener was added in the mixture. Neat epoxy composite demolded from the vacuum mold was coated with this mixture by a paint roller (Figure 3.7). After coating, the coated material was pre-cured in room temperature for 24 hours and post-cured at 49°C for 6 hours. For clay infused carpet composites, 4 wt% clay were dispersed in the epoxy by magnetic stirring for 24 hours at 120°C, and the clay-epoxy mixture was infused through the carpet layers by VARTM process. The resin was pre-cured for 24 hours in room temperature, and then, post-cured at 49°C for 6 hours.
### TABLE 3.1. TIME TAKEN FOR FABRICATION OF CARPET COMPOSITES USING SCALED UP VARTM PROCESS

<table>
<thead>
<tr>
<th>Sr. No.</th>
<th>Process Steps</th>
<th>Time (min.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Mold Preparation</td>
<td>15 min</td>
</tr>
<tr>
<td>2</td>
<td>Cut materials (Carpet, Vacuum bag, breather cloth, resin resin distribution media, peel ply, release film and visually inspect each of these sheets)</td>
<td>45 min</td>
</tr>
<tr>
<td>3</td>
<td>Installation (stacking the carpet layers and adding clay films on its top), connection of vacuum pump to resin trap, cut inlet and outlet helix tubes and connecting outlet pipes to the resin trap.</td>
<td>20 min</td>
</tr>
<tr>
<td>4</td>
<td>Seal the vacuum bag and vacuum the system</td>
<td>40 min</td>
</tr>
<tr>
<td>5</td>
<td>Resin Preparation</td>
<td>15 min</td>
</tr>
<tr>
<td>6</td>
<td>Resin injection</td>
<td>Depends on resin (Generally less than 10 min)</td>
</tr>
<tr>
<td>7</td>
<td>Resin Cure</td>
<td>420 min</td>
</tr>
<tr>
<td>8</td>
<td>Part removal after resin cure</td>
<td>15 min</td>
</tr>
<tr>
<td>9</td>
<td>Cleaning</td>
<td>20 min</td>
</tr>
<tr>
<td>10</td>
<td>Cutting and Machining of the composite</td>
<td>15 min</td>
</tr>
</tbody>
</table>
Figure 3.7: Coating of carpet composite with epoxy-nanoclay
4. MECHANICAL TESTING AND CHARACTERIZATION OF ENVIRONMENTAL DURABILITY OF CARPET COMPOSITES

4.1. ENVIRONMENTAL DURABILITY AND FIRE RETARDANCY OF CARPET COMPOSITES

The carpet composites were evaluated for their environmental durability and fire retardancy. Two different environments were applied on the composites for degradation studies. In the first case, composite panels were soaked in deionized water, termed as “moisture absorption.” In the second case, cyclic exposure of the ultra-violet (UV) and condensed water vapor were applied on the composite. To observe the effect of environmental exposure, flexural tests were performed for each type of the composite before and after exposure. The mechanical test methods and the environmental degradation conditions used in this study are discussed in the following sections.

4.2. FLEXURAL STRENGTH AND MODULUS

Flexural strength and modulus were determined by three point bending test according to ASTM D 790 standard., using an Instron model 5567 with adjustable supporting anvil of radius 4 mm and loading anvil of radius 10 mm (Figure 4.1).

![Figure 4.1: Flexural test set up for carpet composites](image)

Rectangular bars 25 mm wide by 250 mm long were cut from carpet panels for mechanical tests. The bars were rested on two nose supports and span length was kept at 200 mm. The specimen was then loaded in the middle of the supports at a constant strain rate (Figure 4.1). The crosshead motion was calculated using equation 4.1.
\[ R = \frac{ZL^2}{6d} \] \hspace{1cm} \text{(4.1)}

Where:

- \( R \) = rate of crosshead motion, mm/min
- \( L \) = support span, mm
- \( d \) = depth of the beam, mm
- \( Z \) = rate of straining of the outer fiber, mm/mm/min (equal to 0.01)

Based on the nominal dimensions of the specimen and the equation, the crosshead motion was maintained 5 mm/min for every specimen. Equations 4.2 and 4.3 were used to calculate the flexural stress and strain from the applied load and midpoint displacement data.

\[ \sigma_f = \frac{3PL}{2bd^2} \] \hspace{1cm} \text{(2.2)}

\[ \epsilon_f = \frac{6Dd}{L^2} \] \hspace{1cm} \text{(2.2)}

Where:

- \( P \) = load
- \( D \) = deflection of the midpoint of test span

The flexural modulus was obtained from the stress strain curve. For each type of sample, five specimens were used.

**4.3. MOISTURE ABSORPTION TESTS**

The samples were cut according to the required dimensions for the flexural tests. The samples were then submerged in boiling deionized water for 1000 hours. A bulb condenser was attached on the hood of the water vessel to prevent water evaporation and to maintain constant water level (Figure 4.2). Maintaining same water level helped to keep the temperature steady. Edges of the specimens were coated with high temperature wax to ensure uni-directional water diffusion. Before submerging the samples, the temperature of the water was kept stable at 80 °C ±2°C by controlling the hot plate temperature. To reduce evaporation loss during the weight measurement, samples were placed at room temperature water for 10 min to bring them to
ambient temperature before weight measurement. Before every measurement, specimen surface was wiped carefully to remove the surface water.

![Figure 4.2: Moisture absorption test set up of carpet composites](image)

**4.4. CYCLIC EXPOSURE OF UV AND CONDENSATION**

The combined effect of Ultra-violet (UV) and condensation was studied by exposing several composite samples to alternating cycles of 8 hours UV radiation and 4 hours of water vapor condensation, as per ASTM G154-05. An irradiance of 0.89 W/m² and a temperature of 50 °C was applied on the composite surface in the first step of the cycle. In the second step condensed air was applied at 50 °C for 4 hours. These tests were performed using a QUV/se Accelerated Weathering Tester, (Q-Panel Lab Products, Cleveland, Ohio, USA), shown in Figure 4.3. After every 2 cycles (24 hours), the weight of the samples were recorded and the sample were rotated 180°C before the next cycle. High temperature wax was applied on the edge of the composite to ensure unidirectional moisture diffusion.
4.5. FIRE RETARDANCY TESTS

The flammability resistance of different types of composite was determined according to ASTM D 635-10. In this setup, a composite bar of 125 ± 5 mm length was placed horizontally, and the free end of the samples were exposed to a standard test flame for 30 s and then flame was removed, as shown in Figure 4.4. The time of flame propagation in between two-fixed position of the composite bar with 75± 1mm distance was recorded. From the recorded time flame propagation rate was measured in mm/min unit. An average of 11 samples were tested for each type of composite.

Figure 4.3: Test chamber used for alternating exposure of UV and condensation

Figure 4.4: Test setup for determining flame propagation rate
4.6. MOISTURE ABSORPTION

Figure 4.5 represents comparison of the moisture absorption behavior in different carpet composites.

![Moisture absorption graph](image)

**Figure 4.5. Moisture absorption behavior of different carpet composites at 80°C.**

Each error bar indicates standard deviation of five specimens. From the curve, it can be inferred that after 1000 hours of moisture absorption, neat epoxy has reached the saturation or equilibrium water content. On the other hand, clay containing sample sets are showing slightly increasing trend at the end of the observed time period. Relative amount of moisture for the neat epoxy composites was about 11%, where as for clay coated and clay infused samples, it was around 8% and 5%, respectively. The maximum moisture absorption for epoxy and nylon are reported as 7% and 9%, respectively. [92, 93] On the other hand, carpet composites being porous absorb more moisture than the individual components.

It has been reported that clay enhances the saturation moisture level due to interface increase in the composite. [94] From the analysis of moisture absorption graph, it is difficult to predict if the clay increases or decreases water saturation level of carpet composites. However, it is seen that the addition of clay has reduced moisture absorption rate, and the absorption rate is
observed in the following decreasing order: 1) neat epoxy, 2) clay coated, and 3) clay infused carpet composite.

For clay infused carpet composites, clay may affect on the overall moisture behavior of the composites in two ways:

1) Reduction in moisture diffusion rate in epoxy matrix
2) Reduction in permeability of epoxy matrix against moisture to reach nylon fibers

Reduction in moisture absorption rate or diffusivity constant by use of clay has been reported by many authors.[94-97] Others reported that clay reduces moisture saturation level due to: 1) bridging of microvoids, II) increased cross-link density around the clay platelets. [95, 97] Reduction in permeability is due to the tortuosity effect clay against water to pass through the matrix. [95] Because of the effect of clay throughout the matrix, clay infused carpet composites absorbed the lowest amount of moisture than the other two types of composites.

In the case of clay coated carpet composites, although the thickness of the coating is negligible compared to the thickness of the whole composite, a significant effect of clay coating in reducing the moisture absorption rate is observed. The neat carpet composite is highly porous. The coating on the composite with a thin layer of clay-epoxy mixture mends the pores, and acts as a clay reinforced diffusion barrier.

Change in mass due to moisture absorption as a function of square root of time is given in Figure 4.6. The error bars indicate ± standard deviation of five specimens. The diffusion of water inside the composites was made unidirectional by blocking transport of moisture through the edges of the composites by coating with high temperature wax. In the plot, the initial straight line indicates the dominance of one directional Fickian diffusion.[98] Based on the Fickian diffusion, it can be assumed that the movement of moisture molecule is through the micro pores only. In this case, swelling, separation of different phases, and residual stress are not involved in the moisture diffusion process. [98, 99]
Figure 4.6: Change in mass in carpet composites as a function of square root of time.

4.7. MECHANICAL PROPERTIES

Table 4.1 lists the flexural strength of different carpet composites before and after moisture absorption. The flexural strength values of as prepared neat epoxy and clay coated composites are almost same. This shows that the thin clay layer coating did not show any effect in the flexural strength. On the other hand, increase in flexural strength in clay infused samples was around 13% compared to the neat epoxy composite. Similar enhancement of flexural strength has been observed in carbon fiber clay/epoxy [97, 100] and glass fiber clay/epoxy [101, 102] hybrid composites. This improvement in flexural strength for clay reinforced hybrid composites is attributed to the enhancement of strength in epoxy matrix and interfacial interaction of fibers and clay.
TABLE 4.1: FLEXURAL STRENGTH OF DIFFERENT CARPET COMPOSITES BEFORE AND AFTER MOISTURE EXPOSURE

<table>
<thead>
<tr>
<th>Sample Type</th>
<th>Flexural Strength (MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>As Prepared Neat Epoxy</td>
<td>27.1± 0.4</td>
</tr>
<tr>
<td>As prepared Clay coated</td>
<td>27.9± 0.2</td>
</tr>
<tr>
<td>As prepared Clay Infused</td>
<td>30.7± 0.9</td>
</tr>
<tr>
<td>Moist Neat Epoxy</td>
<td>20.9± 1.0</td>
</tr>
<tr>
<td>Moist Clay Coated</td>
<td>22.7± 1.0</td>
</tr>
<tr>
<td>Moist Clay Infused</td>
<td>25.9± 1.1</td>
</tr>
</tbody>
</table>

Significant reduction of flexural strength of the composites has been noticed after the moisture absorption in comparison with the as prepared samples. For neat epoxy composite, reduction in flexural strength was 22.7%. On the other hand, reduction in strength was 18.7% and 15.7%, respectively for clay coated and clay infused composites. Reduction in flexural strength can be due to chemical degradation of both fiber and epoxy matrix, and also due to reduction in interfacial bonding between fibers and epoxy. [103] Reduction of flexural strength was lowest in clay-infused composites. It can be assumed that higher barrier property of the clay-epoxy matrix against water has slowed down the degradation process of nylon and fiber matrix interface. At the same time, presence of clay in epoxy matrix has maintained better mechanical properties of epoxy matrix after the degradation process.

Table 4.2 shows the flexural modulus of different carpet composites before and after moisture absorption. As prepared clay infused composites showed 40.8% increase in flexural modulus in comparison with that of the neat epoxy samples. Increasing flexural modulus in clay-epoxy nanocomposite and for clay-epoxy- fiber composites after moisture absorption have been reported by many authors. The increase in the flexural modulus is attributed to the motion restriction of the polymer chains by the clay platelets, and increase in the cross linking density of polymer around the clay platelets. Another possible explanation for modulus enhancement could be the increase in interfacial bonding between matrix and fiber by clay platelets.
**TABLE 4.2: FLEXURAL MODULUS OF CARPET COMPOSITES BEFORE AND AFTER MOISTURE ABSORPTION**

<table>
<thead>
<tr>
<th>Sample Type</th>
<th>Flexural Modulus (MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>As prepared Neat epoxy</td>
<td>1842± 97</td>
</tr>
<tr>
<td>As prepared Clay coated</td>
<td>1878± 75</td>
</tr>
<tr>
<td>As prepared Clay infused</td>
<td>2594± 64</td>
</tr>
<tr>
<td>Moist Neat Epoxy</td>
<td>898± 45</td>
</tr>
<tr>
<td>Moist Clay Coated</td>
<td>1107± 44</td>
</tr>
<tr>
<td>Moist Clay Infused</td>
<td>1393± 167</td>
</tr>
</tbody>
</table>

Reduction in elastic modulus due to moisture absorption is observed (Table 4.2) in carpet composites. Reduction in modulus is directly related to scission of polymer chain. [102] For neat epoxy carpet composites, reduction was 51.2%, while it was 41.1% and 46.32%, respectively for clay coated and clay infused composites. The improvement in modulus for moisture absorbed clay composites compared to neat composites is attributed to the less scission of polymer chains as a result of low moisture diffusion due to the presence of clay platelets.

**4.8. CYCLIC EXPOSURE OF UV AND CONDENSATION**

Variation in mass in neat and clay carpet composites due to the cyclic exposure to Ultra Violet (UV) and condensation is shown in Figure 4.7. It is observed that during 1000 hours alternative exposure, neat epoxy composites gained more weight than clay composites. Among the clay composites, clay infused composites showed more weight change than clay-coated composites. The reason is attributed to the number of voids present on the surface of composites. The neat composites gained more weight because of the presence of numerous voids. These voids are minimum for clay-coated composites as a result of thin coating of clay-epoxy on the composites’ surface. The voids are not closed for infused carpet composite, but the weight gain is lower than neat composite because moisture diffusion is hindered due to tortuosity by clay platelets. No erosion or decrease of weight has been noticed during exposure of 1000 hours.
The flexural strength of different carpet composites before and after cyclic exposure to UV and condensation is shown in Table 4.3. The results show an unexpected increase in flexural strength after alternating environmental exposure. The possible explanation for this could be an increase in the cross linking density in epoxy matrix due to the high temperature in the degradation chamber. The degradation effect on the mechanical properties as a result of cyclic exposure is not observed. This is due to the high thickness of the carpet composite, and 1000 hr. exposure time is perhaps not sufficient to significantly change the mechanical properties as a result of degradation. UV degradation is a surface-controlled phenomenon. Liau et al. exposed glass fiber composite in UV and effect of UV was not found for the composite with more than one lamina after 180 days exposure. [104] Kumar et al. used sample thickness of 1.27 mm of epoxy carbon fiber composites for UV and cyclic exposure to observe the change in properties, and as a result, found some decrease in the mechanical properties. [105] Allen et al. used nylon test samples of 100-μm thickness to observe the significant effect of exposure. [107] In the case of carpet composite, 14 mm thick composites were used for exposure. Due to the high thickness of the composite, and less exposure time, the UV exposure effect is limited only the surface layer. As a result, significant degradation is not observed in the strength of the carpet composites.
TABLE 4.3: FLEXURAL STRENGTH OF CARPET COMPOSITES BEFORE AND AFTER EXPOSURE TO UV AND CONDENSATION

<table>
<thead>
<tr>
<th>Sample Type</th>
<th>Flexural Strength (MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>As Prepared Neat Epoxy</td>
<td>27.07 ± 0.4</td>
</tr>
<tr>
<td>As Prepared Clay Coated</td>
<td>27.9 ± 0.2</td>
</tr>
<tr>
<td>As Prepared Clay Infused</td>
<td>30.7 ± 0.9</td>
</tr>
<tr>
<td>Degraded Neat Epoxy</td>
<td>37.4 ± 8.6</td>
</tr>
<tr>
<td>Degraded Clay Coated</td>
<td>43.5 ± 1.1</td>
</tr>
<tr>
<td>Degraded Clay Infused</td>
<td>36.9 ± 2.0</td>
</tr>
</tbody>
</table>

It is seen that clay coated composites possess the highest strength after alternating degradation in comparison to other types of carpet composites. This is attributed to the low absorption of water and hence low degradation of the clay coated composites. Clay acted as a strong barrier against moisture penetration, and protected the composite panels degrading from UV radiation and condensation.

The flexural modulus of composites before and after cyclic UV and condensation is given in Table 4.4. The neat and clay-coated composites showed an increase in flexural modulus after alternating environmental exposure. Higher cross-linking density restricts molecular movements. Increase in flexural modulus after alternating degradation indicates higher cross-linking density of epoxy matrix during the degradation process. But, the flexural modulus of clay infused composites decreased after cyclic exposure due to unknown reason.
**TABLE 4.4:** FLEXURAL MODULUS OF CARPET COMPOSITES BEFORE AND AFTER EXPOSURE TO UV AND CONDENSATION

<table>
<thead>
<tr>
<th>Sample Type</th>
<th>Flexural Modulus (MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>As Prepared Neat Epoxy</td>
<td>1842± 97</td>
</tr>
<tr>
<td>As Prepared Clay Coated</td>
<td>1878± 75</td>
</tr>
<tr>
<td>As Prepared Clay Infused</td>
<td>2594± 64</td>
</tr>
<tr>
<td>Degraded Neat Epoxy</td>
<td>2032± 214</td>
</tr>
<tr>
<td>Degraded Clay Coated</td>
<td>2183± 41</td>
</tr>
<tr>
<td>Degraded Clay Infused</td>
<td>2173± 40</td>
</tr>
</tbody>
</table>

**4.9. FIRE TEST**

Figure 4.8 shows the comparison of the linear flame propagation rate for different carpet composites. The error bar shows ± standard deviation of eleven specimens. 20% and 34% reduction in the average linear burning rate for clay-coated and clay infused samples, respectively are observed when compared to neat composites.

Clay coated samples have clay only on the surface and it can be assumed that clay-epoxy coating delayed the ignition time to initiate the flame propagation. The formation of initial flame is important in flame propagation in absence of any other external heat source. Delayed ignition time can be attributed to the thermal stability enhancement of epoxy/ clay coating composites. [71] Among all the prepared composites, lowest flame propagation rate is observed in clay infused composite. This indicates that the presence of clay platelets throughout the matrix in the clay infused composite reduces the flame propagation in the bulk sample. The result is in accordance with the reported observation of reduction in flammability of clay-polymer compared to neat polymer. [53] Nano particles, like clay, when present in the polymer matrix help in forming network like structure against decomposed flammable product and hinder the burning process.
4.10. WIND LOAD TESTS

Wind load tests were carried out to check the wind load bearing capacity of carpet composites using a model as shown below in Figure 4.9 and Figure 4.10. The static wind load test was conducted by subjecting the samples to a uniformly distributed load. The supports were designed to have one support as pin support while the other support was a roller support. As air flows in the airbag it pushes the sample upward with uniformly distributed pressure. According to the pressure and surface area of the P-delta (force vs deflection) diagram can be plotted as shown below in Figure 9. The tests were conducted on a sample of size 13.9573 cm x 59.055 cm and an average thickness of 1.9 cm. It was observed that the samples could hold 448.11 kPa air pressure, without rupture. As seen below in Figure 4.11, the deflection varies linearly to a
pressure of 150 kPa, at this point the material cracks and is exposed to the carpet fibers inside the composite. However the wind load on encountering the perfect fiber matrix combination at 150 kPa cannot break the composite at this pressure. The airbag pressure of 448.11 kPa is required to break the carpet composite, thereby indicating its superior strength. The carpet composite can hold a load of 199.45 kg/cm for a deflection of 10 cm as shown in the Load curve plot below in Figure 4.12.

Figure 4.9. A design model for wind load test
Figure 4.10. A picture taken during wind load test

Figure 4.11. Pressure Deflection wind load plot of a BTTB Epoxy carpet composite
Figure 4.12. Wind Load curve plot of a BTTB Epoxy carpet composite
5. ACOUSTIC PROPERTIES OF RECYCLED CARPET COMPOSITES

5.1 OBJECTIVE

The noise absorption coefficients of carpet composites were measured using the impedance tube method (ASTM C 384-04) to evaluate their feasibility as highway noise barriers.

5.2 BACKGROUND

Sound waves are longitudinal waves which require a medium for travelling, like air or water, and are measured in cycles per second or Hertz (Hz). The human ear can detect a wide range of frequencies from 20 to 20,000 Hz, but it is most sensitive to the sounds over a frequency range of 200 to 5000 Hz. The human ear does not respond in a uniform manner to different frequency sounds – a sound pressure level of 70 dB is perceived as much louder at 1000 Hz than at 100 Hz. The absorption of sound results from dissipation of sound energy as heat and this dissipation mechanism is mainly due to one of two phenomena:

1) Energy loss due to flexural vibrations in the specimen [41].

2) Porosity effects, where the energy is dissipated due to multiple reflections of sound waves within the voids in the structure [41].

For porous materials like sponge and mineral wool which have interconnected pores the incoming sound waves are reflected within the pores, causing them to vibrate and therefore convert sound energy into heat [41]. This is also true in the case of composites as concluded in the study by Hargrove that, “the mechanism that causes composites to reduce noise has been shown to be the reverberations of sound waves against the sides of many voids in the composite. This causes a transformation of acoustic to heat energy” [109]. The nature of composites is a multitude of minute voids in the material. Sound waves reverberate from wall to wall of these voids and are attenuated by the conversion of a large amount of noise to a relatively small amount of heat [91]. Also for a completely porous material, sound is lost isothermally near the walls of the voids and to a lesser extent in or near the center of the hollow spaces [109].

For homogeneous and isotropic materials, acoustic performance is defined by a set of five experimentally determined constants: noise absorption coefficient (NAC), pressure reflection
coefficient (PRC), acoustic impedance, propagation constant and noise reduction coefficient (NRC) [41].

5.3 THEORY

The ability of a highway noise barrier to reduce noise levels depends upon a number of variables including the geometric factors like relative heights and distances of the noise source and receiver to the barrier, and the mass per unit area (surface density) of a wall type barrier [35]. There are two ways to characterize the acoustic properties of a material – by measuring its absorption coefficient or by measuring its transmission loss.

The noise absorption coefficient (NAC) is the most critical parameter in acoustic characterization of a material to determine how well it can absorb sound or noise. When a sound wave hits a material, a portion of it will be reflected or “bounced back.” In simple terms, the lesser the sound energy reflected, the better the noise reduction property of the material [33]. A value of NAC of 0 means the sound energy is being reflected completely, and a value of NAC of 1 means that all the sound energy is being absorbed, which is the best in noise reduction. The American Society of Testing and Materials (ASTM) issued a standard, ASTM C 384 -04, on how to conduct the test of noise absorption coefficient of a material.

In this experiment, sound waves are created with the use of a speaker connected to a function generator. The function generator sends an oscillating current signal to the speaker causing the speaker’s diaphragm to vibrate. As the diaphragm moves outward or toward the sample, the air near the speaker is compressed creating a small volume at relatively high pressure, which propagates away from the speaker. As the diaphragm moves inward, away from the sample, it creates a low-pressure area that propagates away from the speaker. As shown in the figure below in Fig. 5.1, an alternating signal into a speaker creates sound waves that propagate away from the speaker. The dark bands represent areas of high pressure while the light bands represent areas of low pressure [110].
The process of compressions and rarefactions continues with a frequency equivalent to the input signal. The higher the input frequency, the more often the compression/rarefaction cycle would occur at each second. When sound waves move away from the speaker and are detected by the microphone, the entire cycle is reversed: the areas of high and low pressure cause the microphone’s diaphragm to vibrate in both directions, and this creates small electrical signals [110]. This electrical signal from the microphone can be recorded by a recording device or displayed on an oscilloscope [110]. In the method specified by ASTM C 384, “A plane wave traveling in one direction down a tube is reflected back by the test specimen to produce a standing wave that can be explored with a microphone” [111]. The normal incidence sound absorption (NAC), which is a function of the frequency, can be determined from the standing wave ratio at the face of the test specimen. Measurements were made with pure tones at a number of frequencies as described by the third octave band frequency specified in ANSI S1.6 [111].

5.4 SCOPE, STANDARD AND SIGNIFICANCE

ASTM C 384 – 04 mentions in its scope that, “This test method covers the use of an impedance tube, alternatively called a standing wave apparatus for the measurement of impedance ratios and normal incidence sound absorption coefficients of acoustical materials” [111]. The measurements specified in this test are significant because it is useful in basic research and product development of sound absorptive materials [111].

The sound absorbing performance of the material is the ratio of un-reflected sound intensity at the surface to the incident sound intensity, and is known as Noise absorption coefficient (NAC). Other important acoustic parameter include the pressure reflection coefficient
(PRC) which is a ratio of total reflected pressure (sound intensity) to the total incident pressure at the face of the specimen [111]. NAC and PRC vary with frequency and are also a function of material thickness, density and pore sizes.

The standing wave apparatus used for the testing of ASTM C 384 is a simple and less complex method which yields the noise absorption coefficient of the material being tested. The test method can be applied for measuring NAC at normal incidence that is $0^\circ$. NAC calculated by ASTM C 384 is useful in basic research and product development of sound absorptive materials like carpet composites. This method involves the use of impedance tube (with two microphones), a digital oscilloscope, and measurement of complex pressures using the microphones, from the source speaker which has an input from the frequency generator as shown in Fig. 5.2 below.

![Figure 5.2 A schematic diagram to show the apparatus used for acoustic test based on ASTM C-384 to obtain noise absorption coefficient](image)

The approach adopted for this study was to build an acoustic set up rather than purchasing an integrated acoustic set up which is expensive. The tube used as an impedance tube was a rectangular aluminum tube with polished inner ends. The tube is straight 1m in length to ensure that the plane waves are fully developed. The rectangular tube had a cross-section of $0.0762 \text{ m} \times 0.0762 \text{ m}$. The general formula for rectangular cross section as per the ASTM
The standard is as mentioned in the following equation 5.1.

\[ d = \frac{4 \times \text{Area}}{\text{Perimeter}} \]  

(5.1)

The tube should be maintained free of dust to maintain low tube attenuation. Tube attenuation is calculated using the following equation 5.2. Kirchoff developed and Beranek modified the formula for estimating the attenuation constant as given below,

\[ A = \frac{0.02203 f^{0.5}}{C \times d} \]  

(5.2)

Where, \( f \) = frequency in Hz.

\( C \) = velocity of sound in m/s.

\( d \) = diameter (length of one side of the rectangle) in m.

The wavelength of the sound transmitted from the frequency generator is developed using the following equation 5.3.

\[ W = \frac{C}{f} \]  

(5.3)

The velocity of sound is in turn, dependent on the room temperature \( T \) in which the experiment has been conducted and is given by the following equation 4.4.

\[ C = 20.05 (T + 273.1)^{0.5} \]  

(5.4)

Hence the velocity of sound was found to be stable at 345.76 m/s during the experiment process because of a steady atmospheric temperature maintained at 24.24°C.

The microphone is placed at regular intervals of 0.1 m inside the impedance tube with a marking of 0.1 m to 0.9 m on the aluminum tube. A correction factor was used to analyze the distances was used as shown in the following equation 4.5.

\[ x_{cor} = x_{1/4} - x_{mr} - \frac{W}{4} \]  

(5.5)

Where,

\( x_{cor} \) = Correction factor for distance, m.

\( x_{1/4} \) = 0.1 m = observed scale reading with microphone probe at the first minimum, m.

\( x_{mr} \) = 0.005 m = observed scale reading with probe touching face of metal backing plate, m.

The correction factor for distance \( x_{cor} \) was found to be 0.078 m.

This was incorporated in the routine measurements by the following equation 5.6 used to subtract correction factor.
\[ x = (x_{\text{obs}} - x_{sf}) \]  
\[ \text{cor} \]  
\[ (5.6) \]

Where,

\( x \) = true distance from the specimen surface, m

\( x_{\text{obs}} \) = observed scale reading, m, and

\( x_{sf} \) = observed scale reading with probe touching specimen face, m.

For the standing wave within the tube it is necessary to know the distance from the sample face at which each pressure is being measured. The exact location of the mounted sample within the tube can be determined by gently advancing the probe until it makes contact with the sample face and noting the scale reading of the point of contact [93]. The exact location however requires a correction factor applied to the observed scale reading because the acoustic center of the microphone or the microphone probe does not necessarily correspond to its geometric center [93].

The microphone voltage at its minimum voltage should be at least 10 times more than the voltage of the background noise voltage. This would ensure that the noise level would be at least 10 dB more than the noise level of background noise. This can be observed from the following image taken from oscilloscope reading while performing the acoustic test. The sound waves are generated at various frequencies by placing the microphone at a particular distance from the specimen. This can be seen in Fig. 5.3 where the frequency generator was set to produce a sound of 500 Hz and the microphone was placed at the specimen face. The peak to peak voltage was noted at every frequency at different distances from the specimen to obtain a standing wave. Fig 5.3 shows that the noise voltage in green – no.4 is less compared to the microphone voltage observed in the wave no. 3 in purple.

![Figure 5.3 An image of the oscilloscope reading](image-url)
The standing wave ratio (SWR) at the specimen face is calculated using the extrapolation method by measuring the voltages (corresponding pressure levels) at various frequencies and at various distances.

The standing wave ratio at specimen face is measured using the following three cases -

a) **When two or more minima are present in the sine wave obtained**

\[ V_{\text{min}}(0) = V(x_1) - x_1 \left( \frac{V(x_2) - V(x_1)}{x_2 - x_1} \right) \]  

(5.7)

b) **One minimum and one maximum present**

In this case the maximum voltage at the sample face \( V_{\text{max}}(0) \). In this case there is only one minimum and a graphical extrapolation back to the specimen face cannot be used. A valid approximation can be given by the following equation.

\[ V_{\text{min}}(0) = V(x_1) - A \times x_1 \times V_{\text{max}}(0) \]  

(5.8)

where, \( A \) can be calculated from equation B.2.

c) **Only one minimum and no maximum present**

When there is no actual maximum measured in the tube, a maximum level can be inferred from a measurement of the sound pressure level at W/8 distance on either side of the minimum as per the ASTM C - 384 Standard. The following equation B.9 is suggested in the standard for this kind of situations.

\[ V_{\text{max}} = \left( 2 \times V^2_{W/8} - V_{\text{min}}^2 \right)^{\frac{1}{2}} \]  

(5.9)

Thus when no maximum and only one minimum can be measured, an additional voltage measurement at a distance of W/8 from the measured minimum should be taken and used as \( V_{W/8} \) in B.9 to arrive at an estimated \( V_{\text{max}} \) value.

Hence using any of the above three conditions the values can be substituted to calculate SWR (0) at the specimen face.

The impedance tube is used to generate a standing wave in the tube which is enclosed with a speaker at one end and a specimen on the other end. The interference of an incident wave and reflected wave from the specimen creates a standing wave in the impedance tube, which is measured as standing wave ratio (SWR). SWR is a function of the distance between the speaker and the microphone (\( x \)), and is given as,

\[ \text{SWR}(x) = \frac{V_{\text{max}}(x)}{V_{\text{min}}(x)} \]  

(5.10)
Where \( \text{SWR}(x) = \) standing wave ratio at location \( x \), a dimensionless, positive, real number greater than or equal to 1.

In this experiment voltage has been correlated to the change in pressure level and hence the standing wave ratio mathematically indicates,

\[
\text{SWR}(x) = \frac{p_{\max}(x)}{p_{\min}(x)}
\]  
(5.11)

Where \( p_{\max}, p_{\min} \) are the maximum and minimum sound pressures in the impedance tube.

The ASTM C - 384 states that, “The primary purpose for making the measurements described in this test method is to find the standing wave ratio at the face of the specimen, that is, SWR (0).” SWR (0) is determined indirectly by extrapolation of the maximum and minimum voltages actually measured in the tube and its value is critical in this experiment.

The pressure reflection coefficient can be calculated from SWR (0) by the following equation:

\[
P_{\text{RC}} = \frac{\text{SWR}(0)-1}{\text{SWR}(0)+1}
\]  
(5.12)

The noise absorption coefficient is a dimensionless, real number which is directly related to the pressure reflection coefficient by the following equation:

\[
N_{\text{AC}} = 1 - |P_{\text{RC}}|^2
\]  
(5.13)

### 5.5 MATERIAL CHARACTERISTICS

The 11 materials tested for acoustic characteristics and their surface properties are mentioned in Table 5.1. These surface properties were measured using the weigh balance and ruler because of their dimensional stability.

Amongst all the carpet composites the 4 layer carpet composite has the highest specific gravity (SG) of 12.07 while the PE carpet closely followed this number with a specific gravity of 10. The BTTB carpet composite had an SG of 3.611 making it the lowest amidst all the carpet composites tested for acoustic characterization. The TBBT carpet composite had an SG of 6.667 while the carpet with clay configuration that has addition of clay layers in a TBBT carpet composite has an SG of 7.143, indicating just a marginal improvement in SG. This indicates that the use of clay as nanofiller increases the specific gravity of the composite.

“PS” foam was the lightest material tested with an SG of 0.4167 and concrete on the other hand has the highest SG of 24.61.
The choice of the 12 materials was not restrained by thickness or composition as ASTM C 384 has no specifications about thickness and compositions of the sample.

**TABLE 5.1 A TABLE OF 12 SOUND ABSORBING MATERIALS AND THEIR SURFACE PROPERTIES**

<table>
<thead>
<tr>
<th>Sr. No.</th>
<th>Materials</th>
<th>Mass (kg)</th>
<th>Volume (m³)</th>
<th>Specific Gravity</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Virgin Carpet</td>
<td>0.08</td>
<td>0.52</td>
<td>1.54</td>
</tr>
<tr>
<td>2</td>
<td>Virgin Carpet BTTB</td>
<td>0.08</td>
<td>0.52</td>
<td>1.54</td>
</tr>
<tr>
<td>3</td>
<td>Epoxy Matrix</td>
<td>0.21</td>
<td>0.12</td>
<td>1.75</td>
</tr>
<tr>
<td>4</td>
<td>TBBT Carpet Composite</td>
<td>0.24</td>
<td>0.36</td>
<td>0.67</td>
</tr>
<tr>
<td>5</td>
<td>BTTB Carpet Composite</td>
<td>0.13</td>
<td>0.36</td>
<td>0.36</td>
</tr>
<tr>
<td>6</td>
<td>4 layer Carpet Composite</td>
<td>0.35</td>
<td>0.29</td>
<td>1.21</td>
</tr>
<tr>
<td>7</td>
<td>VE Carpet</td>
<td>0.11</td>
<td>0.17</td>
<td>0.65</td>
</tr>
<tr>
<td>8</td>
<td>PE Carpet</td>
<td>0.17</td>
<td>0.17</td>
<td>1.00</td>
</tr>
<tr>
<td>9</td>
<td>Concrete</td>
<td>0.96</td>
<td>0.39</td>
<td>2.46</td>
</tr>
<tr>
<td>10</td>
<td>Wood</td>
<td>0.14</td>
<td>0.19</td>
<td>0.73</td>
</tr>
<tr>
<td>11</td>
<td>Carpet with Clay</td>
<td>0.15</td>
<td>0.21</td>
<td>0.71</td>
</tr>
<tr>
<td>12</td>
<td>PS Foam</td>
<td>0.01</td>
<td>0.24</td>
<td>0.04</td>
</tr>
</tbody>
</table>
5.6 RESULTS

The NAC measurements were conducted using the ASTM C - 384 standards at the 3rd octave band frequency range from 125 Hz to 4000 Hz. The 3rd octave band frequencies chosen for this test were - 125 Hz, 250 Hz, 500 Hz, 1000 Hz, 2000 Hz and 4000 Hz.

Fig. 5.4 shows the result obtained for 11 different materials tested for noise absorption coefficient (NAC) using ASTM C 384-04 and it can be concluded from the result that the virgin carpet (which means carpet without any resin treatment or usage) has the highest noise absorption at low frequencies of 125 Hz but drops down in absorption at frequency range of 250 – 500 Hz. Polystyrene (“PS”) foam is the best absorber material among all the other materials compared in the frequency range of 250 – 500 Hz. However it can also be seen that PS foam gives up at frequency above 500 Hz that is, in the higher frequency range of 1000 Hz to 4000 Hz it performs poorly.

The following observations were made based on the results displayed in Fig. 5.4 for NAC obtained at different frequencies -

- The four layer carpet composite performs exceptionally well in the frequency range from 1000 Hz to 4000 Hz. This proves that among all the materials the best material best suited for a highway noise barrier is the four layer carpet composite material. The material is robust in its noise absorption in the low frequency range of 125 Hz to 500 Hz. This is because it is only third to virgin carpet and PS foam in the low frequency range.

- Virgin carpet BTTB was tested to check if the properties had any difference in properties in comparison to the other virgin carpet sample which has a configuration of TBBT. Results from Fig. 5.4 indicate that probably the backing surface of the carpet is very weak due to lesser porosity in comparison to the carpet fibers. A porous medium can be looked at as a mixture of two phases, air and solid material, both of which react differently to the sound wave. Hence there is a higher NAC indicating greater noise absorption in porous materials like Virgin carpet than in the flat Virgin carpet BTTB.
The material that performed the poorest among the 11 materials is concrete. Concrete has a steady absorption rate of around 0.125 at all frequencies that is significantly low.

There was a steady trend observed at 500 Hz where all the materials displayed high sound absorption. A similar trend was observed for a dip in acoustic absorption for all materials at 1000 Hz. This could probably be because the frequencies could be harmonic frequency of the materials.

It was necessary to understand the behavior of different carpet composites and the effect of different resin systems; layup arrangements that is, TBBT, BTTB, 4 layer; addition of nanocomposite fillers and other general comparisons on the acoustic performance of the composites. Hence a comparison of 6 different carpet composites has been presented in Fig. 5.5 from which the following observations were made -
- The 4 layer carpet composite is the best material among all carpet composite configurations tested. The 4 layer carpet composite performs exceptionally well for high frequency range sound from the range of 1000 Hz to 4000 Hz. This is the range of noise created by the friction of tires on highway roads. The low frequency noise is created by the engine noise of the cars and vehicles. It can be observed that in general the carpet composites do a good job of absorbing the noise at both low and high frequencies. However it is the 4 layer carpet composite that has the highest noise absorption at the high frequency range qualifying it for a product as a highway noise barrier.

**Figure 5.5 Comparison of Noise Absorption coefficients different carpet composite materials calculated at the third octave band range of frequencies using ASTM C 384 -04**

- Addition of clay to the composites has a marginal effect on the noise absorption properties of the carpet composites. Although it is difficult to arrive at conclusion considering the thin margin of difference in the acoustic performance of composites but the comparison suggests that addition of clay has no impact on acoustic performance of carpet composites.
In an attempt to highlight the difference between the proposed highway noise barrier material, the current highway noise barrier material that is, concrete was tested and used in this chart for a direct comparison. Fig. 5.5 clearly highlights the poor noise absorption qualities of concrete in comparison to the carpet composites. Also an interesting observation is the difference in the pattern in which the NAC varies with respect to a change in the frequencies. The NAC variation in concrete is completely different from other carpet composites. For instance the carpet composites have a huge reduction in NAC at 1000 Hz which suddenly increases in 2000 Hz, while this is not observed in concrete. This is interesting in the material science perspective and therefore it is necessary to compare the properties with the parent material, as shown in Fig. 5.6.

A comparison of noise absorption coefficients of carpet composites with its parent materials - virgin carpet and the epoxy matrix is as shown in Fig. 5.6. The following observations were made from the Fig. 5.6.

- It can be observed that the 4 layer carpet composite performs better than the virgin carpet material and the epoxy matrix at higher frequencies, in the range of 1000 Hz to 4000 Hz, which is the most critical sound frequency range for a highway noise barrier.
- It can be observed that the 4 layer carpet composite tries its best to mimic the superior noise absorption properties of virgin carpet.
- The 4 layer carpet composite behave closely to the nature of variation of NAC for carpet and epoxy matrix at lower frequency range. But at higher frequency range it behaves like virgin carpet, rather with higher NAC. The epoxy matrix in itself is poor in noise absorption at higher frequencies like 1000 Hz and 2000 Hz and therefore has minimal influence on the behavior of 4 layer carpet composite NAC variation.

A similar observation can be made for BTTB carpet composite which mimics the NAC behavior of Epoxy matrix over the range of frequencies from 125 Hz to 4000 Hz. This has been presented in Fig. 5.7 to give a comparison of NAC’s of epoxy matrix and BTTB carpet composite.

- The acoustic performance of BTTB Carpet composite is lesser than epoxy matrix in the third octave band range of frequencies but the nature of the plot is similar. It can be
clearly seen that the BTTB carpet composite mimics the NAC behavior of Epoxy matrix at both low and high frequency ranges.

Figure 5.6 Comparison of Noise Absorption coefficients of 4 layer carpet composite with its parent material calculated at the third octave band range of frequencies using ASTM C 384-04.
BTTB carpet composites have the poorest acoustic performance in comparison to the acoustic performance of all the carpet composites. Because BTTB arrangement has Bottom - Top - Top - Bottom, type of arrangement the reinforcement for BTTB carpet composites is provided by the Polypropylene backing of the carpet instead of the nylon fibers. The Polypropylene backing is the flat end of the carpet and therefore has no obstruction for the resin flow during VARTM. This could be a potential reason for the epoxy matrix to play a dominant role in influencing the acoustic performance of the carpet composites instead of the virgin carpet. Hence the BTTB carpet composites mimic the NAC variation behavior of epoxy matrix composites.

Therefore it is safe to conclude that amongst all the carpet composite configurations, it is safe to avoid BTTB carpet composite, because it would not make good highway noise barriers due to poor noise absorption characteristics.
Hence from the test it can be concluded that the noise absorption coefficient of proposed highway noise barrier that is, carpet composites is high in comparison to the currently used concrete highway noise barriers. Fig. 5.8 presents a comparison of noise absorption between the 4 layer carpet composite (the best noise absorption material) and concrete (the worst noise absorption material). The superior noise absorption characteristic of the 4 layer carpet composite is evident from Fig. 5.8.

Figure 5.8 Acoustic property comparison of concrete and 4 layer carpet composite

NAC gives a detailed comparison of the noise absorption of the material over a broad range of frequencies. However there is a need for a parameter, which helps in direct comparison of noise absorption of all the materials. Noise Reduction Coefficient (NRC) represents a single number, which is the average of value of all the absorption coefficients of the material at the frequencies 125, 250, 500, 1000, 2000 and 4000 Hz. NRC was calculated for all the materials and the plot in Fig. 5.9 illustrates the difference in NRC’s for all the 12 materials tested for noise absorption.
The following observations can be made on the basis of analysis of the values of NRC from Fig. 5.9 -

- It can be seen that the NRC of 4 layer carpet composite (best absorbing material) is the highest and the NRC of concrete (worst absorbing material) is the lowest.

- The NRC of BTTB carpet composite is close to the NRC value of epoxy matrix and therefore confirms the influence of epoxy matrix in determination of noise absorption characteristics of BTTB carpet composite. This also indicates that there is no significant acoustic contribution by the reinforcement which has occurred in the BTTB composite formed by VARTM of carpet in BTTB lay - up with epoxy matrix as the thermoset matrix.

- The NRC of virgin carpet is the highest amongst all the 12 materials tested for noise absorption. But poor mechanical properties of the virgin carpet would disqualify it as a good highway noise barrier. Hence the 4 layer carpet composite which is only next to
virgin carpet with a high NRC value of 0.72 can be proposed as a good highway noise barrier.

- The carpet composites have good noise absorption properties compared to wood and concrete which are the traditional highway noise barrier materials.

An attempt was made to understand the correlation between the specific gravity of the material and its acoustic properties. It can be roughly said that the acoustic properties are better for a material with higher specific gravity. This can be inferred from Fig. 5.10 below, where the 4 layer carpet composite material with highest specific gravity of 12.07 also had the highest NRC of 0.72 among all the carpet composites tested for acoustic characterization.

![Graph showing correlation between NRC and SG]

**Figure 5.10** Correlation of Noise Reduction Coefficients (NRC) with Specific Gravity (SG)

The following observations can be made from Fig. 5.10 -

- A material with higher specific gravity would have lower noise reduction coefficient. This can be seen on a comparison of high specific gravity material like Concrete with a
low specific gravity material like PS Foam, where PS Foam yielded better noise reduction characteristics in comparison to concrete.

- Carpet composites have lower specific gravity and therefore have higher noise reduction coefficient in comparison to materials like concrete and wood. Hence it can be concluded that carpet composites have better noise reduction capabilities than concrete or wood.
6. CONCLUSIONS AND OTHER SIGNIFICANT RESULTS

The acoustic results for six different set of carpet composites indicated excellent absorption properties of the carpet composites and therefore it could have a potential application as highway noise barriers. The contributions of this work include:

1) Scaling up of VARTM process for fabrication of large-scale carpet composites. The capabilities of the scale up include:
   1. Feasibility of making large composites of size more than 0.61 m x 0.61 m with uniform thickness.
   2. Fabrication of curved shaped panels by the scale up of VARTM process.
   3. One of the powerful features of the process scale-up of VARTM practiced is the exclusion of hot press used in VARTM process. Hot press restricts the size and shape of the final part of the composite and by avoiding it, the scaled up the process.

2) Product scale up included the addition of PVP-clay 3% nanocomposite films in the carpet composite by the scaled up VARTM process. The clay nanofiller can act as a fire retardant and probably prevent the carpet composites from undergoing environmental degradation by UV and moisture. The PVP Clay 3% nanocomposite film used as filler was a well-exfoliated film.

3) Application of the carpet composites prepared by scaled up VARTM process as Highway Noise Barriers:
   a. It was demonstrated that Carpet composites have excellent absorption properties with a noise absorption coefficient in the range of 0.70.
   b. Addition of clay to the carpet composites did not make a significant difference in the noise absorption capabilities of the carpet composites.
   c. Changing the resin system did not make any significant improvement of decline in the noise absorption of carpet composites.
   d. The BTTB carpet composites have an acoustic performance similar to the epoxy matrix – which indicates that the noise absorption properties of carpet have no contribution in this type of configuration of composites.
e. The acoustic test results indicated that the carpet composites could have an acoustic performance better than the epoxy matrix, but they do not seem to match the sound absorption capabilities of virgin carpet material. However on the basis of the experiments, the performance of the carpet composites is estimated to be better than both the virgin carpet and the epoxy matrix at higher frequencies.

f. On the basis of the acoustic measurements of concrete and other materials it can be estimated that the carpet composites can absorb more sound than concrete. Hence there is a possibility for carpet composites as alternative highway noise barrier material.

For evaluating the effect of the environment and fire, composites were fabricated with neat epoxy, and further coated with a mix of epoxy and I.28E nanoclay. Composite samples were also fabricated by infusing them with a mixture of epoxy and 4 % I.28E nanoclay. Composites were then exposed to moisture and cycling exposure of ultra violet radiation and condensed vapor. Flexural properties were tested before and after each degradation process.

Significant reduction in flammability in the carpet composites was observed due to use of clay. The linear burning rate as per ASTM D 635-10 was reduced by 20 % for composite samples coated with the epoxy-clay coating, while it was reduced by 34 % for composite samples infused with 4 % nanoclay.

Moisture exposure for these samples indicated that both clay coating and clay infusion reduced the deleterious effect of moisture on the composites. Significant reduction in diffusivity was found in clay coated and clay infused composites. All composites showed a reduction in flexural strength and modulus, where severity of degradation was less in clay coated and clay infused carpet composites. After 1000 hours exposure of moisture clay infused carpet composite demonstrated higher strength and modulus than other other two composites. It was also observed that moisture absorption was a Fickian diffusion mechanism.

For alternating moisture and UV exposure cycles, the clay-coated samples exhibited the lowest change in mass, while the neat epoxy recycled carpet composites showed the highest mass gain. However, It was also observed that the combined moisture/UV exposure resulted in an increase in strength in the case of all composite types. It can assumed that high temperature exposure of composites will lead to higher cross linking of the polymer. However, the exposure time for cyclic environmental condition was not long enough to observe the effect of
degradation. Longer exposure and molecular analysis can be conducted to determine the effect of UV on the carpet composite.

Finally, wind load tests proved that the composite is strong and withstands extreme wind conditions similar to tornadoes.

Three graduate students have completed their Masters work on this funded project. A patent based on this work was applied for during the project and has recently been issued – A. K. Singh, G. Pandey, R. Vaidyanathan, and R. P. Singh, “Fabrication Method of Multi-functional Composites from Pre/Post-Customer Carpet,” U.S. Patent Application No. 12/695730, August 5, 2010. An Oklahoma manufacturer (K. T. Plastics, Calera, OK) has signed an option to license for this patent. It is expected that KT Plastics will be providing the technology to another Oklahoma manufacturer of highway sound barriers (Apache Precast, Jenks, OK).

Several papers have been published and presented, while also giving the opportunity to intern for undergraduate students at Oklahoma State University.
REFERENCES


