ME 450
Winter 2009

Project 4: Fine Needle Aspiration

Final Report
April 21, 2009

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EXECUTIVE SUMMARY

Design Problem
Current fine needle aspiration (FNA) devices have a “pistol grip” style interface, with a trigger to apply suction to the needle tip. In this configuration, the user’s entire arm is moved to position the needle. There is a need for a device that can be operated using the fingertips, where fine motor control can sense differences in tissue consistencies and control suction levels.

Customer Requirements and Engineering Specifications
The device must perform all the same functions as the existing devices, while allowing for greater fine motor control and sensitivity during the aspiration process. Fine motor control can be accomplished by using a smaller device, operable with the fingertips on one hand (either hand), that allows the user to adjust the amount of suction applied to the needle during aspiration. The device must collect a sufficient amount of tissue for testing (at least the volume of a needle), and must be able to express the sample with the same ease as current devices. A disposable device should cost less than $3 per unit, and a reusable device should cost no more than $100. Lastly, the device must be safe.

From these customer requirements, we defined our engineering specifications. The most important specification for the new design is a pen-like cylindrical shape that can be manipulated with the fine motor control of the fingers on one hand. The table below shows the measured engineering specifications for our design. Additional specifications are incorporating the standard Luer-Lok™ feature that enables the attachment of various sized needles, and that the pressure vacuum is controlled by a single action.

<table>
<thead>
<tr>
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<td>&lt; 5 kPa</td>
<td>≥ 0.04 cc</td>
<td>&lt; 20N</td>
<td>&lt; 250 g</td>
</tr>
</tbody>
</table>

Concepts Considered
After generating and evaluating several concepts in the mechanical (valve design, push slider, direct slider, spring and locking slider, spring loaded, lever arm, live hinge, and rack and pinion), electrical (linear actuator), and pressure (shoe pump and diaphragm valve) categories, we decided to pursue the valve design, specifically considering a pinch, gate, and ball valve.

Concept Selection Methodology
Our concept selection methodology involved using Quality of Functional Deployment charts to rank which concepts met customer requirements most effectively. We also used scoring matrices, evaluated with the requirements of team members and Professor Davenport.

Engineering Challenges
In developing mock-ups, our main challenge was to create effective seals in our pinch, gate, and ball valve design concepts. Appropriate materials selection was an important consideration, so that adhesives would bond and proper shapes could be manufactured.

Rationale for the Final Concepts
Mock-ups and trial and error testing were used to create the final pinch and gate valve design concepts.

Deliverables for Design Expo
For the Design Expo, we re-built our pinch and gate valves into more polished final prototypes by making slight modifications to the preexisting mockups. We also prepared verbal presentation material, in addition to a poster and demonstration for Expo attendees.
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INTRODUCTION

Our sponsor, Professor Robertson Davenport, performs fine needle aspiration (FNA) biopsies in the Department of Pathology at the UM Hospital. He has experimented with various devices and methods, but has yet to find a fine needle aspiration device that meets all of his needs as a cytopathologist. Professor Davenport currently uses a Cameco® Syringe Pistol to perform his biopsies, although this device is less than ideal for several reasons (Figure 1). The most frustrating aspect of this design is that its use of a hand grip to apply suction ultimately requires whole arm movement for needle placement. When using the whole arm, even slight movements in the arm have significant effects on the sensitive needle/tissue interface and make needle placement more difficult. The absence of fine motor control also prevents the user from distinguishing between varying tissue consistencies, which is a crucial step in FNA procedures. Another shortcoming of this design is that the size of the device can make it very challenging to maneuver around certain areas of the body without rotating the hand into awkward positions while performing the procedure. Professor Davenport believes the procedure could be greatly improved by developing a device that possesses a more pen-like shape that can be manipulated with the fine motor control of the fingers on either hand. Given the compact nature of this idea, the device would have to incorporate a convenient method for applying suction to extract the tissue, maintaining a vacuum, and then emitting the tissue sample.

Figure 1: Cameco Syringe Pistol® [1]

Considering all parts of this design problem, a specific set of customer requirements was defined. The new fine needle aspiration device must: be operated with fine motor control, be operated with the fingertips of one (and either) hand, have user-controlled suction, be simple to operate, be sterile, be safe for the operator, have the ability to break up tissue, be affordable, be able to collect a large sample size in the needle, have user-controlled suction release, and be able to express the sample easily. Our ultimate goal is to produce a completed device that could be immediately implemented into use in the medical community and would outperform existing devices with its ease of use.

INFORMATION SOURCES

Our main source of information regarding FNA procedures and current FNA devices is our sponsor, Professor Davenport. Also, as mentioned in design review #1, Dr. Stewart Knoepp has assisted our team by helping us better understand the procedure, describing to us his own experiences with the current device and explaining his own ideas about changes that could be made to improve the current procedure [2].

Additionally, since design review #1, our team gained access to the ME 395 Laboratory at the University of Michigan. In the lab, we used testing equipment to finalize the engineering specifications for our design problem. Specifically, we used an Instron 4466 force transducer with a testing rate of 1 in/min to determine the forces required to extract the plunger in a 1cc, 3cc, and 10cc syringe to 2cc (1cc for the 1cc
syringe) of displacement. We also used a standard mass scale in the laboratory to take mass measurements of the syringes.

Technical Benchmarks
We would like to recap the three selected FNA devices that are already in existence as a basis for comparison. The Cameco Syringe Pistol® is known as the industry standard for FNA devices. The syringe pistol, shown in Figure 1 on page 4, is a reusable device that incorporates a standard disposable syringe and needle. The syringe pistol is operated by manually squeezing the “trigger,” which pulls the plunger back, creating the vacuum necessary to capture the tissue samples. The Tao Aspirator®, shown in Appendix A.1 on page 3 is a commercially available design that is designed to be held like a pencil. It is another example of a reusable design that uses the standard disposable needle and syringe. It is a finger-gripped style, which should allow for greater fine motor control. The plunger in this design is pulled back by a pre-loaded spring that is released by pushing a button on the device. The Cytec® device, shown in Appendix A.2 on page 3 is similar to our target design, but is not available in the US. This non-disposable design incorporates the use of a unique disposable vacuum chamber and standard disposable needle. The basic design has a pen-like shape that is easily manipulated by the fine motor control of the fingers. The device is “cocked” to a certain suction level before the aspiration takes place. Once the needle is inserted, a button turns on the suction, while a second button relieves the pressure when the procedure is complete.

Colleague Information
We were also fortunate enough to receive a reference table of maximum hand strengths of the human hand from our classmate and colleague, Max Bajcz. We used the information in this table as a guideline for the maximum activation force for our device. In particular, we used the maximum finger gripping strength, or strength of pinching your thumb and index finger together, for a “weak woman” as a starting point for a maximum activation force.

ENGINEERING SPECIFICATIONS
Engineering specifications, the major theme of our first design review, are the critical foundation upon which all of our concept designs are centered. The success of our final product will not be judged on how cool it looks, but rather on how well it satisfies the desired design specifications. It is important that we remain continuously aware of the specifications requested by our sponsor and those we have instituted ourselves. Many of our design specifications were discussed at length in design review #1, but a number of specifications have undergone further analysis since that time so it seems wise to review all of the specifications pertaining to our project.

User Requirements
Through meetings and ongoing correspondence with our sponsor, Professor Davenport, we have determined several of his key specifications. The device must perform all the same functions as the existing devices, while allowing for greater fine motor control and sensitivity during the aspiration process. Fine motor control can be accomplished by a device operable with the fingertips on one hand (and either hand) that allows the user to adjust the amount of suction applied to the needle during aspiration. In the current devices, the needle tip breaks up tissue in the mass so it can be collected for testing. The user must be able to release the suction while the needle is still in the patient, preventing the suction from taking in blood or fat from areas surrounding the mass while the needle is being pulled out of the skin. The device must collect a sufficient amount of tissue for testing (at least the volume of a needle), and must be able to express the sample with the same ease as current devices. A disposable device should cost less than $3 per unit, and a reusable device should cost no more than $100. Lastly, the device must be safe. For the patient, this means that it must be sterile, and therefore, a reusable device
should be autoclave-able (for sterilization). For the doctor, the device must present no more opportunity for an accidental needle prick than the risk already present with syringes and needles.

Relative Importance of User Requirements
Professor Davenport has led us to understand that fine motor control is his primary requirement for this device. He has described the aspiration process to us and shared some of his experiences performing the procedure. His descriptions have indicated that lack of fine motor control is the greatest disappointment in all of the current devices. As such, creating a device that can be operated by fine motor control is of utmost importance, and all other potential design features unrelated to this outcome are secondary concerns. Naturally, it is important that any device we design is capable of performing all the same tasks as or more effectively than performed by current devices. Designing a disposable device is certainly a point of interest for Professor Davenport, though he has assured us that this is not a crucial feature for our design.

Previously, we compared the customer requirements and technical specifications to define a set of engineering requirements in a Quality of Function Deployment chart (shown in Appendix D.1 on page 42). The results of that chart agreed with Professor Davenport’s requirements, ranking fine motor control (including user stability and device weight) as the top priority.

Determining Specifications
Various levels of testing were performed to determine numerical specifications for the final device shown in Table 1 below. An obvious specification is the size of the design. We assumed a cylindrical shape for the device and practiced holding and manipulating cylinders of different sizes to determine an acceptable size for a finger-operated device. Next, we specified the weight of the device by holding objects of different weights and determining qualitatively the maximum weight that still allows for fine motor control. The force required to activate the suction in the needle is a driving specification for this design. We tested how much force is required to retract syringe plungers to displacements of 2cc in the ME 395 lab with the assistance of Tom Bress. The results of our tests, shown in Table 2 on page 7, indicated that the force required to retract a 10cc syringe is nearly ten times the force required for a 3cc syringe. We also must insure that the required user input force is within the capability of a human hand. We were especially interested by the amount of force that can be exerted by a pinching motion between two fingers. Thanks to some data provided by classmate Max Bajcz, we determined a maximum activation force of 20N (See Appendix B on page 34). This includes a safety factor of 2 to insure we remain well below the limit of human capability. Since the existing devices can collect up to several cubic centimeters of tissue and blood, we decided to specify a minimum volume of tissue the device must be able to collect. Analysis of the sample requires no more tissue than what can be collected in the needle of a syringe, so we determined the volume of a standard gauge needle and made it a minimum requirement.

Table 1: Engineering Specifications

<table>
<thead>
<tr>
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<td>≥ 0.04 mL</td>
<td>&lt; 20N</td>
<td>&lt; 250 g</td>
</tr>
</tbody>
</table>
Table 2: Force to pull back plungers

<table>
<thead>
<tr>
<th>Syringe Capacity (cc)</th>
<th>1</th>
<th>3</th>
<th>10</th>
</tr>
</thead>
<tbody>
<tr>
<td>Force to pull plugged plunger to 2cc displacement (N)</td>
<td>1.74*</td>
<td>5.2</td>
<td>45.95</td>
</tr>
<tr>
<td>Force to pull plunger to 2cc displacement in an apple (N)</td>
<td>2.25*</td>
<td>5.4</td>
<td>---</td>
</tr>
<tr>
<td>Syringe Weight (g)</td>
<td>5.44*</td>
<td>9.98</td>
<td>12.47</td>
</tr>
</tbody>
</table>

* 1cc syringe plunger pulled back all the way to 1cc displacement

Benchmarks
We compared our design to the technical benchmarks (Cameco Syringe Pistol®, Tao Aspirator®, and Cytec device®), described in detail on page 5. The size and shape of all our concepts were specifically selected in order to allow for maximum control with the hand and fingers, rather than the arm. Because many of our concepts are relatively small compared to the benchmarks, they are likely to weigh less. The suction pressure and collection volume are more dependent on the user than they are on the design of the device. In our concepts, the user can apply the same amounts of pressure and collect the same volume of tissue as with the benchmarked devices. The method of creating suction does vary between concepts, and because some concepts may be operated by finger forces rather than hand forces, it may be more difficult to apply suction in some. All of the benchmarked designs are reusable devices, but a number of our concepts are disposable, which is desirable, according to Professor Davenport.

Tradeoffs
The specifications had not undergone much evolution in the project. From the beginning, the importance of a small device was greatly emphasized and many of our specifications were determined accordingly. Ultimately we were seeking to design a compact device and often designing small devices can become rather challenging, particularly when they are technologically complex. Our ideas to create a simply-operated device allowed us more breathing room where size was concerned. The activation force, or force required to retract the plunger of the syringe back to 2cc, had been a strong point of interest during early stages of concept generation. If the user is required to apply this force during the procedure we want it to require the least amount of effort possible, but until originally we didn’t know how much force would actually be necessary. Having gathered this data, we were more aware of our limitations and took them into consideration as we moved forward in the design process. For example, the possibility of a design that incorporated a linear actuator had brought to light a number of obvious tradeoffs. While the linear actuator would allow the device to operate by the simple push of a button, it is somewhat large and heavy compared to other designs and would also require an external power source. Because “large” and “heavy” are very much in disagreement with our number one priority to design a small device that is operable by fine more control, such trade off”s would not be considered worthwhile at this stage.

CONCEPT GENERATION
Whether we follow standard engineering design methods shown in an IDEO video in our ME 450 design class, or read over our lecturer Professor Skerlos’ posted presentation, the first two steps necessary to generate design concepts are functional decomposition and brainstorming.

Functional Decomposition
In the functional decomposition step of the concept generation process, the desired outcome is broken down into the operations required to get to the outcome. In the case of designing a fine needle aspiration device, the device must 1) pierce the skin, 2) break up the tissue, 3) retrieve a tissue sample, and 4) discharge the tissue sample.
Brainstorming
In the brainstorming step of the concept generation process, we compiled a list of various ways to accomplish each of the functional decomposition steps. We generated several ideas that would work well for one functional decomposition step, but did not seem to directly fulfill the needs of all the other functional decomposition steps. Nevertheless, we still listed all of our ideas, some reasonable, and some outrageous, in hopes that a few unlikely ideas could spur on our creativity into one state-of-the-art idea.

Some of the ways we brainstormed to pierce the skin include using a guitar string, bullet, porcupine needle, screw, nail, laser, thin straw, needle, pin, tack, briar, nettle, bee stringer, plastic needle, and capillary tube. Then, to break up the tissue, a screw, reverse auger, ultrasound, chemical reaction, tweezers, scissors, jackhammer, sandblaster, squiggle pen, sharp needle, sharp plastic needle, or fishhook could be used. The third functional decomposition step, retrieving a tissue sample, could be accomplished using a straw with suction, conveyor belt, screw, gravity, pressure increase in the tissue, siphon, pulsed low pressure, or esophagus contraction. Various ways of accomplishing the final function, discharging the tissue sample, include using reverse pressure, a mechanical push, a siphon, chemical precipitation, or gravity.

In addition the the exact functional decomposition steps, we also brainstormed different types of energy sources that could be harnessed to remove the tissue. A mechanical energy source could be the force of a finger, electrical energy sources could include AC power and a battery, and a chemical energy source could involve a gas producing or gas consuming reaction. Looking into more abstract energy sources, we hypothesized that a thermal energy source could include freezing or ‘melting’ the tissue, an ultrasonic or microwave energy source could use sound waves to vibrate tissue cells loose, and a solar energy source could involve focusing light to potentially burn through the tissue. A magnetic energy source was considered, but could only work if tissue was polarized.

Concept Generation Results
From the functional decomposition and brainstorming steps, we drew conclusions and created concepts for our design. Looking at the list of ways to pierce the skin, we concluded that using a needle will be the best option. Because a needle is currently the standard device used to pierce the skin, we know that it meets the strength requirements necessary for the fine needle aspiration process. A needle is sterile and biocompatible. Also, a needle is minimally invasive, creating a minimal amount of pain and scarring for the patient.

Similarly, looking at the list of ways to break up the tissue, a sharpened needle was concluded to be the best option. Again, the needle’s strength, biocompatibility, and minimal invasion set it apart from some of the other ideas for breaking up tissue. Additionally, we concluded that it would be better to use one device (a needle) to complete two steps in the functional decomposition (pierce the skin and break up the tissue) than it would be to attach an additional device for breaking up the tissue. We plan to use the Luer-Lok™ design for our needle attachment because it is a readily-accessible, existing standard (Luer-Lok™ is a trademark of Beckton-Dickson Co).

Although we concluded that a Luer-Lok™ needle will be the best idea for piercing the skin and breaking up the tissue, we still created many concepts using our various ideas for retrieving and discharging the tissue sample. All of our concepts, including the obviously infeasible ones, are documented in Appendix C on pages 35-41. We classified the concepts into three main categories: mechanically actuated, electrically powered, and pressure driven. Some of the main concepts included in the appendix are the mechanical valve design and push slider design, the electrical linear actuator design, and the pressure-driven shoe pump design.
Mechanical
The valve design (Appendix C.1 on page 35) is a mechanical design, and also is the only potentially disposable design. It would be placed between the standard needle attachment and the Luer-Lok™ of the syringe. The valve could be opened and closed throughout the fine needle aspiration procedure to control the suction level. The push slider design (Appendix C.3 on pages 35-36) converts a vertical finger force on the syringe pen to a horizontal force on the syringe plunger, to control suction. Several variations of this direct finger force, mechanically translated into syringe slider action, were created. Some other concepts in this category included the direct slider, spring and locking slider, spring loaded, lever arm, live hinge, and rack and pinion, and are shown in Appendix C.

Electrical
The electrical linear actuator design (Appendix C.2 on page 35) is our only design to involve an electrical energy source. It would use an electrical power source and linear actuator to move a syringe plunger up and down, controlling the pressure/suction level.

Pressure
The shoe pump design (Appendix C.10 on page 38) is similar to the push slider design because it converts a finger force perpendicular to the syringe pen axis to a pressure differential to control suction. However, the force translated in this concept involved pressure rather than a mechanical set-up. Another pressure-related concept included the diaphragm valve, which is shown in Appendix C.

CONCEPT SELECTION
Following the functional decomposition, brainstorming, and concept generation activities, our team was faced with choosing between eleven potential designs. In order to evaluate each of the designs in a qualitative manner, we made use of a scoring matrix, similar to a Quality of Function Deployment diagram. However, in this case, each of the designs was weighed against the technical requirements of the design.

Technical Requirement Weights
The first step in setting up a design scoring matrix is to clearly define the technical requirements used in the scoring matrix. First, the device should be lightweight in order for the user to maintain fine motor control during the procedure. Secondly, the diameter of the device is a crucial requirement and a big part of our design problem. A device that can be grasped like a pencil rather than held like a pistol would greatly benefit the user. We also defined a requirement to help distinguish the user stability, as well as one that defined the amount of motion required to use the device, user range of motion. Device flexibility reflects the way the device can conform to the user’s fingers. Another important requirement to maintain fine motor control was the characteristic length. The characteristic length is defined by the distance from the user’s fingertips during the procedure to the needle tip. We felt that a device that would minimize this length would give the user better control and a sense of feel during the procedure. The device should also have a relatively low market price, especially if it would be a feasible solution to our design problem. Related to this requirement was the manufacturability of the device. If the design were to be mass produced, a disposable device should cost less than $3 per unit, and a reusable device should cost no more than $100. Next we considered the pressure activation force which is the force required by the user to activate the suction pressure during the procedure in order to collect the tissue cells that are removed. Finally, the aesthetics of the device should always be considered so the device’s appearance is not intimidating for a patient.
Constructing the Design Scoring Matrix
To develop the design scoring matrix, we began with the set of technical requirements that were defined above. Namely, these are device: weight, diameter, flexibility, characteristic length, user stability, manufacturability, user range of motion, relative market price, pressure activation force, and aesthetics. Next, each of the technical requirements was given a weight based on their importance for overall design success. Then, we scored all of the eleven designs outlined above based on their correlation (weak, moderate, or strong correlation) to each of the technical requirements. The sum of the ranked correlations defined which design would most sufficiently meet the technical requirements. The results of the design scoring matrix revealed that the valve design was a clear favorite, followed by the electrical linear actuator. The third place position resulted in a tie between the direct slider and the shoe pump design. To solve this problem, we administered an engineering survey to each member of our design team. Specifically, each engineer independently ranked the designs numerically (from one to eleven) based on their personal intuition as to which of the designs would best meet the technical requirements. We felt that incorporating human intuition may also help us decide which designs may be difficult to manufacture and prone to failure, even if they closely met the technical requirements. The engineering survey served as tie breaker and consequently, the third place design was decided to be the push slider rather than the shoe pump. See Appendix D.2 on page 43 for a figure of the design scoring matrix diagram.

The Third Place Design
The results of our design scoring matrix suggested that the third best design was a push slider design. As stated earlier in concept generation, the basic idea behind this device is translating a vertical force into a horizontal displacement. Such a design would allow a user to press a button on the side of a pen style syringe and translate that force into expanding the syringe’s plunger and creating a vacuum chamber within the barrel of the syringe (Appendix C.3 on pages 35-36).

The most desireable characteristic of this design is its simplicity. Geometry as simple as a triangular block and a circular button could accomplish the motion necessary to make this design work. Secondly, because of the direct force to displacement interface of the design, the user would have direct control over suction level, and in turn the stroke length that the plunger is extended. Additionally, a seasoned veteran of such a device would be able to sense the amount of suction created during the procedure after developing a feel for the device’s button force pressure relationship.

On the other hand, the design also has its shortcomings. First of all, depending on the syringe size used, there would be a direct tradeoff between the stroke length necessary to achieve a 2cc displacement (accepted extension for FNA procedures) and the application force. Specifically, a larger syringe would require a greater activation force to achieve a 2cc displacement while a smaller syringe would require a longer stroke length. Since a smaller syringe diameter is desirable in our case, it may be necessary for the user to press the button more than once to achieve an acceptable plunger displacement for adequate vacuum pressure. In this case, we would need to add a locking device to the plunger end of the syringe to lock each subsequent displacement. Doing so would greatly complicate the simplicity of the design which was its greatest asset.

The Second Place Design
Coming in second place was an electrical device that incorporated a linear actuator. The basic idea behind this reuseable device is an envelope or casing that would enclose the syringe, while an actuator would be attached to the plunger of the syringe. After locking the device around the syringe, the user would have complete control over extension and compression of the plunger by means of a “car window” type switch controlling the actuator (Appendix C.2 on page 35). Needless to say, the actuator would require either an internal or external power source to function.
At first glance, the electrical linear actuator design seems like it may be an unrealistic and infeasible option for our design challenge. However, it has a few notable characteristics that allowed it to score exceptionally well in our design scoring matrix. First of all, incorporating a machine (actuator) into the design would allow for the user of the device to concentrate solely on performing the FNA procedure rather than worrying about having to squeeze or apply a force to achieve suction. Along the same lines, since the actuator can be controlled by a switch, the characteristic length (distance from finger tip to needle tip) could be minimized. Such a feature could increase the sense of feel during the procedure. Lastly, in this design the user has direct control over the amount of suction used during the procedure and could modify this as they saw fit.

The downsides of the electrical linear actuator design are quite obvious. First of all, our specification for a weight limit places major constraints on the size of our actuator, button, and power source. Second, heavier objects have a tendency to be bulky and uncontrollable, especially when the center of gravity is not positioned carefully. This could drastically compromise the fine motor control of the device. Additionally, the need for incorporating a power source, either internal or external, adds extra cost, maintenance, and replacement issues that the other designs escaped. Furthermore, the speed of the actuator could compromise the time it takes to apply suction during in the procedure which would be undesirable.

**CHosen DESIGN DESCRIPTION**

Based on our specifications and the customer requirements, we have decided that the valve design, shown in Figure 2, is the best design to achieve the desired results. This section will describe the device, its operation, and how it was selected as the best design.

**Figure 2: Valve Design**

![Valve Design Image]

**Device Description**

The valve design device will make use of the standard equipment already used in fine needle aspiration procedures. The needle and 10cc syringe are considered to be the standard for this procedure, and we do not want to change that.

The device will consist of two main parts. The first part is a valve with Luer-Lok™ connectors on either side. The valve will be cylindrical, with a diameter roughly equivalent to the diameter of the syringe. On the exterior of the cylinder, there will be a control to open and close the valve. The second part of the device is a chock designed to fit between the syringe plunger pull tab and the syringe body tabs, or some other type of syringe plunger locking mechanism.

**Device Operation**

To operate the device, the cylindrical valve will first be threaded onto the syringe tip, and the aspiration needle will be threaded onto the opposite end of the valve. The cylinder will function as an on/off valve for the syringe, allowing syringe pressure to reach the needle tip when the valve is open.

Once the valve, needle, and syringe are assembled, the syringe plunger will be pulled back to about 0.5cc displacement, and the valve placed in the closed position. Then, the plunger will be pulled back to the
full 10cc displacement. This is where the second part of the device comes into play. While holding the plunger out at the 10cc position, the chock will be inserted between the syringe body and the plunger tab, holding/locking the plunger at the 10cc position. The syringe barrel now contains a vacuum, which can be released with the valve.

The user will then hold the device like a pen or marker, and proceed to insert the needle into the lumped tissue. Once the needle has been inserted, the user will press the valve button to open the vacuum present in the syringe barrel to the needle tip. The user will be free to move the needle around in the lump with the vacuum applied. Once the user has collected enough sample tissue, the button will close the valve, shutting off the vacuum to the needle tip. The user will remove the needle from the lump without danger of sucking up blood in the needle tip.

Once the needle has been removed from the lump, the chock/lock will be removed, allowing the plunger to return to its position of 0.5cc displacement. The valve will then be opened, and the plunger depressed to express the desired amount of sample onto a slide for testing.

**Valve Type**

By Design Review #2, a final valve design had not yet been determined; however, we had generated three valve concepts that we believed could easily be activated and maintain a pressure differential of nearly 100 kPa for several minutes. The three designs are the gate valve, ball valve, and pinch valve. After completing a valve scoring matrix (Appendix D.3 on page 43), we decided to use the pinch valve as our target design.

The concept of the pinch valve uses a piece of flexible tubing as the method of sealing the valve in the open and closed positions. Our conceptual design can be seen in Figure 3 below. A hard outer shell protects the flexible tube on the inside. The rocker switch has a bump on the bottom of one side, and is positioned so that when in the closed position, the bump on the rocker switch will pinch the flexible tube so that it will not allow any air to pass through. When the rocker switch is in the open position, the bump is removed from the tube, allowing it to remain open, so air can flow. Although we did not yet know how to accomplish its locking feature, the rocker switch should be such that it can snap from the “on” or “off” position and hold itself there.

**Figure 3: Pinch Valve Concept**

![Pinch Valve Concept](image)

**Justification**

This device was selected as the best choice because it stands out from the rest of our designs in several areas. As shown in the design scoring matrix (Appendix D.2 on page 43, the valve design scored well relative to the other designs in several key categories: diameter, stability, and activation force. The diameter is important because it affects the ability of the user to grip the device like a pen. Since the diameter of the valve design will be no greater than that of the syringe itself, the valve scored very well. The stability rating reflects a user’s ability to keep the device in the desired position during the procedure. Since the valve design will be light and small, we expect that the user will easily be able to maintain position. The activation force refers to the relative force required to activate the vacuum pressure to the
needle tip during the procedure. Since the valve requires only a minimal force to press the button, the design scored well in the activation force. The complete scoring can be seen in Appendix D.2 on page 43.

Note that, although the pinch valve was selected as the best choice through the use of a scoring matrix, the gate valve and pinch valve are still potentially feasible designs. We decided to pursue the development of all three of these designs unless they proved to be infeasible at some point.

MOCK-UP DESCRIPTIONS

Our mock-ups are full scale, fully developed models of our final design. Each mock-up exhibits the same functionality as is expected of the final design, so we will not repeat the details of what the mock-up design is, how it works, and why it works in this section. We will simply discuss the relationship between our current mock-up and our final design in detail, explaining the larger points of similarity and difference. We will also state how each mock-up proves the most important elements of our final design.

Pinch Valve

The pinch valve mock-up, shown in Figure 4, proves that we derived a good solution to the fine needle aspiration device problem presented by Professor Davenport. The pinch valve mock-up proves the most important elements of the final design. The fine motor control force of a fingertip can control the valve’s push button dowel. The push button is able to pinch a pressurized tube, allowing suction to be held and released throughout the fine needle aspiration procedure. Also, the mock-up has a saw-tooth, locking syringe plunger, as will be used to hold and release suction in the final design.

Figure 4: Pinch Valve

The pinch valve mock-up validates our final design. It is not a scale up or a scale down of our actual design, but is scaled appropriately to resemble our final design within engineering specifications. The mock-up uses the standard Luer-Lok™ fittings, syringe barrels, and needle sizes as will be used in the actual, final design.

Although the mock-up proves the validity of the most important elements of our design, it was anticipated that there would be significant differences between the final design and the mock-up. For example, in our mock-up, we added circular end caps to the valve casing ends to provide more area to attach the Luer-Lok™ fittings. Next, we attached the latex tubing to the Luer-Lok™ barbs and glued the Luer-Lok™ ends to the end caps, and the end caps to the valve casing. In the final design, it was likely the Luer-Lok™ fittings would be glued directly to the valve casing, which would be made from solid polypropylene stock. This would eliminate the need for an end cap. Another area of improvement in the pinch valve design would be using a solid pinch valve button base, not a cross section, as used in the mock-up. The last area of improvement in the pinch valve final design would be adding a locking
mechanism to the pinch valve button, which is not securely fastened to the current pinch valve mock-up. In the final design, each saw-tooth of the locking syringe plunger could be manufactured in a repeating stamp pattern, rather than individually and inconsistently cutting out with a blade. In terms of aesthetics, our final design may include a smooth outer surface and painted finish, which our mock-up does not exhibit.

**Pinch valve special challenges:** Maintaining airtight seals throughout this design was challenging because of the number of interfaces we had to seal. Upon completing the valve for the first we discovered we had failed to create the necessary seal and air was leaking from one of the interfaces. We placed the valve under water to locate the leak. It appeared that the seal between the inner tubing and the Luer-Lok™ attachment had failed. We removed the Luer-Lok™ tip and added more epoxy to repair the seal. Because the function of the design is highly dependent on the user’s ability to pinch off the inner tubing it was important to find tubing easy to collapse. Initially we used softer vinyl tubing with an unnecessarily large inner diameter. Not only was the vinyl a little too hard, but the size of the inner diameter meant the user had to push the dowel further and with a greater force to fully collapse the tube. We relieved this problem by replacing the vinyl tubing with latex tubing that had a significantly smaller inner diameter. The softness of the latex tubing makes it easier to pinch, while decreased inner diameter means less tubing has to collapse and less force is required. After making the corrections to seals and replacing the tubing, everything appeared stable and our design was ready to move on to further stages of testing.

**Gate Valve**
The gate valve mock-up, shown in Figure 5, proves that we derived a second promising solution to the fine needle aspiration device problem presented by Professor Davenport. The pinch valve mock-up proves the most important elements of the final design. The fine motor control force of a fingertip can control the valve plunger. The plunger (with a lubricated seal) is able to slide through the gate, allowing suction to be held and released throughout the fine needle aspiration procedure. Also, the mock-up has a saw-tooth, locking syringe plunger, as would be used to hold and release suction in the final design.

**Figure 5: Gate Valve**

The gate valve mock-up validates our final design. It is not a scale up or a scale down of our actual design, but is scaled appropriately to resemble our final design within engineering specifications. The mock-up uses the standard Luer-Lok™ fittings, syringe barrels, and needle sizes as will be used in the actual, final design.
Although the mock-up proves the most important elements of our design, it was still expected that there would be significant differences between the final design and the mock-up. For example, in our mock-up, we drilled a rough hole for the gate dowel, used a purchased rod of given diameter for the dowel, added rubber seal rings to the dowel, and lubricated the seals with liquid soap. In the final design, more precise material selection and manufacturing methods could be used to insure a tight seal between the gate hole and the valve plunger in the gate valve. Another area for improvement in the gate valve final design would be manufacturing a solid tube for the gate valve body, rather than filling an annulus with epoxy or fiberglass resin, as was done for the gate valve mock-up. Lastly, in the final design, each saw-tooth of the locking syringe plunger could be manufactured in a repeating stamp pattern, rather than individually and inconsistently cut out with a blade. In terms of aesthetics, our final design would include a smooth outer surface and painted finish, which our current mock-up does not exhibit.

**Gate valve special challenges:** Similar to the pinch valve, the greatest challenge of this design was to maintain air tight seals between interfaces. Adding sufficient amounts of sealant and epoxy allowed us to successfully seal each interface on the main valve component. The next challenge however was to insure an airtight seal between the valve interfaces with the dowel pin that regulates air flow through the valve. Smooth surfaces are critical for the airtight seal, and so are tolerances. We pulled a piece of latex tubing over the dowel in hopes of creating a better seal at the interface.

**Ball Valve**

Figure 6 shows our attempt at manufacturing a ball valve mock-up. We were not able to create a ball valve mock-up that solves the fine needle aspiration device problem provided by Professor Davenport. Because rubber is not an easily-manufactured material, we were not able to create a rubber stop of the correct sealing shape for the ball valve. Also, because a flexible adhesive for a polypropylene-PVC-rubber interface does not exist, we were not able to create an effective seal in this valve. At this point, we decided not to continue developing a final design for the ball valve.

**Figure 6: Ball Valve**

**Ball valve special challenges:** The most critical element of the ball valve was the surface interface between the ball bearing the conical interior shape that provides the airtight seal. Because the seal is broken by squeezing the cone and dislodging the ball bearing, it was important that the material used for the cone be soft enough to deform. We found that the first rubber component we used was too soft to be machined. As a result, we couldn’t accomplish a perfectly symmetrical cone and the airtight seal could not be established. Our second attempt was to use a harder plastic component that already possessed the conical shape we required. This presented a new challenge when we attempted to create a seal between the plastic component and the vinyl tubing surrounding it. Initially we used a marine-grade sealant; however after drying we noticed that upon squeezing the vinyl tubing to deform the cone, the sealant simply broke apart from the surface. In other words there sealant wasn’t actually having any bonding effect. Next we used a hot glue gun only to experience the same outcome; no bonding was taking place. We decided the only way to create a bond might be to melt the tubing and plastic cone together. First we attempted to melt a small stretch of plastic on the perimeter of the cone component with a soldering gun.
While the vinyl tubing melted rather successfully, we found that the plastic was very difficult to melt and it was impossible to create a large enough melt surface for the two pieces to adhere to one another. Our final attempt to bond the two surfaces consisted of taking a lit match and holding it up to the surface of the plastic. As before, the plastic melted very slowly, and rather than transforming to molten plastic it began to burn off. Given our inability to create an airtight seal between the cone and vinyl tubing which is a mandatory feature of the design, we have decided the ball valve design is not worth pursuing any further at this time.

ENGINEERING DESIGN PARAMETER ANALYSIS

The majority of the designs for both the gate valve and the ball valve had been determined through the process of building mock-ups. We knew ahead of time that many of the parameters that were needed to determine the performance of the devices would not be available, and we would need materials and properties testing to determine the parameters. One major concern for any valve is sealing, which is difficult to quantify for moving parts. The original concept for the gate valve involved a rigid-to-rigid surface that would seal and slide at the same time. We quickly realized that this would require very low surface roughnesses which are probably unattainable on machined plastic parts. In the ball valve concept, the seal was dependent on the interface between the ball bearing and the rubber cone. Whether or not this interface would seal was dependent on the roughness of the ball, rubber, and the amount of compression on the rubber. We had no way of quantifying the roughnesses, and in reality, slight imperfections in either surface would lead to a faulty seal, and therefore device failure.

Knowing that so many of the parameters would be difficult to determine, we decided to begin experimenting with different ideas and see what worked well. In the following paragraphs, we will describe the design evolution based on our experimentation for the ball valve, gate valve, and pinch valve.

Ball Valve

The original concept for the ball valve required a rubber cylinder with a conical indentation to cradle the ball bearing. We bought a rubber stopper and immediately found that rubber is not easily machinable. We attempted to use a drill bit with a point tip to tap-drill the top of the cylinder to cut a conical indentation. In order to secure the stopper in the drill press, it had to be compressed in a vise, which distorted its shape. After the top was drilled and removed, there were two issues. First, the surface of the cut was rough, since rubber does not form chips, like rigid materials, but instead stretches and tears. Second, the conical indentation was not perfectly round, as it had been distorted by the vise. These two problems led us to search for another method of creating a seal with the rubber.

We purchased a rubber gasket that had a preexisting conical indentation of the right size to fit the valve. Some basic testing showed that this combination of gasket and ball could create a seal if the ball was pressed into the cone. The next step was to attach a flexible tube to the gasket, to serve as the chamber that could be deformed to break the seal. We immediately found that the issue of sealing rubber to any other material would be very difficult. We tried to use silicon sealant, but the sealant would not hold to the flexible joint. The issue was that the joints in the ball valve would require a rigid material to be bonded to a flexible material that would deform in use. To bond the rubber to the tube (PVC), we tried epoxy, silicon, hot glue and melting, of which none worked.

The combination of the two issues noted above led us to abandon the concept of the ball valve. We determined that creating an airtight seal between a flexible part and a rigid part would be too difficult for a small, cheap, easily manufacturable device, so we gave up and pursued the other designs.
Gate Valve
The gate valve concept called for a solid cylinder with a small through-hole along the axis of the cylinder, with a slot cut in the cylinder perpendicular to the axis (see Appendix G.1 and G.2 on pages 46-47). A plastic plate would slide in the slot, closing off the through-hole, and sealing the valve. After some consideration, we decided that a cylindrical hole, rather than a slot, would be significantly easier to machine. To create the cylinder with through-hole, we glued a piece of 1cc syringe barrel concentrically inside a piece of 10cc syringe barrel, and filled the annulus with fiberglass resin. When we drilled a perpendicular hole through the cylinder, we found that the resin tended to splinter and chip. This prevented a seal from forming between the cylinder and the rod which would function as the gate. After repeating the process with epoxy resin and getting the same results, we decided that drilling through epoxy or fiberglass will never give the smooth surface necessary for a seal.

We decided to instead use a preexisting smooth cylinder (another syringe barrel) as the surface to be sealed (see Figure 7). We also decided to take a trick from the syringe designer’s book, and use a rubberized plunger to create the seal between the hole and the gate. We know that this is the technique for syringe plungers, and they seal fairly well so we assumed this method would create a good seal for our valve as well. After some simple tests, we found that the rubber in the tube does seal, and we decided to move forward with that course of action. The dimensions of the hole, rod, and rubber will remain the same from the mock-ups to the final prototype, so those parameters are already determined.

Figure 7: Gate valve mock-up

Pinch Valve
The pinch valve gave us the least trouble as we built a mock-up. We first knew that a flexible tube would be needed, but we did not know how to quantify the softness or flexibility of a tube, so we bought the first thing we found at the hardware store -- a piece of small diameter vinyl tubing. The vinyl tubing proved to be too difficult to squeeze closed by finger force, so we searched McMaster-Carr and bought the softest tube we could find, which was latex rubber. This was essentially the only design parameter specific to the pinch valve that we needed to redefine from the initial concept.

General Notes
Both of the chosen valve designs require a Luer-Lok™ connector on either end. Originally, we planned to use the Luer-Loks™ on the ends of the provided syringes, and glue them onto the valves. However, they proved to be weak, and broke off easily after being glued. We then found that Luer-Lok™ fittings are readily available on the internet, and we purchased fittings with hose barbs, which attach easily to the latex tube. The first Luer-Loks™ we purchased were nylon. After several failed attempts at gluing many
of our parts with epoxy, we did some research and found that polypropylene is a low surface energy plastic, and will not bond without special adhesives. These special adhesives do not bond to nylon, so we are purchasing a new batch of Luer-Loks™ that are polypropylene as well.

All of the mock-ups have proven to have more than enough strength to withstand the loads they will experience, so we have determined that the solid mechanics of the situation are a non-issue. In terms of the pressures achieved in the syringe, we have an engineering specification target of less than 5 kPa when the plunger is in the full-back position. This means that the ratio of the initial volume to the full-open volume has to be less than 5 kPa / 101 kPa, or approximately 1/20. The pinch valve, which has the larger of the two internal volumes, has a ratio of less than 1/100, so we can be sure that the pressure achieved in the syringe will be sufficient.

**FINAL DESIGN DESCRIPTION**

This section describes our two solutions to the fine needle aspiration device design problem. The final design descriptions include what they are, how they work, and why they work. Details such as final design dimensions, materials, and operation are included. Appendix F on page 45 shows a Bill of Materials list of all off-the-shelf parts (along with manufacturer, part number, and cost) and all parts made in-house that were used to make our prototypes and were used in our final designs.

**Locking Syringe Plunger**

*Figure 8: Locking Syringe Plunger*

The locking syringe plunger is a mechanism developed to lock the plunger in a desired position in order to maintain a vacuum inside. As shown in Figure 8, the plunger itself has been altered to possess a saw-tooth shape and a wire has been attached to the flange of the syringe barrel. The mechanism operates by twisting the plunger so the teeth are parallel to the wire, retracting the plunger to the desired volume and then twisting the plunger back to the starting position such that the teeth are directed perpendicular to the wire and catch when the plunger is released to lock it in place. After testing our design hundreds of times we are convinced it of its robustness and ability to function effectively each time it’s used.

**Pinch Valve**

The pinch valve design is a valve situated between the syringe tip and the needle base (see Appendix G.3 and G.4 on pages 47-48). The polypropylene valve connects to the heat-treatable stainless steel needle and polypropylene syringe tip via a female, polypropylene Luer-Lok™ tip on one side, and a male, polypropylene Luer-Lok™ tip on the other; this is the traditional method used in today’s medical settings for connecting standard needles to syringes. Vinyl tubing runs between both ends of the valve to allow for air and substance flow between the needle and the syringe. There is a polypropylene passage that runs from the top surface of the valve down into the chamber containing the vinyl tube. The passage acts as the casing that houses the polypropylene push button dowel. The dowel possesses a small, J-shaped
cutout that will help lock to the dowel in the pinching position by a frictional contact with a secondary stationary dowel situated perpendicular to the motion of the push button dowel.

To operate the valve, the user will cut off airflow between the needle and syringe by pinching the vinyl tube with the push button dowel. After locking the dowel in the pinching position the user will then be free to pull the plunger of the syringe back to the desired capacity to create the vacuum inside. The syringe plunger will then be locked in the desired position. At this point, the user can insert the needle into the tissue lump of interest and activate suction by releasing the push button dowel from its locked position. When the required tissue has been retrieved the pinch can be re-engaged, to hold pressure, and the needle can be retracted from the skin. To eject the tissue sample, the pinch valve can be removed from the needle, and a positively-pressurized syringe can be attached to the needle. Additional procedural information for the pinch valve can be found in Appendix H, pages 49-52.

The successful functionality of the pinch valve is dependent upon its ability to maintain a vacuum and for the user to have control over the activation of the vacuum. Maintaining the vacuum requires the pinching act of the push button dowel to sufficiently compress the vinyl tubing and to keep it compressed. After testing our prototype a significant number of times, we found the pinch action repeatedly provided sufficient compression of the vinyl tube to prevent the flow of any air and could maintain the compressed state for the sufficient length of time; this ultimately means that the valve is capable of maintaining the necessary vacuum. In each test that we disengaged the pinch action, the vacuum was exposed to the needle tip as desired. This was sufficient evidence to us that the user will have control over the application of the vacuum.

Gate Valve
The gate valve design will also be situated between the syringe tip and the needle (see Appendix G.1 and G.2 on pages 46-47). The polypropylene valve connects to the heat-treatable stainless steel needle and polypropylene syringe tip via a female, polypropylene Luer-Lok™ tip on one side, and a male, polypropylene Luer-Lok™ tip on the other; this is the traditional method used in today’s medical settings for connecting standard needles to syringes. An open passage runs between both ends of the valve to allow for air, and substance flow between the needle and the syringe. There is an additional passage that runs from the top surface of the valve all the way through to the bottom surface. This passage acts as the gate casing that houses the polypropylene dowel plunger. The dowel plunger is slightly longer than the outer diameter of the valve casing and has two small Viton™ gaskets on it separated by a short distance.

To operate the valve, the user will cut off airflow between the needle and syringe by positioning the plunger such that one of the Viton™ gaskets is in line with the through-passage of the valve, creating an airtight seal. After the dowel plunger is placed in the blocking position in the gate, the user will then be free to pull the syringe plunger back to the desired capacity to create the vacuum inside. The syringe plunger will then be locked in the desired position. At this point, the user can insert the needle into the tissue lump in question and activate suction by forcing the dowel plunger into the valve so the gasket is moved past the through-passage and the airtight seal is broken. When the required tissue has been retrieved, the blocked position can be re-engaged by forcing the dowel plunger further inward so the second gasket creates a seal and then the needle can be retracted from the skin. Once the needle is out of the lump, the dowel plunger can be pressed until it hits the bottom of the stop, releasing the negative pressure of the syringe by venting it to the atmosphere. Finally, to express the tissue sample, the syringe can be pulled back to create a positive pressure chamber inside the syringe, and the plunger can be returned to the position that creates an airtight seal around the through-passage. This can be done by simply rotating the device 180 degrees and pressing from the bottom part of the plunger dowel. Additional procedural information of the gate valve can be found in Appendix I, page 53-56.
Like the pinch valve, the successful functionality of the gate valve is dependent upon its ability to maintain a vacuum and for the user to have control over the activation of the vacuum. Maintaining the vacuum requires the blocking action of the rubber gasket to allow for an airtight seal. After testing our prototype a significant number of times we found the seal created by the gasket repeatedly provided sufficient blockage to prevent the flow of any air and could hold the seal for the sufficient length of time; this ultimately means the valve is capable of maintaining the necessary vacuum. In each test that we disengaged the blocked position, the vacuum became exposed to the needle tip as desired. This was sufficient evidence to us that the user will have control over the application of the vacuum.

PROTOTYPE FABRICATION

There are two major differences between the prototypes that our team built as verification mock-ups and the final prototype designs. First, in each case, the valve casings were made from round solid polypropylene stock as opposed to cannibalized syringes, tee brackets, and bonding agents. This change gave each device more rigidity, and minimized the chance of air leaks at bonded interfaces in the device, which would result in failure. These changes also preserved a more pleasing aesthetic appearance to each of the devices.

Secondly, there were minor design changes to each prototype that are device specific. For the pinch valve prototype, two changes were made. First, the pinch button itself was made of solid polypropylene stock similar to the valve casing, as opposed to cannibalized portions of a syringe plunger. This slight modification allowed for a better pinch interface between the bottom of the push button and the vinyl tubing. The bottom of the mock-up pinch button had an “x” shaped cross-section, which needed to be aligned correctly to create a good pinch and seal. For the prototype, we made this bottom a hemispherical shape, which ensured that it would always seal the flexible tube against the bottom of the tubular channel. Secondly, the pinch button now incorporates a retaining and locking mechanism. This feature ensures that the pinch button does not separate from the valve casing, and also gives the operator the option to lock the valve in the closed position.

There were three design modifications between the gate valve mock-up and the prototype. First, we drilled a through hole into the gate valve casing, rather than inserting a 3cc syringe to serve as the interior surface for the gate seal. Second, the gate plunger for the gate valve design was turned on the lathe to put grooves in the rod, in which the sealing gaskets sit. Incorporating these notches guaranteed that the gaskets would not slide on the plunger during operation of the device. We also added an end button on the top of the plunger rod to give the user a larger surface on which to push. We also incorporated a lip around the bottom of the through hole in the valve casing. These design alterations help with user operation of the device by defining stops to mark the full extension of the plunger in each direction.

Fabrication

To begin the fabrication process, we first organized all of the materials we would be using into a bill of materials (see Appendix F on page 45). To simplify this section, each valve device was considered separately. The bill of materials includes components for each design; they are labeled accordingly. Note: the following sections include tool sizes in English units because that is what was available in our machine shop.

Prototype I: Pinch Valve

As a beginning note, the engineering drawing of the pinch valve (Appendix G.3 and G.4 on pages 47-48) may help visualize the descriptions of the following parts and their manufacturing procedures. The following is the procedure we followed to construct our pinch valve prototype. We began by manufacturing the pinch valve casing. We selected 19.05 mm diameter polypropylene stock as the
material for this part. The first step was to cut this rod stock down to the correct length. This was a simple cutting operation and was done on a band saw. Once this piece was cut down to 30.8 mm in length, we drilled a through hole through the axis of the round stock to house the flexible tubing. This operation was done on a drill press with a 0.295 inch drill bit. Following the drilling of the through hole, we drilled a countersink on either end of the body, using a 0.5 inch Forstner bit. This increased the surface area for the glue bonds to the Luer-Lok™ fittings, ensuring that the Luer-Loks™ would not be pulled off of the body during use. Next, we drilled a blind hole perpendicular to the axis of the stock, running halfway through the diameter of the body, until it reached the axial hole, as the shaft for the pinch button. We used a drill press with 0.315 inch twist drill bit to complete this procedure. Finally, we drilled a hole through the side of the pinch valve casing perpendicular to both the axial and vertical shafts to serve as a latch hole for the locking mechanism. We used a drill press with a center drill to first start the hole, and a 0.0625 inch twist drill bit to finish the hole once it had been started.

Following the completion of the pinch valve casing, fabricated the pinch valve push button. As mentioned earlier, we created this piece out of polypropylene rod, as opposed to a cannibalized plunger end, as we had done for the mock-ups. The rod had a diameter of 6.8 mm, and we cut this to a length of 13 mm on the band saw. Next, we cut a J-shaped slot into the side of the push button, perpendicular to the axis of the stock. We used a 0.0625 inch drill bit in a mill to cut the slot. Although that is not the appropriate tool for the job, we did not have access to a mill bit that was long enough to reach all the way through the rod. Next, we cut out an end cap for the top of the push button to increase the button diameter. We used a laser cutter to cut an 8 mm diameter disc from a sheet of acrylic, 2.36 mm thick. Our final manufacturing procedure for the pinch valve was to cut a piece of flexible PVC tubing to a length of 26.8 mm; this was done with an X-acto™ knife.

After fabricating the parts, we began the assembly of the pinch valve. The first step was to test-fit the parts together and see if the valve works when the parts are not glued together. We slipped one end of the flexible PVC tube over the barb tip of one Luer-Lok™, and then slid the whole tube into the valve body. We then pushed the other Luer-Lok™ barb into the other side of the flexible tube. Although that is not the appropriate tool for the job, we did not have access to a mill bit that was long enough to reach all the way through the rod. Next, we inserted the button into the button hole, and then the locking wire through the button and valve casing. Through experimenting with the setup and making small adjustments to ensure its proper functionality, we achieved a valve that consistently closed, locked, and sealed before doing any real assembly. Once we had test-fitted and dry-run the valve, we bonded the Luer-Loks™ on either end with ScotchWeld DP-8005 polyolefin bonder. We chose this glue because it bonds well to polypropylene, which is something that most glues and epoxies cannot do. Prior to gluing, we roughened all the surfaces to be bonded with low grit sandpaper. We also glued the button top onto the push button, again with the ScotchWeld. To finalize the device, we used a combination of files and high grit sandpaper to smooth out edges and the surface finish.

Prototype II: Gate Valve
Once again, viewing of the engineering drafting of the gate valve (Appendix G.1 and G.2 on pages 46-47) may help visualize the descriptions of the following parts and their manufacturing procedures. The fabrication of the gate valve began similarly to the pinch valve. We began making the gate valve casing, with the same 26 mm round polypropylene stock for material. Next we cut it to size with a simple cutting operation on a band saw. Once this piece was cut down to 30.6 mm in length, we drilled a through hole through the round stock axially to serve as our valve chamber. This operation was completed on a drill press with a 0.0625 inch drill bit. Following the drilling of the through hole, we manufactured a counterbore on each entrance to the through holes. Again, this helped increase the surface area for our connections to the Luer-Lok™ fittings and came in handy when bonding these joints during assembly. The same process as used in the pinch valve was used. Next, we proceeded by drilling a stop-hole perpendicular to the axis of the stock to serve as the shaft for the plunger button. Then, we used a V-
block to fasten the round stock and a simple drilling operation on a drill press with a 0.339 inch twist drill bit to complete this procedure. However, for the gate valve, we drilled almost completely through the casing, leaving only a small lip on the bottom edge; this will serve as a bottom end cap for the plunger.

Following the completion of the gate valve casing, we began fabrication of the plunger button. Again, we created this piece from polypropylene rod that has an outer diameter 6.8 mm. Next, we cut this to a length of 30.7 mm on the band saw. Then, we incorporated an end cap on the top of the push button to increase the button diameter, as we did with the pinch valve button. We used the laser cutter to cut a 10 mm diameter hole from an acrylic sheet with a width of 2.36 mm to accomplish this. The final fabrication of this part consisted of cutting slots into the outer edge of the plunger button to hold the rubber gaskets. We used a lathe to cut these grooves. Our final manufacturing procedure for the gate valve consisted of cutting the rubber gaskets from the Viton™ tubing (0.125 inch ID and 0.25 inch OD). One is 2.5 mm in length, and the other is 4.5 mm, (the slots turned in the plunger are the same length). This was completed with a pair of industrial scissors.

Following these operations, we began the assembly of the gate valve. Our first step consisted of bonding the end cap to the middle on the top point of the push button, as done with the gate valve. Next, we attached the male and female Luer-Loks™ to the ends of the gate valve casing, much like we had done for the pinch valve, minus the need for the vinyl tubing. Then we threaded the rubber gaskets into their corresponding holes on the plunger button. To complete the prototype, we lubricated the gaskets and insert the plunger into the hole. Once again, to finalize the device, we used a combination of files and high grit sandpaper to smooth out the edges and the surface finish. For each of our valve devices, our greatest concern with assembly was not the level of difficulty, but rather time. The structural plastic bonding took 24 hours to cure completely, and served as our only major time consumer during the fabrication of valve design.

**Tolerances**
A majority of the manufacturing necessary for each of our designs was not dependent upon high tolerances. However, each of our devices has certain areas in which tolerance needs to be considered.

For our pinch valve, the only area of high tolerance was in the locking mechanism. Without precise manufacturing in this area, trouble with the geometry of pinch valve operation would occur. Therefore, when manufacturing the J-shaped locking mechanism, we machined to achieve high tolerances in order to minimize the possibility of friction, rubbing and misalignment. As stated above, to combat this problem, we machined this part with a CNC milling machine to maintain high tolerance.

For the gate valve design, the tolerances for the end connections of the valve casing to the Luer-Loks™ as well as the gate plunger hole were considered. In this design, there is no vinyl tube to serve as a pressure chamber. Instead, our seals between the barbed ends of the Luer-Loks™ and the ends of the valve casing need to be matched precisely in order to obtain a seal. Additionally, the inside of the plunger hole needs to have an exact match with the outer diameter of the rubber gaskets on the plunger. In each case, precisely manufactured holes were crucial for our seal.

**Critical Surfaces**
Most of the areas in which small tolerances are an issue are also areas where surface finish was an issue. For our pinch valve, recall that the trouble area was the interface between the locking mechanism on the push button and the wire keystone of the casing. Without a smooth surface finish at this interface, we would have run into heavy friction between these materials, preventing flow and operation of the device. For the gate valve design, the inside of the plunger hole needed to have a smooth surface finish to facilitate the seal with the plunger gaskets. During fabrication for our prototype mock-ups, this surface
was a concern. Drilling the epoxy and resin fillers used to make our casings resulted in a chipped inner surface to this interface. Needless to say, anything but a smooth surface on this inner face resulted in device failure, and caused several early failures for the device.

Two other areas of interest in terms of surface finish are the areas between bonded surfaces, as well as the overall surface finish of our devices. As mentioned early, we prepared all bonded surfaces by roughening them with low grit sand paper. A rough surface area in these regions served as a better surface for the structural plastic ScotchWeld DP-8005 to bond. Finally, we made sure that the finish of our valve was balanced between a sleek and graspable surface. Aesthetically we wanted to maintain a semi-gloss appearance to the valves so that they mesh nicely with the syringes that they will be used with. However, we also needed to account for the fact that handling this device is essential for operational purposes, and therefore did not want to create an excessively polished outside surface.

PROTOTYPE DESCRIPTION

Our prototypes are full scale, fully developed models of our final designs. Each prototype exhibits the same functionality as is expected of its final design, so we will not repeat the details of what the prototype design is, how it works, and why it works in this section. We will simply discuss the relationship between our current prototypes and our final designs in detail, explaining the larger points of similarity and difference. We will also state how the prototypes prove the most important elements of our final designs.

Both prototypes are valves, which attach via Luer-Loks™ to the end of a syringe. A standard Luer-Lok™ needle can be attached to the other end of each valve. When the user wants to perform a fine needle aspiration, he or she will attach the selected valve and close the valve. The user will then pull back the plunger of the modified syringe and lock the plunger in the full-back position. The vacuum is now contained in the syringe barrel. Once the needle is inserted into the tumor or tissue in question, the user will release the vacuum with a button on the valve, allowing the needle to start applying suction to the tissue. When a sufficient sample has been collected, the user simply closes the valve, and can safely remove the needle from the tissue, without danger of spraying blood into the syringe barrel. Once the sample has been ejected from the needle, the valve can be disposed.

Locking Syringe

The syringe to which the valves attach must be a modified version of the standard Beckton-Dickson syringe used in hospitals. Figure 9, below, shows the modifications that we made to the syringe to adapt it to the procedure.

Figure 9: Locking Syringe
To use the syringe, the user simply pulls back the syringe as usual, except that when the plunger is pulled to the desired displacement, the user can rotate the plunger to engage the locking teeth. The syringe is now locked back, and the vacuum can be maintained in the valve without needing to hold the plunger. Once the procedure has been completed, the plunger is rotated back to the original angle, and it can be pushed back to the closed position.

**Pinch Valve**
The pinch valve prototype, shown in Figure 9, is a good solution to the fine needle aspiration device problem presented by Professor Davenport. The fine motor control force of a fingertip can control the valve’s push button dowel. The push button is able to pinch a pressurized tube, allowing suction to be held and released throughout the fine needle aspiration procedure. Additional pictures of the pinch valve can be seen in Appendix H, pages 49-52.

**Figure 10: Pinch Valve**

The current pinch valve prototype validates our final design. It is not a scale up or a scale down of our actual design, but is scaled appropriately to resemble our final design within engineering specifications. The prototype uses the standard Luer-Lok™ fittings, syringe barrels, and needle sizes as will be used in the actual, final design.

A significant number of changes were made in the parts, production and assembly of our prototype from our initial mock-up device. The general housing/casing of the valve was machined from a solid piece of polypropylene rod. It was cut to the desired length and then the through-hole and pinch button slot were drilled. The inner tubing that was originally Latex in our mock-up design was changed to a vinyl hose. We used a different variety of Luer-Lok™ attachments that included barbs on the end, helping to improve the connection between the Luer-Loks™ and tubing, making it more rigid and airtight. The pinch button itself was comprised of two parts; the shaped rod, and the button on top. The shaped rod was also made from polypropylene and the button was cut from acrylic and adhered on top. The pinch button also possesses a J-shaped slot used in the locking mechanism. The second component of the locking mechanism is a small wire that runs the diameter of the push button chamber and passes through the J-slot. When button is compressed downward and pushed forward, the wire locks in the rounded portion of the J-slot and holds the inner tubing in a closed state. We used ScotchWeld DP-8005 as our general adhesive for any bonding and sealing functions.

**Gate Valve**
The gate valve prototype, shown in Figure 10, is a second viable solution to our design problem. The fine motor control force of a fingertip can control the valve plunger. The plunger (with a lubricated seal) is able to slide through the gate, allowing suction to be held and released throughout the fine needle
aspiration procedure. Also, the current prototype has a saw-tooth, locking syringe plunger, as was used to hold and release suction in the final design. Additional pictures of the gate valve can be found in Appendix I, pages 53-56.

Figure 11: Gate Valve

The current gate valve prototype validates our final design. It is not a scale up or a scale down of our actual design, but is scaled appropriately to resemble our final design within engineering specifications. The prototype uses the standard Luer-Lok™ fittings, syringe barrels, and needle sizes as will be used in the actual, final design.

A number of changes were made in the parts, production and assembly of our prototype from our initial mock-up device. The general housing/casing of the valve was machined from a solid piece of polypropylene rod. It was cut to the desired length and then the through-hole and gate plunger slot were drilled. The gate plunger, also made from polypropylene rod, was machined with appropriately sized and positioned grooves to help grip the seals to the plunger. To eliminate the use of latex in the device, we replaced the old seals with Viton™ rubber. To keep the gate plunger moving more smoothly, we have been adding an oil-based lubricant to the seals. We used ScotchWeld DP-8005 as our general adhesive for any bonding, and sealing functions.

VALIDATION

Engineering Specifications vs. Prototype Parameters
As mentioned earlier in this report, the engineering specifications of our design project were the major drivers of our design process. They were critical foundations upon which all of our concepts, designs, and ideas were centered. Satisfying the engineering specifications, especially with our prototype designs, would allow our team to validate the approach taken to solve our fine needle aspiration device design problem. Additionally, some of the success of our prototypes will be evaluated on how well they satisfy the desired design specifications.

Now that the prototyping stage is complete, we have had a chance to verify the engineering specifications that were set earlier in the design process. Recall from earlier in the report that the pertinent engineering design specifications were the device diameter, length, suction pressure, collection volume, activation
force and weight. The table below lists the engineering specifications as well as the device parameters for our two possible prototype designs. Verification of the specification is marked in the right column.

### Table 3: Validation of Prototypes to Engineering Specifications

<table>
<thead>
<tr>
<th>Engineering Specification</th>
<th>Target</th>
<th>Prototype A Pinch Valve</th>
<th>Prototype B Gate Valve</th>
<th>Target Satisfied?</th>
<th>A</th>
<th>B</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diameter</td>
<td>( \leq 2.5 \text{ cm} )</td>
<td>2.5 cm</td>
<td>2.3 cm</td>
<td>YES</td>
<td>YES</td>
<td>YES</td>
</tr>
<tr>
<td>Length</td>
<td>11.25-15 cm</td>
<td>15 cm</td>
<td>14.7 cm</td>
<td>YES</td>
<td>YES</td>
<td>YES</td>
</tr>
<tr>
<td>Suction Pressure</td>
<td>(&lt; 5 \text{ kPa} )</td>
<td>4.1 kPa*</td>
<td>( \approx 0 \text{ kPa} )</td>
<td>YES</td>
<td>YES</td>
<td>YES</td>
</tr>
<tr>
<td>Collection Volume</td>
<td>( \geq 0.04 \text{ cc} )</td>
<td>( \geq 0.04 \text{ cc} )</td>
<td>( \geq 0.04 \text{ cc} )</td>
<td>YES</td>
<td>YES</td>
<td>YES</td>
</tr>
<tr>
<td>Activation Force</td>
<td>(&lt; 20 \text{ N} )</td>
<td>11.6 N</td>
<td>36 N</td>
<td>YES</td>
<td>NO</td>
<td>YES</td>
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<tr>
<td>Weight</td>
<td>(&lt; 250 \text{ g} )</td>
<td>21.3 g</td>
<td>28.3 g</td>
<td>YES</td>
<td>YES</td>
<td>YES</td>
</tr>
</tbody>
</table>

*These values were calculated, not measured.

### Experiments Used to Verify Engineering Specifications

Laboratory testing similar to what was used to determine the original engineering specifications was reused to determine the prototype specifications. In particular, device diameter was assumed as the maximum outer diameter of the prototype and it was measured using a standard metric caliper. Similarly, a measurement was taken with a standard metric ruler to determine the prototype length. Recall this length was defined as the distance from male Luer-Lok™ tip on the front end of the device to the syringe flange on the butt end of the device. Suction pressure is a specification of the prototypes that was individually considered because of our difficulty in quantifying it. See the section below for our procedure and method of calculation. Collection volume is directly related to suction pressure and was determined each of the devices with a benchmarking test. Although this may not seem like the best way to measure such a parameter, as long as adequate suction exists (and it does), we do not foresee collection volume to be an issue. Finally, device weight was determined using a standard mass scale and included all components of the design (needle, valve and syringe).

### Engineering Specifications Discussion

The measurement of the suction pressure parameter for each of the prototype designs was more of a challenge than anticipated. Our team took a numerical and idealized approach to measure the suction pressure. Specifically, we determined the volume of air that we could displace which each device (included both the volume within the syringe and the valve), then used an ideal gas law (Boyle’s law) to determine the (negative) pressure that could be achieved. This approach assumed perfectly sealed devices as well as pure air as our fluid medium, which may introduce some error in our pressure calculations, but we believe the error would be insignificant.

Hinging upon the suction pressure parameter was the collection volume. To briefly recap, in order to successfully collect tissue samples removed during a fine needle aspiration procedure, a suction pressure of about five kilopascals is required; this being the upper limit. Based on the trial-and-error of our prototypes, we anticipated that our general ballpark number for the suction pressure for each device was more than sufficient to apply enough pressure to collect adequate sample during the procedure. As a safety precaution, our team worked with a safety factor to make sure that the operator of the prototypes has the option to increase suction as he or she sees fit. To supplement our calculations on the collection volume, our team ran a benchmarking test on each of the valves. The benchmarking test consisted of creating a vacuum of 10 cc volume within the chamber of a syringe with the each valve design, and then testing the amount of water each syringe could collect. In each case, the collection of water was greater than 9 cc, which by far exceeds our specified 0.04 cc collection volume.
Lastly, the activation force necessary to apply the suction pressure for each prototype was determined. These measurements were completed using a scale from a University of Michigan Mechanical Engineering lab. For each of the two valves, we anticipate that the activation force will be substantially smaller for a device that would be mass produced. The prototypes that we constructed were somewhat limited by our ability to manufacture precise tolerances within our machine shop. Specifically, for the gate valve, we feel that an interface similar to the one that exists between the rubber seals on a plunger in a current syringe could be integrated into our gate hole and plunger design. Such a change would undoubtedly reduce the activation force substantially, accounting for the fact that our gate valve prototype did not meet the activation force target at this point.

**Why the Design Approach Works**
The approach that our team of engineers used to solve our design problem was very systematic, yet problem solving-oriented. First, by vigorously describing our design problem, our team was able to largely grasp the design problem in its entirety. Understanding every aspect of a design problem allowed our team to explore every possible solution to the problem. From this detailed problem description, our team brainstormed in a non-stress environment, exploring every solution to the problem, even those outside of the box. These ideas were then organized into categories by their solution type, and design scoring matrices were used to choose the best type of solution to our problem as well as the best device to solve our problem. Stemming from these exercises, our team was left with three valve devices (ball, gate, and pinch valves) for prototype mock-up verifications. At this stage, we wanted to make sure that the design itself was indeed feasible to produce. This step proved to be fatal for the ball valve, and it was dropped from our list. Our team of engineers has tried to consider every possible design in our approach as well as verify the best resulting design solution. Consequently, we feel that our design approach was planned, creative, and precautionary in order to achieve success in our design problem.

**MASS PRODUCTION MANUFACTURING**
Our prototypes were built using stock materials that we machined down to their final shapes. We also glued together a variety of purchased parts to create the final prototypes. If this design were to be produced in large scale, the fabrication would be very different. In this section, we will discuss what changes would need to be made to the designs for large scale manufacturing.

Using the CES Manufacturing Selector, we found that injection molding would be the best way to mass produce the polypropylene valve bodies that we fabricated from rod stock. In our prototypes, we glued polypropylene Luer-Loks™ to the ends of the valve bodies. In a mass production setting, these Luer-Loks™ would be incorporated into the valve body, so it would all be molded in one shot. This would eliminate the need for any adhesive bonding of parts.

The gate valve consists of a main body piece and the plunger piece. The body piece could be injection molded in one piece. The plunger is essentially a rod with grooves in it. These grooves serve as seats for the Viton™ tube that acts as a seal against the valve body. The grooved rod could also be injection molded as a separate piece from the gate valve body. The Viton™ seals could be cut from a continuous length of tube, and slipped over the plunger rod. This makes a total of two injection molded parts and two parts cut from a continuous tube. The assembly would be the most labor-intensive part of the process for the gate valve. The Viton™ seals would need to be slipped over the rod and into their grooves. This could probably be done as an automated process. Our sponsor gave us a target of less than $3 per unit for a disposable device. Based on the cost of a comparably sized syringe with a similar number of parts, we believe that a price under $3 would be easily attainable, were the gate valve to be mass manufactured.
The pinch valve body could also be molded as a single piece. However, we do not know what the best way to incorporate the flexible tube would be. The tube must run through the inside of the body, and be sealed to the Luer-Lok™ on either end. The solution could be to make the valve body in two halves that could snap together in the middle. The button, which was very difficult to manufacture in our machine shop, could again be simply injection molded as a single piece, and could be made much more precisely than we were able to achieve. The flexible vinyl tube and the wire lock could both be cut from continuous rolls of material. This brings the part count for the pinch valve up to five, with three injection molded parts. Again, the assembly would be the issue with mass production of this device.

The syringe lock would be a relatively easy device to build. The design is very similar to existing syringes, and could be created simply by modifying the existing mold for the syringe plunger to include a saw tooth pattern on the edge of one fin of the plunger. In our prototype, the wire lock was threaded through a hole in the syringe body. In mass production, it would probably be more cost-effective to incorporate the lock into the molded syringe body as a plastic ratchet edge. It would be fairly simple to produce and would not require any more equipment than what syringe manufacturers already have.

**PROJECT TIMELINE**

Part of the ongoing planning component of our project included the upkeep of an active Gantt chart, used to track team progress and keep us aware of upcoming deadlines. The Gantt chart also served as an accountability tool for individual team members to complete project components for which they were responsible for.

**Mileposts for the Project**

At the milepost of Design Review #2, we decided that the valve design was our first-choice (alpha) concept design. From Design Review #2 until Design Review #3, our team was responsible for thoroughly exploring the concept design for engineering feasibility and defining rigorous specifications for it. If, anywhere along this process of verifying the validity of our valve design, we were to discover that it was not a feasible design, then we would have explored one of our secondary concept designs, the electrical linear actuator design or the push slider design.

At the milepost of Design Review #3, we had produced mock-ups of the pinch, gate, and ball valve designs. We had decided that the pinch and gate valves were our first-choice designs. By Design Review #4, our team had finalized many of the details of the pinch and gate valves, and had created two final designs for the Design Expo. By the Expo, we had completed the fabrication of our prototypes, and completed all of the auxiliary assignments to be turned in with the final report, such as the Design Expo poster layout (Appendix J, page 57), materials selection assignments (Appendix K, pages 58-60), environmental impact assignment (Appendix L, pages 61-68), process selection assignment (Appendix M, pages 69-75) and the safety report (Appendix N, pages 76-109).

**Summary of Completed Tasks since Concept Selection**

In solidifying the feasibility of our alpha valve design after Design Review #2, several tasks were completed. The first step in exploring this design space was to create mock-ups of our valve design. We considered that our University of Michigan laboratory resources may not have been sufficient in producing very small designs and that we may have needed to work with outside vendors to create mock-ups of the designs. Therefore, we purchased parts from the Home Depot, Ace Hardware, McMaster-Carr, and Cole-Parmer to create mock-up prototypes of the pinch, gate, and ball valves. These mock-up prototypes allowed us to physically test our designs’ quality. We met with Professor Davenport to survey his opinion of the designs’ functions and forms. The valve designs did not prove to be insufficient for any reason, we did not need to make a mock-up of one of our secondary concept designs and repeat this.
part of the process. Since these pinch valve and gate valve design mock-ups were approved by Professor Davenport and appeared to meet our engineering specifications, we were able to move ahead in our process.

After deciding on the appropriate concept designs to produce, based on our mock-ups, our next step was to finalize all of the engineering specifications, in preparation for making the final design prototypes. We used UniGraphics NX 5.0 CAD (Computer Aided Design) software to create orthographic and three-dimensional views of the designs with specific dimensions [3].

In order to prepare our final prototypes, we needed to purchase additional parts that we did not already have. After the parts were purchased and delivered, we worked in the ME machine shop to replicate our pinch and gate valve mock-ups as polished, final prototypes for the Design Expo. After creating our final prototypes and presenting them in Design Review #4, we worked to prepare our alpha prototype presentations for the Design Expo.

While building our final prototypes, we were also working on key components for the final report. We completed the documentation for the Safety Report as we bought parts and built mock-ups. Additionally, we had to continually have our design approved for safety by Professor Im. We completed the Material Selection Assignment (functional performance and environmental performance) using the CES (Cambridge Engineering Selector) material selection software to confirm that the materials we used were the optimal choices for the specifications of our final design. We also completed the Manufacturing Process Selection Assignment, again using CES. Individually, we all completed an individual ethics essay.

As has been the case all semester, we worked as a team on all of the tasks above (with the exception of the individual ethics reports, of course). We collaborated to complete each task in our design process, working together and exchanging ideas, rather than using a divide and conquer strategy.

**Gantt Chart**
To organize the milestones of our project into a neat and organized manner, the major and minor deadlines of our fine needle aspiration project were organized into a Gantt chart. This tool allowed our team to easily visualize progress, satisfy task deadlines on time, and assign future work to prevent procrastination. See Appendix E on page 44 for our Gantt chart.

**Budget**
At the conclusion of our design process, we had spent a total of $174.96, well below the allotted budget of $400. In addition to the parts we purchased, we used syringes and needles that Professor Davenport supplied to our team free of cost. We estimated that the total value of these donated syringes was no more than $10. We have also used force transducers and mass scales from the AutoLab laboratories at the University of Michigan, provided free of cost, to take measurements for our engineering specifications. All of the parts that were used in the final prototypes were purchased from McMaster-Carr Supply Company. For the mock-ups and intermediate devices, we used parts bought from local hardware stores and Luer-Lok™ connections purchased from Cole-Parmer Instrument Company.

**RECOMMENDATIONS**
Although we did produce two working prototypes, the job of fabricating these devices was long and arduous. We believe that our designs are well conceived and are an improvement over previous devices for fine needle aspiration. Were someone to attempt to recreate our project, we recommend that they use our design, although the fabrication process could be improved.
When we designed our gate valve, our goal was for the inside of the valve body to mimic the inside of a syringe, such that the rubber seal effectively creates an airtight seal against the valve body, yet still slides easily along the length of the tube. Naturally, the best way to recreate this functionality is to use the same materials as used in a syringe. The syringe body is polypropylene, and the plunger seal is butyl rubber. We used polypropylene in our valve bodies, but we were unable to obtain butyl rubber of the correct size and shape to use in our prototypes. Instead, we used Viton™ tubing for our seals, which is firmer and less elastic than butyl rubber. We recommend that butyl rubber be pursued as the seal to lower friction and ensure an airtight seal. Also, the size and shape of the seals could be altered slightly to make the device more compact and more easily held by the fingertips. Additionally, to further reduce friction, the formula of the polypropylene could be altered. It is common practice to use additives in plastics to enhance certain properties. It would be simple to incorporate a lubricating additive to the polypropylene to lower the friction between the gate valve and the seals.

Our chief concern with the pinch valve was the locking mechanism. We struggled to fabricate the hole in the button with the required level of precision. We recommend that anyone who pursues this device further looks into better facilities for fabricating plastic parts. We used the equipment available in the ME machine shop: drills, lathes, and saws. These tools are designed for metalworking and are too large and lack the precision necessary for the fabrication of plastics. We regret not researching the plastics facilities available at the University of Michigan, and recommend that they be researched before further work.

The syringe plunger could be improved by incorporating the locking feature into the syringe body. The plunger itself is easy enough to produce, but our device uses a wire wrapped around the syringe body to catch the saw teeth. It would be an improvement if some other method of retaining the saw teeth could be developed and built into the syringe body; that would reduce assembly time and complexity of the part. One simple solution would be a partial lid that covers a small sector of the circular area of the syringe body. When a saw tooth is rotated over the lid, it would catch the edge and not be able to proceed downward.

These are a few changes that could be made to the design and fabrication processes of the devices that would lead to better quality and repeatability of the designs.

CONCLUSIONS

The current devices for performing fine needle aspirations do not meet doctors’ needs as they perform this procedure. The devices typically use a pistol grip that requires movement of the large muscle groups in the hand and arm to maneuver the needle tip. Our sponsor, Professor Robertson Davenport, M.D., of the University of Michigan Hospital’s Department of Pathology, asked us to design a new device to allow doctors to use the finger tips to position and activate the device. Using the fingertips allows for much finer control over the positioning of the needle during the procedure, and also increases tactile feedback from the tissue. That way, the doctor can more accurately direct the needle, and can also feel the consistency of the tissue through which the needle is passing.

We carried out the steps of the standard design process, researching and defining the problem, then defining the success criteria for the design and the engineering specifications. Then, through brainstorming and some broadening of the design concepts, we developed a wide range of concepts to fulfill our sponsor’s request. We then went through the process of concept selection, narrowing the field down to the most feasible concepts. The concept that emerged as the design to be pursued was a valve design, which decouples the action of creating a vacuum in the syringe and the movement of the needle tip. The vacuum is created in a syringe with a closed valve before the procedure begins, and the vacuum
is activated to the needle tip once it has been inserted into the skin. The device is held like a pen, and the vacuum is activated with a single button press.

We developed two distinct valves that can be used in this procedure, a gate valve and a pinch valve. The gate valve’s design is more likely to be produced in a mass manufacturing setting since it can be injection molded in two main pieces with two seals. The gate valve also has an advantage in the way it operates. With the gate valve, the doctor does not need to remove the needle after the procedure to express the sample. The pinch valve is simpler, but would be more difficult to manufacture, since it requires a flexible piece inside of a rigid shell. All of the details of these devices can be found in the body of this report. After developing the designs on paper, we fabricated prototypes using stock materials. The prototypes serve as proofs of concept that the designs function the way we wanted, and working models to show interested parties.

Both of our final valve prototypes meet the engineering requirements we set out to achieve at the beginning of the design process (gate valve activation force aside; explanation in validation section on page 26). More importantly, they satisfied the goals that Professor Davenport has asked us to achieve. They can be easily held and manipulated with a single hand, and the vacuum can be activated or deactivated with one finger. These devices mark a significant improvement over the existing options in terms of fine motor control, tactile feedback, and needle stability.

ACKNOWLEDGEMENTS

We would like to thank and acknowledge the following individuals for their contributions throughout the course of our project:

Professor Roberston Davenport, M.D.
Doctor Stewart Knoepp, M.D.
Professor Hong G. Im
Professor Steven Skerlos
Robert Coury
Marv Cressy
Daniel Johnson
Tom Bress
Max Bajcz
REFERENCES AND INFORMATION SOURCES


APPENDIX A: Technical Benchmarks

A.1: Tao Aspirator® [4]

A.2: Cytec® System [5]
### Reference Human Strength Levels for $F_i$

<table>
<thead>
<tr>
<th></th>
<th>maximum force (men)</th>
<th>max. hand grip strength</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thumb</td>
<td>164N (133 - 191N range)</td>
<td></td>
</tr>
<tr>
<td>Finger</td>
<td>106N (75 - 137N range)</td>
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</table>

<table>
<thead>
<tr>
<th></th>
<th>maximum finger gripping strength</th>
</tr>
</thead>
<tbody>
<tr>
<td>weak woman</td>
<td>40N</td>
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<tr>
<td>weak man</td>
<td>57</td>
</tr>
<tr>
<td>strong woman</td>
<td>93</td>
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<tr>
<td>strong man</td>
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### Reference Operational Forces

<p>| | |</p>
<table>
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<tr>
<th></th>
<th></th>
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<tbody>
<tr>
<td>Typewriter key</td>
<td>0.25 - 1.5N</td>
</tr>
<tr>
<td>Push buttons</td>
<td>1.1 - 8.3N range</td>
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</table>
APPENDIX C: Concept Sketches

C.1 Valve Design

C.2 Electrical Linear Actuator

C.3a Push Slider Sketch 1
C.3b Push Slider Sketch 2

C.4 Direct Slider

C.5 Spring and Locking Slider

C.6 Spring Loaded Design
C.7 Lever Arm Design

C.8 Live Hinge Design

C.9 Rack and Pinion Design
C.10 Shoe Pump Design

C.11 Diaphragm Design

1. Set pressure
2. Break up tissue / get into needle
3. Release pressure / retain tissue

Diaphragm Valve - disposable

- Valve holds air pressure below
C.12 Ball Valve Design
C.13 Gate Valve Design
C.14 Pinch Valve Design

[Diagram images showing various views and sections of a pinch valve design, including labeled parts such as flexible tube, rocker switch, and sections A-A and B-B.]
APPENDIX D: Concept Selection Aids

D.1 Quality Function Deployment

Fine Needle Aspiration QFD

<table>
<thead>
<tr>
<th>Customer Needs</th>
<th>Weight</th>
<th>User-Loc Cap</th>
<th>Device Diameter</th>
<th>Length</th>
<th>Reusability</th>
<th>Capacity</th>
<th>Material</th>
<th>Color</th>
<th>Device Flexibility</th>
<th>User Stability</th>
<th>Canvco</th>
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<th>No</th>
<th>Outside</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fine Motor Controlability</td>
<td>9</td>
<td>o</td>
<td>●</td>
<td>●</td>
<td>△</td>
<td>△</td>
<td>o</td>
<td>1</td>
<td>3</td>
<td>5</td>
<td></td>
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<td>o</td>
<td>●</td>
<td>●</td>
<td>△</td>
<td>△</td>
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<td>User-controlled suction</td>
<td>3</td>
<td>o</td>
<td>●</td>
<td>●</td>
<td>o</td>
<td>5</td>
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<td>Simplicity of use</td>
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<td>o</td>
<td>o</td>
<td>△</td>
<td>o</td>
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<td>●</td>
<td>o</td>
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<td>△</td>
<td>o</td>
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<td>△</td>
<td>△</td>
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<td>Expresses sample easily</td>
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<td>△</td>
<td>△</td>
<td>o</td>
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Raw score: 17 16 15 14 14 15 3 15 21
Scaled: 1 0.76 0.71 0.76 0.67 0.67 0.76 0.14 0.71 1
Relative Weight: 1% 1% 1% 1% 1% 1% 1% 2% 1% 1%
Rank: 2 3 6 3 8 8 3 10 6 1

Requirement Benchmarking

Best in Class: A\E
Average: A\E
Worst in Class: Not Yet Completed

Legend:
- ○: Benefit
- ●: Satisfactory
- △: Satisfactory
- #: Conflict
### D.2: Design Scoring Matrix

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<tr>
<td>Device Diameter</td>
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<tr>
<td>Length (grip to needle tip)</td>
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<td>Device Flexibility</td>
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<td>Manufacturability</td>
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<td>User Range of Motion</td>
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<td>Relative Market Price</td>
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### D.3: Valve Scoring Matrix

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<td>Simplicity</td>
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### APPENDIX E: Gantt Chart

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<tr>
<td>Research SIM Devices</td>
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<td>Thu 1/23/09</td>
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<tr>
<td>Literature and Patent Research</td>
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<td>Thu 1/23/09</td>
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<td>Individual Meeting</td>
<td>Tue 1/20/09</td>
<td>Tue 1/23/09</td>
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<tr>
<td>Group Discussion - Specifications and Literature Review</td>
<td>Thu 1/22/09</td>
<td>Thu 1/22/09</td>
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<tr>
<td>CFD development</td>
<td>Thu 1/22/09</td>
<td>Thu 1/22/09</td>
</tr>
<tr>
<td>Individual Meeting</td>
<td>Tue 1/27/09</td>
<td>Tue 1/27/09</td>
</tr>
<tr>
<td>Design Review #1</td>
<td>Thu 1/28/09</td>
<td>Thu 1/28/09</td>
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<tr>
<td>Take remaining engineering/force measurements in ME</td>
<td>Mon 2/2/09</td>
<td>Mon 2/2/09</td>
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<tr>
<td>Brainstorm Additional FEA Concepts</td>
<td>Thu 1/28/09</td>
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<td>Individual Meeting</td>
<td>Tue 2/3/09</td>
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<tr>
<td>Progress meeting with Dr. Davenport</td>
<td>Wed 2/4/09</td>
<td>Wed 2/4/09</td>
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<tr>
<td>Group Discussion - Additional Concept Generation</td>
<td>Thu 2/5/09</td>
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<tr>
<td>Choose Material (Cambridge Engineering Selector)</td>
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<tr>
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<tr>
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<td>Individual Meeting</td>
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<tr>
<td>Engineering Drawings CAD</td>
<td>Thu 2/12/09</td>
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<td>Push Chart</td>
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<td>Design Review #2 - Concept Generation and Selection</td>
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<td>Winter Break</td>
<td>Mon 2/23/09</td>
<td>Fri 2/27/09</td>
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<td>Do CES analysis</td>
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<td>Wed 3/4/09</td>
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<tr>
<td>Create Mock-Up</td>
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<tr>
<td>Create CAD models</td>
<td>Mon 3/6/09</td>
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<td>Safety Review</td>
<td>Mon 3/16/09</td>
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<td>Tue 3/17/09</td>
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<td>Design Review #3 - Final Design</td>
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<td>Purchase parts for final pinch and gate valve design</td>
<td>Fri 3/27/09</td>
<td>Fri 3/27/09</td>
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<td>Do Material Selection Assignment (functional performance)</td>
<td>Mon 3/30/09</td>
<td>Fri 4/3/09</td>
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<tr>
<td>Do Material Selection Assignment (environmental performance)</td>
<td>Mon 3/30/05</td>
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<td>Do Manufacturing Process Selection Assignment</td>
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<td>Fri 4/3/09</td>
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<tr>
<td>Build Final Prototypes</td>
<td>Tue 3/3/09</td>
<td>Wed 3/4/09</td>
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<tr>
<td>Review prototypes with Professor Davenport</td>
<td>Fri 4/3/09</td>
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<tr>
<td>Design Review #4 - Alpha Prototype</td>
<td>Tue 4/7/09</td>
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<td>Prepare presentation and pinch and gate valves for the prototype</td>
<td>Tue 4/7/09</td>
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<td>Design Expo</td>
<td>Thu 4/10/09</td>
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<td>Individual Ethics Essay due</td>
<td>Tue 4/21/09</td>
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<td>Final Report and Safety ReportDue</td>
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<td>Prototype Delivered</td>
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## APPENDIX F: Bill of Materials

### Materials List

**FINE NEEDLE ASPIRATION**  
20-Apr-09

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45
APPENDIX G: Final Prototype Drafting [5]

G.1 Gate Valve Design

G.2a Gate Valve Dowel Design
G.2b Gate Valve Dowel Design

G.3 Pinch Valve Design
G.4 Pinch Valve Push button Design
APPENDIX H: Pinch Valve Operation and Procedural Instructions

Step 1: Locate needle and the male Luer-Lok™ of the pinch valve casing.

Step 2: Attach needle to the male Luer-Lok™ of the pinch valve casing by inserting and turning clockwise.

Step 3: Attach syringe to opposite end of the pinch valve casing by inserting and turning clockwise.
Step 4: Close the valve by pushing down on the push button and lock the button by pulling back on the button in the pushed position.

Step 5: Now that the valve is sealed, to create a vacuum in the syringe barrel:
1. Twist the syringe plunger counterclockwise
2. Pull back to the desired suction level
3. Twist the plunger clockwise to lock the plunger within a sawtooth slot.

Step 6: Device is ready for procedure.
Step 7: Insert needle into lump and release lock to apply vacuum to the needle tip by pressing down and pushing forward on the push button then releasing.

Step 8: After sufficient amount of tissue cells are collected, deactivate vacuum (before removing needle) by pressing and locking the button as in step 4.

Step 9: Remove needle from the lump.
Step 10: Remove needle containing tissue sample from valve casing by twisting counter clockwise.

Step 11: Connect needle to another syringe to express tissue onto slide.
APPENDIX I: Gate Valve Operation and Procedural Instructions

Step 1: Locate needle and the male Luer-Lok™ of the gate valve casing

Step 2: Attach needle to the male Luer-Lok™ of the gate valve casing by inserting and turning clockwise

Step 3: Attach syringe to opposite end of the gate valve casing by inserting and turning clockwise
Step 4: Gate valve packaged in sealed position. To create a vacuum in the syringe barrel:
1. Twist the syringe plunger counterclockwise
2. Pull back to the desired suction level
3. Twist the plunger clockwise to lock the plunger within a sawtooth slot.

Step 5: Device is ready for procedure.

Step 6: Insert needle into lump.
Step 7: Activate vacuum to needle tip by pushing gate plunger to middle position (second line).

Step 8: After sufficient amount of tissue cells are collected, deactivate vacuum (before removing needle) by pressing gate valve button entirely down.

Step 9: Remove needle from the lump.
Step 10: Unlock plunger to prepare for tissue extraction by twisting plunger clockwise.

Step 11: Flip gate valve 180 degrees and return gate plunger to middle position by pushing the exposed button (uncapped side of plunger) until it is flush with the bottom of the valve casing.

Step 12: Express tissue sample onto a slide for analysis by forcing plunger down.
APPENDIX J: Design Expo Poster

FINE NEEDLE ASPIRATION DEVICE

ME 450 TEAM 4
Nathan Brown, Mary Kay DuBay, Jeff Otto, Joel Van Sloten

WHAT IS FINE NEEDLE ASPIRATION?

A biopsy technique in which a needle is inserted into a potentially cancerous lump or tumor, and tissue is extracted by a vacuum (typically created by a syringe) at the needle tip.

WHAT’S WRONG WITH THE CURRENT DEVICES?

Even a small movement by the doctor’s hand creates a large movement at the needle tip.

Our Goal:

Create a device that can be held and manipulated with the fingertips.

Customer Requirements:

- Fine motor controllable
- Operated with one (dominant) hand
- User-controlled suction activation and release
- Expresses sample easily
- Simple to use
- Sterile
- Safe for operator
- Affordable

The New Design:

- Controlled with fingertips
- Held like a pen in either hand
- Valve controls vacuum suction to needle tip
- Disposable
- Incorporates a ratcheting locking plunger to maintain a vacuum in the syringe

We developed two valve designs that fulfill the required specifications.
APPENDIX K: Materials Functional Performance

Materials Selection Assignment
Functional Performance
Project 4: Fine Needle Aspiration

Flexible tube for pinch valve

Function: This flexible tube, inside of the pinch valve casing, connects between the syringe needle and the syringe tip. It serves as a flexible passageway for air and tissue samples in the valve. By pinching/creasing the flexible tube with the user-controlled pinch valve button, the sealed, flexible tube holds and releases air pressure from the syringe.

Objective: minimize cost (must be below $3 per total design unit)
Constraints: must flex to seal (~35A durometer)

Material Indices:
Since we cannot define the durometer in terms of other parameters, we were not able to define material indices for the flexible tube for the pinch valve. We decided to simply limit our search to elastomers in the durometer range of Shore 30A-40A. The proof-of-concept we built used a latex tube of 35A, which worked very well, so we tried to match that (while avoiding latex for allergy issues).

The objective is to minimize cost, so to select our first five options, we looked at the five cheapest materials in the range of 30A-40A.

CES Top 5 Material Choices: 1: PVC-elastomer
2: Butadiene rubber (nitrile)
   Softest readily available grade of nitrile tube is about 60A
3: Ethylene propylene diene (EPDM)
   Softest readily available grade of EPDM tube is about 60A
4: SIS (Styrene Isoprene Styrene)
   Not readily available
5: Butyl Rubber
   Expensive in tube form

Final Choice: PVC
PVC is readily available, low cost, and meets the hardness/durometer requirement. Although CES rated Butadiene Rubber (nitrile) and Ethylene Propylene Diene (EPDM) within the 30-40 A hardness range, they were only available for purchase at 60 A hardness in reality. The Styrene Isoprene Styrene (SIS) is not readily available. The Butyl Rubber highly exceeds our minimal cost objective. According to CES, PVC is our best option, and it also is available on McMaster in the right size and hardness.
Gate Valve Casing

Function: This valve casing/structure connects between the syringe needle and the syringe tip. It serves as a strong, rigid structure to withstand user forces and internal pressures.

Objective: minimize cost (must be below $3 per total design unit)
Constraints:
- maximum mass specification = 250 g
- fixed volume = 0.0000135 m³
- minimum ultimate tensile strength = 450 kPa

Material Indices:
- ↑Material Index = ↑Performance
- Cost = V*c (V = volume, c = unit cost)
- m = ρ*V (m = mass, ρ = density)
- Eliminated free variables: V
- Material Indices: M = ρ/c (to be maximized)
- Cost = m*(c/ρ)
CES Top 5 Material Choices:

1: Polypropylene
   Our first-choice material

2: Polybutylene (PB)
   Only bonding adhesives, manifolds, and tube fittings available

3: Polymethylpentene (PMP)
   Only film readily available; would have to special order

4: Thermoplastic Polyolefin Elastomer (PP+EP(D)M)
   Only plastic welding rod available

5: Ethyl Methyl Acrylate (EMA)
   Only available by special order overseas

Final Choice: Polypropylene is our first choice because it meets all of our CES objectives and constraints, and is readily available to order in the rod form we need. Even if we were to order Polymethylpentene (PMP) in its available film form, Thermoplastic Polyolefin Elastomer (PP+ER(D)M) in its available plastic welding rod form, or Polybutylene (PB) in its available bonding adhesive/manifold/tube fitting form, we couldn’t machine the material to our desired form. It was possible for us to special order a Polymethylpentene (PMP) rod or an Ethyl Methyl Acrylate (EMA) rod, but the added cost and time of shipping these special orders from overseas companies was not worth it. Polypropylene will do the job.
APPENDIX L: Environmental Impact

Flexible tube for pinch valve

1. Materials Considered:
   Polyvinylchloride (PVC) (Shore 35a)
   Ethylene Propylene Diene M (EPDM)

2. Determination of Mass in Final Design:
   Determine the volume of the tubing necessary using geometry
   Determine the density of each material using Cambridge Engineering Selector
   Multiply volume by density of selected materials to determine mass

   Volume of tubing needed: \( \forall = \left( \pi \left( \frac{D_o}{2} \right)^2 - \pi \left( \frac{D_i}{2} \right)^2 \right) \times L \)
   \[ \forall = \left( \pi \left( \frac{1/4}{2} \right)^2 - \pi \left( \frac{1/8}{2} \right)^2 \right) \times 1.05314961 \text{ [in}^3\text{]} \]
   \( \forall = 0.0387723 \text{ [in}^3\text{]} \)

   Density of Polyvinylchloride (PVC) (Shore 35a): \( \rho = 0.03955 \text{ [lb/in}^3\text{]} \)
   Density of Ethylene Propylene Diene M (EPDM): \( \rho = 0.03145 \text{ [lb/in}^3\text{]} \)

   Mass of Polyvinylchloride (PVC) (Shore 35a) tubing:
   \[ \text{mass} = \rho \times \forall \]
   \[ \text{mass} = 0.03955 \text{ [lb/in}^3\text{]} \times 0.0387723 \text{ [in}^3\text{]} \]
   \( \text{mass} = 0.001533 \text{ [lb]} (\approx 0.0006954 \text{ kg}) \)

   Mass of Ethylene Propylene Diene M (EPDM):
   \[ \text{mass} = \rho \times \forall \]
   \[ \text{mass} = 0.03145 \text{ [lb/in}^3\text{]} \times 0.0387723 \text{ [in}^3\text{]} \]
   \( \text{mass} = 0.0012194 \text{ [lb]} (\approx 0.000553 \text{ kg}) \)

3. Corresponding Materials in SimaPro
   Material Polyvinylchloride (PVC)
   SimaPro Material: PVC film E

   Ethylene Propylene Diene M (EPDM)
   SimaPro Material: EPDM rubber ETH S
4. Calculate Total Pollution Emissions

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5. Determine which material is more environmentally damaging

The EcoIndicator 99 damage classification reveals that using a Polyvinylchloride film E tubing results in fewer grams of damaging effects on the environment compared to that of the Ethylene Propylene Diene M rubber ETH S. Total mass of damage to the environment caused by Polypropylene totaled 15.4 grams, compared to 16 grams caused by Propylene Diene M rubber ETH S. The damage cause by Polyvinylchloride film E is focused primarily on raw and air pollutants. For the air, water, and waste categories, Polyvinylchloride film E was more damaging than the Ethylene Propylene Diene M rubber ETH S, but the amount of pollution was quite insignificant compared to raw pollution. Ethylene Propylene Diene M rubber ETH S was more damaging in the raw and soil categories.

Based on the EI99 point values, the Human Health is the most notable and important meta-category, followed by Resources and then Ecosystem Quality. For both materials, Human Health has point values approximately two to six times greater that of Resources and ten to twenty times that of Ecosystem Quality.

6. Since the application of both materials is for a disposable device, each will have the same lifecycle. Because of this there is no reason to change the current evaluation and EPDM is still considered the most environmentally unfriendly material.
Casing for Gate Valve

1. Materials Considered:
   - Polyvinylchloride (PVC)
   - Polybutylene

2. Determination of Mass in Final Design:
   Computationally determine the volume of the casing using UGS NX 5.0
   Determine the density of each material using Cambridge Engineering Selector
   Multiply volume by density of selected materials to determine mass

   Volume of tubing needed: \( V = 0.827209 \text{ in}^3 \)

   Density of Polyvinylchloride (PVC): \( \rho = 0.03252315 \frac{lb}{in^3} \)
   Density of Polybutylene: \( \rho = 0.03235 \frac{lb}{in^3} \)

   Mass of Polyvinylchloride (PVC) tubing:
   \[
   mass = \rho \times V
   \]
   \[
   mass = 0.03252315 \frac{lb}{in^3} \times 0.827209 \text{ in}^3
   \]
   \[
   mass = 0.026903 \text{ lb} \approx 0.012203 \text{ kg}
   \]

   Mass of Polybutylene:
   \[
   mass = \rho \times V
   \]
   \[
   mass = 0.03235 \frac{lb}{in^3} \times 0.827209 \text{ in}^3
   \]
   \[
   mass = 0.0267602 \text{ lb} \approx 0.0121382 \text{ kg}
   \]

3. Corresponding Materials in SimaPro
   - Material: Polyvinylchloride (PVC)
     SimaPro Material: PVC I
   - Material: Polybutylene
     SimaPro Material: PB I

4. Calculate Total Pollution Emissions

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5. The EcoIndicator 99 damage assessment indicates that Polypropylene is less harmful to the environment by generating less pollution than amount created by the Polybutylene. Specifically the Polypropylene generates a total of approximately 750 g of pollution, while the Polybutylene generates 1290 g. The major source of pollution for both materials comes from raw state pollutants; however the Polybutylene produces a rather significant quantity of waste pollutants as well. The only category where Polypropylene generates more waste than the Polybutylene is air pollutants, however these pollutants account for only a small percentage of the total pollution.

Based on the EI99 point values, it is rather obvious that Human Health is the most notable and important meta-category. For both materials, Human Health has point values approximately ten times greater than Ecosystem Quality and ten to one hundred times that of Resources.

6. Since the application of both materials is for a disposable device, each will have the same full life cycle. Because of this there is the initial material assessment remains unchanged and Polybutylene is still considered the most environmentally unfriendly material.
We assumed that a batch size of 10,000 – 100,000 units of our Fine Needle Aspiration device will be useful to society. The best manufacturing process for the PVC flexible tube for the pinch valve was polymer extrusion. The best manufacturing process for the polypropylene valve casing for the gate valve was injection molding. These “best” processes were chosen using the CES Manufacturing Selector, comparing legitimate values for mass range, section thickness range, tolerance range, and economic batch size. The graphs below (CES printouts) justify these choices.

Flexible Tube — polymer extrusion

Since this is a piece to be produced, it is a primary shaping process. We defined the tolerance for the tube radius to be 0.02 – 0.0055 in. This number is somewhat arbitrary, since the tube is flexible, and will stretch if it is the wrong size, but for the sake of eliminating processes with high or low (and expensive) tolerances, we chose the above range, shown as a box on the graph below.
Next, we defined the range of section thickness, which in this case is the wall thickness of the tube, 0.0625 in. Including a tolerance of 0.02 in, the range of section thickness becomes 0.0425 – 0.0825 in, shown as a box on the chart below. Also, the part is a circular prismatic, which is reflected on the chart as being on the right half of the chart.

Next, since the part is a short tube of constant diameter, it makes sense to produce as a continuous process and cut the continuous tube to the correct length. Therefore, we added a limit stage to the CES selection template that limited the processes to only continuous processes.
Finally, we determined what the mass of a continuously produced tube would be, based on a batch size of 10,000 to 100,000 units produced. Based on the number of tubes needed and the density of the material, we found that we would need to produce between 15lb and 150lb of tubing. The final stage, shown below, limits the processes to elastomers (since the part must be flexible), with a unit size of 15lb to 150lb.

![Diagram showing mass range and polymer extrusion](image)

Polymer extrusion is the only process that passed all four of the above stages, so we determined that it would be the best manufacturing option for the flexible tube.

In reality, the tube would probably be purchased from a supplier. Tubes are already mass-produced, and it would likely be more cost effective to buy a tube from a manufacturer that is already set up to produce large amounts. And the manufacturer, having done an analysis similar to the one we did above, would produce the tube by polymer extrusion.
Valve Casing – Injection molding

Again, the valve casing needs to be produced from some type of material stock, so it is a primary process. To determine the tolerance for the part, we looked to the sections of the casing that would need to be most precise, which are the Luer tips. According to ISO 594-1 (1986) for a 6% Luer-Lok standard conical fitting, the diameter tolerance for semi-rigid material is given as 0.102mm, which equals 0.004 in. We used a range of 0.003 – 0.005 in, as shown by the box below.
The part is a 3-D hollow part (a tube with a perpendicular hole), and the thickness ranges from 0.25 – 0.5 in. We displayed this range of thicknesses on the chart below as a box.
Next, we defined the material as a thermoplastic (x-axis of the chart below) and calculated the mass for each part, based on its volume and the density of polypropylene, our chosen material. The mass range is 0.03-0.06 lb per unit, accounting for potential variations in the design to optimize mass manufacturing. This range is boxed on the chart below.
Finally, defined the parts as discrete parts, rather than continuous, and selected a batch size of 10,000 to 100,000, as shown by the box below.

The only process that passed all of the above stages for the valve casing was injection molding.
APPENDIX N: Safety Report

ME 450 Safety Reporting: Winter 2009

Project #: 4
Date: 4/21/2009
Report Version #: A
Project Title: Fine Needle Aspiration Device

Nathan Brown, Mary Kay DuBay, Jeff Otto, Joel Van Sloten

Team Member Names: nathanab, mkdubay, jmotto, javs

Team Member Uniquenames: nathanab, mkdubay, jmotto, javs

Attach your Safety Report to this cover page and instructions found on Pages 2 and 3.

The Safety Report is to be completed by your team and must be approved by your section instructor (or approved substitute) prior to any hands-on experimentation, manufacturing or testing of your prototype.

The safety hazards inherent in your experimental plans, component selection, manufacturing methods, assembly techniques, and testing must be expressed and evaluated before any hands-on work with safety consequences will be allowed to proceed.

The purpose of this safety report is to assure that you have thought through your hands-on work before it begins, and that you have shared your plans with your Section Instructor. You may submit more than one version. This will likely be necessary as your project evolves.

APPROVAL:

Name: Hong G. Im

Signature: __________________________

Date: __________________________
1 EXECUTIVE SUMMARY
This report covers the safety considerations for the design, manufacturing, and use of the devices designed by ME 450 Team 4.

Experimental Plans Prior to Design Completion
We will perform some very basic measurements on existing devices and equipment for fine needle aspiration to give us a baseline against which we can compare our final designs. The design process will be mostly completed through experimental testing of proof-of-concept devices and comparing the performance of those mock-ups to the desired outcome.

Design Elements
This report contains a list of all purchased materials for the project and an FMEA matrix for all of the components used in the final design. A DesignSafe report is included for each of the syringe, gate valve, pinch valve, and the device assembly. The major risks for each component are addressed with what preventative measures we have taken to eliminate those risks, as well as the measures we think will reduce risk in the use of the device. The CAD drawings for each designed component are included as well.

Manufacturing and Assembly Elements
The majority of the manufacturing undertaken for the project will be cutting, drilling, and joining. The processes that required each type of machining are described in this report. Since most of the parts are plastics, the machining feeds and speeds are flexible, and the safety concerns are not as great as if the parts had been metal. The assembly issues will be intensified because of the small and precise nature of the parts required for the design.

Design Testing and Validation
The design validation will happen step by step as we proceed through different stages and test individual aspects of the valves along the way. Most of these tests will not be quantitative as much as qualitative. To ensure that our engineering specifications are met, we will measure various physical properties of the devices.
2 EXPERIMENTATION PLANS PRIOR TO DESIGN COMPLETION

The first thing we will test is the current setup used by doctors to perform fine needle aspirations. The current equipment is typically a 10cc syringe with a needle on the tip, held by a pistol-grip type syringe holder. We will begin by testing the force required to pull back on the syringe plunger in a 10cc syringe. We have contacted Tom Bress of the ME 395/495 labs, who has agreed to help us perform these experiments. We also would like to compare those results to the forces required to pull back the plunger on smaller syringes, such as a 3cc and 1cc syringe. These force measurements will show us what our target value is for activation force on our device. Ideally, we could make the activation force less than what already exists in the current setup. Since the device will ultimately be controlled with the fingertips, the activation force will be important. We do not foresee any safety risks in the measurement of the mass, dimensions, and force required to pull back a syringe.

3 PURCHASED COMPONENT AND MATERIAL INVENTORY

Appendix A is a list of all the materials purchased for the project. Some will be used in building mock-ups and proof-of-concepts, and some will be used for the final prototypes. The materials are organized by their application, and each includes a description of its final use. None of these materials are inherently dangerous in their handling or use, unless used inappropriately. The only components that could be considered hazardous are the ScotchWeld DP-8005 adhesive and the fiberglass resin, both of which need to be used in a well-ventilated area, as it has fairly volatile fumes. The plastic epoxy and the silicone sealant should be well-ventilated as well, although they do not pose a threat.

FMEA Analysis Results

We performed an FMEA analysis on each of the purchased components to be used in the final prototypes. The table we used for the FMEA is shown in Appendix B. The FMEA results showed that the components that were most likely to cause a failure or safety hazard are the Viton tube in the gate valve and the PVC tube in the pinch valve. These, not surprisingly, are the only flexible parts in the either of the designs. They are the most likely to cause an issue because the both relate directly to the functional performance of the devices, and they also are the primary defense in ensuring that blood does not leak from the valves. The Viton tube in the gate valve serves as the seal that slides in the valve, allowing pressure to reach the needle or blocking it. If it fails, the valve will not hold pressure, and will not function correctly. To mitigate this issue, we have decided that it would be best to test each valve before shipping in a production setting. This testing could ensure that no valves with bad seals would go out. Since it is easy enough for us to simply look at the seals and determine whether they are damaged, we do not foresee and safety or failure issues for our prototype seal.

The PVC tube in the pinch valve is the vessel through which the vacuum flows, allowing the vacuum to reach the needle tip. The tube is pinched by a rounded rod, sealing off the passage and not allowing vacuum to pass. If the rod should tear the tube, the valve would not hold pressure. If the sample of tissue was already collected at the time of tearing, some blood could exit through the failure in the tube, and the operator could be exposed to the blood. We can ensure in our prototype that the tube is intact by inspecting it before installation. However, once the procedure is in progress, an operator would have no indication that the tube had ruptured, which would lead to a test that did not collect any sample.
4 CAD DRAWINGS AND DESIGNSAFE ANALYSIS

Pinch Valve
Pinch Valve Button Detail

PINCH BUTTON
ALL RADI 1.0
Gate Valve
Gate Valve Rod Details
DesignSafe Analysis
Our analysis with DesignSafe shows that the highest safety risks are related to proper use. For the assembly as a whole, the highest risks were from mechanical failure of the device, and the potential for contact with blood-borne disease. Both valve designs should not experience mechanical failure in normal use. Therefore, if they were to actually fail, it would be due to improper use, if the devices were abused or used in another way than their intended use. The best way to ensure that abuse doesn’t happen is to make sure that the users understand the proper way to employ the device. This is standard training, and could be included in the form of an instruction booklet, or even a diagram showing how to correctly operate the devices.

The other issue is that of blood-borne disease. This could happen if the seals or the flexible tube failed, depending on the valve type. In both cases, there is a small chance that blood could seep or drip out of the valve after the seal was ruptured. In either case, the user would be protected from any disease by wearing the appropriate equipment. Gloves are standard for doctors anyway, and would prevent their hands from coming in direct contact with patient blood. Also, respiratory masks are readily available in hospitals, so if the doctor wishes to protect himself further, that is always an option.

The full DesignSafe reports can be found in Appendix C.

5 MANUFACTURING PROCESSES
This section lists the manufacturing procedure for each mock-up and for each final prototype. The process is listed with the part to which it applies.

Manufacturing Procedure List—Mock-Ups
Pinch Valve
1. Size and cut vinyl tubing (1/2 x 0.170)
2. Cut nylon tee to size
3. Trim off male and female luer lock (with cone) from a syringe
4. Cut two body syringe parts to size
   a. Trim circular hole for tee top
5. Feed tube through tee
6. Seal tube to luer lock connections with silicon
7. Attach body to luer lok parts with plastic epoxy
8. Attach center body to body interface with plastic epoxy

Gate Valve
1. Cut syringe 10cc syringe and 1cc syringe to desired valve length
2. Insert 1cc syringe into 10cc syringe, center and fill void with Fiberglass Resin
3. Cut slot for gate valve
4. Cut out gate from acrylic sheet or dowel
5. Trim hole in sheet or dowel
6. Coat gate valve with Plasti Dip
7. Insert into slot and troubleshoot
**Ball Valve**
1. Cut vinyl tubing (5/8 x 1/2) to desired length
2. Machine rubber cork to correct outer diameter and inner shape
3. Attach rubber cork to inside of vinyl tubing with adhesive
4. Attach tubing to male Luer lock with adhesive
5. Attach spring to ball bearing (solder?)
6. Attach opposite edge of spring to Luer lock
7. Attach other Luer lock to vinyl tubing
8. Troubleshoot

We will assemble all components in the ME machine shop, as we do not have appropriate machines and tools to assemble the mock-ups on our own.

**Manufacturing Procedure List—Final Prototypes**

**Pinch Valve**
1. Cut a piece of ¾” polypropylene cylinder, 30.8mm long
2. Drill through length of ¾” polypropylene cylinder, 7.5mm diameter (500 RPM drill)
3. Drill through length of ¾” polypropylene cylinder, 12mm diameter, 2mm deep (500 RPM drill) for luer-loks
4. Drill through diameter of ¾” polypropylene cylinder, 6.0mm diameter, 21mm deep (500 RPM drill)
5. Cut vinyl tubing (½” x 0.170”)
6. Run vinyl tubing through length of ¾” polypropylene cylinder
7. Seal male and female luer-loks to end of vinyl tube with Scotch-Weld DP8005
8. Seal ¾” polypropylene cylinder ends to luer-lok connection with Scotch-Weld DP8005

**Pinch Valve Push Button**
9. Cut 16 gauge galvanized steel wire, 10mm long
10. Drill hole through ¾” polypropylene cylinder to run steel wire through, 1/16” diameter
11. Cut 6.8mm polypropylene cylinder, 20mm long
12. Drill J-shape out of 6.8mm polypropylene cylinder
13. Run steel wire through one ¾” polypropylene cylinder hole, through the J-shape of 6.8mm polypropylene cylinder, through other ¾” polypropylene cylinder to secure push button
14. Cut 8mm diameter button cap, 2mm deep
15. Adhere 8mm diameter button cap to 6.8mm polypropylene cylinder with Scotch-Weld DP8005

**Gate Valve**
1. Cut a piece of 1” polypropylene cylinder, 30.6mm long
2. Drill through length of 1” polypropylene cylinder, 1.6mm diameter (500 RPM drill)
3. Drill through length of 1” polypropylene cylinder, 12mm diameter, 2mm deep (500 RPM drill) for luer-loks
4. Drill through diameter of 1” polypropylene cylinder, 6.8mm diameter, (500 RPM drill)
5. Seal 1” polypropylene cylinder ends to luer-lok connection with Scotch-Weld DP8005
**Gate Valve Button**
6. Cut 2 pieces of ¼” OD, 1/8” ID Viton rubber, 2.5mm long and 4.5mm long
7. Cut 6.8mm diameter polypropylene cylinder, 30.7mm long
8. Etch out holders on 6.8mm diameter polypropylene cylinder for Viton rubber seals
9. Slide Viton rubber seals onto 6.8mm diameter polypropylene cylinder
10. Insert gate valve button into through hole of 1” polypropylene cylinder with Banana Boat dark tanning oil lubrication
11. Cut 8mm diameter button cap, 2mm deep
12. Adhere 8mm diameter button cap to 6.8mm polypropylene cylinder with Scotch-Weld DP8005

**Syringe Plunger**
1. Remove standard syringe plunger from syringe barrel
2. Carve a saw-tooth edge out of one syringe plunger face with an Exact-o knife
3. Drill two 1/16” holes out of syringe barrel flange
4. Run 16 gauge galvanized steel through flange holes
5. Re-insert saw-tooth syringe plunger into syringe barrel

We will assemble all components in the ME machine shop, as appropriate machines and tools to assemble the prototypes are not available elsewhere.

The cutting processes listed above are either cutting on a bandsaw or using a hobby knife. For the larger plastic parts that need to be cut to length, the pieces will be cut on the bandsaw with a speed of about 300 fpm. The usual concerns with the bandsaw apply. Round parts will need to be put in a vise before they are cut, and all pieces should be pushed with a sacrificial piece of wood. The smaller, flexible pieces will be cut using a handheld knife, to enhance precision. The user will need to take care to hold the knife carefully (cut away from the body, not towards it!) and to safely put it away when not cutting.

The drilling processes will be performed on a drill press in the ME shop. Since the Machinist’s Handbook does not specify cutting speeds for plastic, we will set the speed to 500RPM for all cuts. We assume that this would be safely low enough for any size bit. As usual, when drilling, we will need to put all workpieces in a vise, and clamp the vise to the drill press table to ensure that it will not move.

**6 ASSEMBLY**
The assembly of the devices will occur in the ME Machine shop. The CAD drawings in Section 4, above, show the final assembly design, and Section 5 includes details on how the components will be fitted together. The only safety concern in the assembly of the devices will be the use of the ScotchWeld glue. We will need to be mindful not to inhale its fumes, and not to glue ourselves to anything. Most of the assembly for these devices include nothing more than gluing together plastic parts. We do not foresee any dangers in the assembly process.

**7 EXPERIMENTAL/VALIDATION PLAN**
Much of the testing will occur during manufacturing, checking each part of the prototype along the way. For example, the pinch valve rod will be testing by squeezing a PVC tube to see if it will make it seal. Also, all of the components will be tested in a “dry fit” to make sure that all the parts fit together before
they are glued or attached. We anticipate that our production process will include lots of on-the-fly tests to check each component and see whether it performs the job it is supposed to do.

We do not have access to the appropriate devices to test the pressure that can be sustained with our valves. We have talked to Tom Bress, and he has said that he does not have any pressure transducers, or any other method with which we can ensure that the pressure in the syringe meets the engineering requirements that we set for our devices. The best alternative we can come up with is simple: start with the syringe closed and the attached valve closed. Pull the syringe plunger back to the full displacement, and wait for some amount of time before releasing the plunger. If the plunger returns to its fully-closed position, then we can deduce that the valve is not leaking, and that the vacuum created in the syringe will be constant. By holding the plunger back for longer amounts of time, we can ensure that the valves hold pressure for the longer lengths of time and do not suffer from slow leaks. The activation force is the specification that will be most difficult to test. We plan to take the completed prototypes to Tom Bress again, and possibly use an Instron machine to test the force required to push the buttons on both valve types. This will be a definitive test to see whether our devices meet the design specifications. We do not foresee any safety issues or concerns with any of this testing.

The true test will be whether or not our sponsor believes these devices will perform in the real world. Professor Davenport has the experience with the procedure to recognize whether our devices will work or not. And, in his words, the ultimate test will be clinical trials, if the devices make it that far into the development process. At this point, the safety concerns will be all of the concerns for normal use. Training for the doctors would be important for their safety, as well the appropriate personal protective equipment.

8 ADDITIONAL APPENDICES
The Materials Data and Safety Sheet (MSDS) for the ScotchWeld DP-8005 can be found in Appendix D.
# Appendix A: Bill of Materials

## Materials List

**FINE NEEDLE ASPIRATION**

15-Apr-09

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<th>Part #</th>
<th>Part Name</th>
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## Appendix B: FMEA Table

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<th>Material or Item</th>
<th>Functions</th>
<th>Potential Failures</th>
<th>Causes or Mechanisms of Failure Modes</th>
<th>Effects of Failure</th>
<th>Likelihood of Occurrence</th>
<th>Potential Severity</th>
<th>Controls Tests for detecting Failure</th>
<th>Risk Priority Number (RPN)</th>
<th>Actions for Failure Modes</th>
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<td>16 Gauge Galvanized Wire</td>
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<td>Manufacturing defect, tensile stress</td>
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<td>Polypropylene Rod</td>
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<td>27</td>
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<td></td>
</tr>
<tr>
<td>Polypropylene Rod</td>
<td>Plunger rod</td>
<td>Bending</td>
<td>Bending stress</td>
<td>Takes more effort to push valve</td>
<td>1</td>
<td>3</td>
<td>1</td>
<td>3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Polypropylene Luer-Loks</td>
<td>Interface between valve and syringe/needle</td>
<td>Bond delamination</td>
<td>Improper surface prep</td>
<td>Valve does not hold pressure</td>
<td>3</td>
<td>9</td>
<td>1</td>
<td>27</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Clear PVC tube</td>
<td>Flexible sealing tube</td>
<td>Tearing</td>
<td>Shear</td>
<td>Valve does not hold pressure</td>
<td>1</td>
<td>9</td>
<td>4</td>
<td>36</td>
<td>Pre-test devices before sending</td>
<td>27</td>
</tr>
<tr>
<td>16 Gauge Galvanized Wire</td>
<td>Button Lock</td>
<td>Shear, Fatigue</td>
<td>Bending</td>
<td>Button locking mechanism fails</td>
<td>2</td>
<td>6</td>
<td>2</td>
<td>24</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ScotchWeld DP-8005 Glue</td>
<td>Glue Luer-Loks to valve body</td>
<td>Bond delamination</td>
<td>Manufacturing defect, improper surface prep</td>
<td>Valve does not hold pressure</td>
<td>3</td>
<td>9</td>
<td>1</td>
<td>27</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Acrylic Sheets</td>
<td>Button Top</td>
<td>Shatter, Chip</td>
<td>Shear, Bending</td>
<td>Smaller button top</td>
<td>2</td>
<td>4</td>
<td>2</td>
<td>16</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
**APPENDIX C: DesignSafe Reports**

Gate Valve

---

**desigmsafe Report**

<table>
<thead>
<tr>
<th>Application:</th>
<th>Gate Valve</th>
</tr>
</thead>
<tbody>
<tr>
<td>Description:</td>
<td>Gate Valve</td>
</tr>
<tr>
<td>Analyst Name(s):</td>
<td>ME 450 Team 4</td>
</tr>
<tr>
<td>Company:</td>
<td>University of Michigan</td>
</tr>
<tr>
<td>Facility Location:</td>
<td></td>
</tr>
<tr>
<td>Assessment Type:</td>
<td>Detailed</td>
</tr>
</tbody>
</table>

**Limits:**

**Sources:**

Guide sentence: When doing [task], the [user] could be injured by the [hazard] due to the [failure mode].

<table>
<thead>
<tr>
<th>User / Task</th>
<th>Hazard / Failure Mode</th>
<th>Initial Assessment</th>
<th>Final Assessment</th>
<th>Status / Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Severity</td>
<td>Exposure Probability</td>
<td>Risk Level</td>
</tr>
<tr>
<td>All Users</td>
<td>mechanical : pinch point</td>
<td>Minimal</td>
<td>Remote</td>
<td>Unlikely</td>
</tr>
<tr>
<td>All Users</td>
<td>mechanical : impact could dislodge gate</td>
<td>Catastrophic</td>
<td>Remote</td>
<td>Negligible</td>
</tr>
<tr>
<td>All Users</td>
<td>ergonomics / human factors : excessive force / exertion arthritic hands could be stressed</td>
<td>Slight</td>
<td>Occasional</td>
<td>Unlikely</td>
</tr>
<tr>
<td>All Users</td>
<td>biological / health : blood borne diseases blood could leak from valve</td>
<td>Serious</td>
<td>Occasional</td>
<td>Unlikely</td>
</tr>
<tr>
<td>All Users</td>
<td>fluid / pressure : vacuum could ingest exterior fluids/particulate</td>
<td>Serious</td>
<td>Remote</td>
<td>Negligible</td>
</tr>
<tr>
<td>All Users</td>
<td>fluid / pressure : fluid leakage / ejection blood could leak from valve</td>
<td>Serious</td>
<td>Occasional</td>
<td>Unlikely</td>
</tr>
</tbody>
</table>
**Pinch Valve**

<table>
<thead>
<tr>
<th>Application:</th>
<th>Pinch Valve</th>
<th>Analyst Name(s):</th>
<th>ME 450 Team 4</th>
</tr>
</thead>
<tbody>
<tr>
<td>Description:</td>
<td></td>
<td>Company:</td>
<td>University of Michigan</td>
</tr>
<tr>
<td>Product Identifier:</td>
<td></td>
<td>Facility Location:</td>
<td></td>
</tr>
<tr>
<td>Assessment Type:</td>
<td>Detailed</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Limits:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sources:</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Guide sentence: When doing [task], the [user] could be injured by the [hazard] due to the [failure mode].

<table>
<thead>
<tr>
<th>User / Task</th>
<th>Hazard / Failure Mode</th>
<th>Initial Assessment</th>
<th>Final Assessment</th>
<th>Status / Severity</th>
<th>Risk Level</th>
<th>Exposure</th>
<th>Probability</th>
<th>Risk Reduction Methods / Comments</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>All Users</td>
<td>mechanical: pinch point</td>
<td>Minimal</td>
<td>Minimal</td>
<td>Remote</td>
<td>Unlikely</td>
<td>Low</td>
<td>small button</td>
<td></td>
<td></td>
</tr>
<tr>
<td>All Tasks</td>
<td>finger could get caught under button</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>All Users</td>
<td>mechanical: break up during</td>
<td>Catastrophic</td>
<td>Catastrophic</td>
<td>None</td>
<td>Unlikely</td>
<td>Moderate</td>
<td>different tube material</td>
<td></td>
<td></td>
</tr>
<tr>
<td>All Tasks</td>
<td>operation</td>
<td>Remote</td>
<td>Remote</td>
<td></td>
<td>Unlikely</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>All Users</td>
<td>mechanical: impact</td>
<td>Catastrophic</td>
<td>Catastrophic</td>
<td>None</td>
<td>Negligible</td>
<td>Moderate</td>
<td>standard procedures</td>
<td></td>
<td></td>
</tr>
<tr>
<td>All Tasks</td>
<td>could jar lock loose</td>
<td>Remote</td>
<td>Remote</td>
<td></td>
<td>Unlikely</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>All Users</td>
<td>ergonomics / human factors:</td>
<td>Slight</td>
<td>Slight</td>
<td></td>
<td>Remote</td>
<td>Low</td>
<td>scheduled rest periods</td>
<td></td>
<td></td>
</tr>
<tr>
<td>All Tasks</td>
<td>excessive force / exertion</td>
<td>Remote</td>
<td>Remote</td>
<td></td>
<td>Unlikely</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>hand could be too weak to squeeze button</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>All Users</td>
<td>heat / temperature: severe</td>
<td>Serious</td>
<td>Serious</td>
<td>None</td>
<td></td>
<td>Low</td>
<td>warning label(s)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>All Tasks</td>
<td>cold could make flexible tube brittle</td>
<td>None</td>
<td>None</td>
<td></td>
<td>Negligible</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>All Users</td>
<td>fluid / pressure: fluid leakage /</td>
<td>Serious</td>
<td>Serious</td>
<td>Remote</td>
<td></td>
<td>Low</td>
<td>gloves</td>
<td></td>
<td></td>
</tr>
<tr>
<td>All Tasks</td>
<td>ejection could occur if tube fails</td>
<td>Remote</td>
<td>Remote</td>
<td>Negligible</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Guide sentence: When doing [task], the [user] could be injured by the [hazard] due to the [failure mode].

<table>
<thead>
<tr>
<th>User / Task</th>
<th>Hazard / Failure Mode</th>
<th>Initial Assessment</th>
<th>Final Assessment</th>
<th>Status / Responsible</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Severity</td>
<td>Exposure</td>
<td>Probability</td>
</tr>
<tr>
<td>All Users</td>
<td>mechanical : crushing</td>
<td>Minimal</td>
<td>Occasional</td>
<td>Negligible</td>
</tr>
<tr>
<td>All Tasks</td>
<td>Crushing from syringe pulling in</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>All Users</td>
<td>mechanical : cutting / severing</td>
<td>Serious</td>
<td>Occasional</td>
<td>Unlikely</td>
</tr>
<tr>
<td>All Tasks</td>
<td>Moving sawtooth edge could cut</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>All Users</td>
<td>mechanical : pinch point</td>
<td>Slight</td>
<td>Occasional</td>
<td>Possible</td>
</tr>
<tr>
<td>All Tasks</td>
<td>Moving sawtooth edge and lock could catch a finger</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>All Users</td>
<td>mechanical : break up during operation</td>
<td>Catastrophic</td>
<td>None</td>
<td>Possible</td>
</tr>
<tr>
<td>All Tasks</td>
<td>The sawtooth could fail</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>All Users</td>
<td>mechanical : impact</td>
<td>Catastrophic</td>
<td>None</td>
<td>Negligible</td>
</tr>
<tr>
<td>All Tasks</td>
<td>Could fail under impact</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>All Users</td>
<td>ergonomics / human factors : excessive force / exertion</td>
<td>Minimal</td>
<td>Remote</td>
<td>Negligible</td>
</tr>
<tr>
<td>All Tasks</td>
<td>User could potentially not be strong enough to retract plunger</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>All Users</td>
<td>ergonomics / human factors : human errors / behaviors</td>
<td>Slight</td>
<td>Remote</td>
<td>Negligible</td>
</tr>
<tr>
<td>All Tasks</td>
<td>Improper use could lead to injury</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>All Users</td>
<td>fluid / pressure : vacuum</td>
<td>Minimal</td>
<td>Occasional</td>
<td>Negligible</td>
</tr>
<tr>
<td>All Tasks</td>
<td>Vacuum is contained in syringe barrel: could create unexpected movement</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>User / Task</td>
<td>Hazard / Failure Mode</td>
<td>Initial Assessment</td>
<td>Final Assessment</td>
<td>Status / Responsible / Reference</td>
</tr>
<tr>
<td>-------------</td>
<td>-----------------------</td>
<td>--------------------</td>
<td>------------------</td>
<td>-------------------------------</td>
</tr>
<tr>
<td>All Users</td>
<td>fluid / pressure : fluid leakage / ejection</td>
<td>Moderate/none</td>
<td>Serious/Remote/Unlikely</td>
<td>Moderate</td>
</tr>
<tr>
<td>All Tasks</td>
<td>If the syringe seal leaks, it could possibly allow blood or tissue to escape</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
**designsafe Report**

**Application:** Fine Needle Aspiration Device Assembly

**Analyst Name(s):** ME 450 Team 4

**Company:** University of Michigan

**Facility Location:**

**Limits:**

**Sources:**

Guide sentence: When doing [task], the [user] could be injured by the [hazard] due to the [failure mode].

<table>
<thead>
<tr>
<th>User / Task</th>
<th>Hazard / Failure Mode</th>
<th>Initial Assessment</th>
<th>Risk Reduction Methods /Comments</th>
<th>Final Assessment</th>
<th>Status / Responsible /Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>All Users All Tasks</td>
<td>mechanical : crushing syringe plunger</td>
<td>Slight Remote Unlikely</td>
<td>Low</td>
<td>smaller syringe</td>
<td>Minimal Remote Unlikely Low</td>
</tr>
<tr>
<td>All Users All Tasks</td>
<td>mechanical : cutting / severing plunger sawtooth and needle</td>
<td>Serious Remote Unlikely</td>
<td>Moderate</td>
<td>rounded sawtooth edge, needle guard</td>
<td>Slight Remote Unlikely Low</td>
</tr>
<tr>
<td>All Users All Tasks</td>
<td>mechanical : pinch point valve buttons and sawtooth</td>
<td>Minimal Occasional Unlikely</td>
<td>Low</td>
<td>rounded sawtooth edge, small buttons</td>
<td>Minimal Remote Negligible Low</td>
</tr>
<tr>
<td>All Users All Tasks</td>
<td>mechanical : stabbing / puncture needle</td>
<td>Serious Occasional Unlikely</td>
<td>Moderate</td>
<td>needle guard</td>
<td>Serious Remote Negligible Low</td>
</tr>
<tr>
<td>All Users All Tasks</td>
<td>mechanical : break up during operation luer-locks</td>
<td>Catastrophic Remote Unlikely</td>
<td>Moderate</td>
<td>one-piece valve</td>
<td>Catastrophic None Unlikely Moderate</td>
</tr>
<tr>
<td>All Users All Tasks</td>
<td>mechanical : impact whole device</td>
<td>Catastrophic Remote Unlikely</td>
<td>Moderate</td>
<td>standard procedures</td>
<td>Catastrophic None Negligible Low</td>
</tr>
<tr>
<td>All Users All Tasks</td>
<td>ergonomics / human factors : excessive force / exertion valve, plunger</td>
<td>Slight Occasional Unlikely</td>
<td>Moderate</td>
<td>scheduled rest periods</td>
<td>Slight Remote Unlikely Low</td>
</tr>
<tr>
<td>All Users All Tasks</td>
<td>biological / health : blood borne diseases valve, luer-locks</td>
<td>Serious Remote Unlikely</td>
<td>Moderate</td>
<td>one-piece valves, gloves</td>
<td>Serious Remote Unlikely Moderate</td>
</tr>
<tr>
<td>All Users All Tasks</td>
<td>fluid / pressure : vacuum syringe, valve</td>
<td>Slight Remote Unlikely</td>
<td>Low</td>
<td>smaller syringe</td>
<td>Minimal Remote Unlikely Low</td>
</tr>
<tr>
<td>All Users All Tasks</td>
<td>fluid / pressure : fluid leakage / ejection syringe, valve, luer-locks</td>
<td>Serious Remote Unlikely</td>
<td>Moderate</td>
<td>one-piece valves, gloves</td>
<td>Serious Remote Unlikely Moderate</td>
</tr>
</tbody>
</table>
APPENDIX D: ScotchWeld DP-8005 MSDS

Material Safety Data Sheet

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PRODUCT NAME: Scotch-Weld(TM) Structural Plastic Adhesive DP-8005
MANUFACTURER: 3M
DIVISION: Industrial Adhesives and Tapes
ADDRESS: 3M Center
St. Paul, MN 55144-1000

EMERGENCY PHONE: 1-800-364-3577 or (651) 737-6501 (24 hours)

Issue Date: 09/12/2003
Supercedes Date: 05/22/2001

Document Group: 08-8288-6

ID Number(s):

This product is a kit or a multipart product which consists of multiple, independently packaged components. An MSDS for each of these components is included. Please do not separate the component MSDSs from this cover page. The document numbers of the MSDSs for components of this product are:

08-8286-0, 08-8284-5

No revision information is available.

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SECTION 1: PRODUCT AND COMPANY IDENTIFICATION

PRODUCT NAME: 3M(TM) Scotch-Weld(TM) Structural Plastic Adhesive DP-8005 (Part A)
MANUFACTURER: 3M
DIVISION: Industrial Adhesives and Tapes Division
ADDRESS: 3M Center
St. Paul, MN 55144-1000

EMERGENCY PHONE: 1-800-364-3577 or (651) 737-6501 (24 hours)

Issue Date: 05/20/2008
Supercedes Date: 10/15/2007
Document Group: 08-8284-5

Product Use:
Specific Use: part A of two part adhesive
Intended Use: Structural adhesive

SECTION 2: INGREDIENTS

<table>
<thead>
<tr>
<th>Ingredient</th>
<th>C.A.S. No.</th>
<th>% by Wt</th>
</tr>
</thead>
<tbody>
<tr>
<td>Polyester Adipate - N.J.T.S. Reg No. 800928-5001</td>
<td>Trade Secret</td>
<td>40 - 70</td>
</tr>
<tr>
<td>Amine Borane Complex</td>
<td>223674-50-8</td>
<td>10 - 30</td>
</tr>
<tr>
<td>Polyfunctional Aziridine</td>
<td>64265-57-2</td>
<td>10 - 30</td>
</tr>
<tr>
<td>Amorphous Silica</td>
<td>67762-90-7</td>
<td>0.5 - 1.5</td>
</tr>
</tbody>
</table>

SECTION 3: HAZARDS IDENTIFICATION

3.1 EMERGENCY OVERVIEW

Specific Physical Form: Paste
Odor, Color, Grade: mild odor, white
General Physical Form: Liquid
Immediate health, physical, and environmental hazards: Combustible liquid and vapor. Closed containers exposed to heat from fire may build pressure and explode. Vapors may travel long distances along the ground or floor to an ignition source and flash back. May cause chemical eye burns. May cause allergic skin reaction. May cause severe skin irritation. May cause allergic respiratory reaction.
3.2 POTENTIAL HEALTH EFFECTS

Eye Contact:
Corrosive (Eye Burns): Signs/symptoms may include cloudy appearance of the cornea, chemical burns, severe pain, tearing, ulcerations, significantly impaired vision or complete loss of vision.

Vapors released during curing may cause eye irritation. Signs/symptoms may include redness, swelling, pain, tearing, and blurred or hazy vision.

Dust created by cutting, grinding, sanding, or machining may cause eye irritation. Signs/symptoms may include redness, swelling, pain, tearing, and blurred or hazy vision.

Skin Contact:
Severe Skin Irritation: Signs/symptoms may include localized redness, swelling, itching, dryness, cracking, blistering, and pain.

Prolonged or repeated exposure may cause:
Allergic Skin Reaction (non-photo induced): Signs/symptoms may include redness, swelling, blistering, and itching.

Inhalation:
Respiratory Tract Irritation: Signs/symptoms may include cough, sneezing, nasal discharge, headache, hoarseness, and nose and throat pain.

Dust from cutting, grinding, sanding or machining may cause irritation of the respiratory system. Signs/symptoms may include cough, sneezing, nasal discharge, headache, hoarseness, and nose and throat pain.

Prolonged or repeated exposure may cause:
Allergic Respiratory Reaction: Signs/symptoms may include difficulty breathing, wheezing, cough, and tightness of chest.

Ingestion:
Gastrointestinal Irritation: Signs/symptoms may include abdominal pain, stomach upset, nausea, vomiting and diarrhea.

SECTION 4: FIRST AID MEASURES

4.1 FIRST AID PROCEDURES

The following first aid recommendations are based on an assumption that appropriate personal and industrial hygiene practices are followed.

Eye Contact: Immediately flush eyes with large amounts of water for at least 15 minutes. Get immediate medical attention.

Skin Contact: Remove contaminated clothing and shoes. Immediately flush skin with large amounts of water. Get medical attention. Wash contaminated clothing and clean shoes before reuse.
Inhalation: Remove person to fresh air. If signs/symptoms develop, get medical attention.

If Swallowed: Do not induce vomiting unless instructed to do so by medical personnel. Give victim two glasses of water. Never give anything by mouth to an unconscious person. Get medical attention.

**SECTION 5: FIRE FIGHTING MEASURES**

5.1 FLAMMABLE PROPERTIES

<table>
<thead>
<tr>
<th>Property</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Autoignition temperature</td>
<td>No Data Available</td>
</tr>
<tr>
<td>Flash Point</td>
<td>180 °F [Test Method: Closed Cup]</td>
</tr>
<tr>
<td>Flammable Limits - LEL</td>
<td>No Data Available</td>
</tr>
<tr>
<td>Flammable Limits - UEL</td>
<td>No Data Available</td>
</tr>
<tr>
<td>OSHA Flammability Classification</td>
<td>Class IIIA Combustible Liquid</td>
</tr>
</tbody>
</table>

5.2 EXTINGUISHING MEDIA

Use fire extinguishers with class B extinguishing agents (e.g., dry chemical, carbon dioxide).

5.3 PROTECTION OF FIRE FIGHTERS

Special Fire Fighting Procedures: Water may not effectively extinguish fire; however, it should be used to keep fire-exposed containers and surfaces cool and prevent explosive rupture. Wear full protective equipment (Bunker Gear) and a self-contained breathing apparatus (SCBA).

Unusual Fire and Explosion Hazards: Combustible liquid and vapor. Closed containers exposed to heat from fire may build pressure and explode. Vapors may travel long distances along the ground or floor to an ignition source and flash back.

Note: See STABILITY AND REACTIVITY (SECTION 10) for hazardous combustion and thermal decomposition information.

**SECTION 6: ACCIDENTAL RELEASE MEASURES**

Accidental Release Measures: Observe precautions from other sections. Call 3M- HELPS line (1-800-364-3577) for more information on handling and managing the spill. Evacuate unprotected and untrained personnel from hazard area. The spill should be cleaned up by qualified personnel. Ventilate the area with fresh air. For large spill, or spills in confined spaces, provide mechanical ventilation to disperse or exhaust vapors, in accordance with good industrial hygiene practice. Warning! A motor could be an ignition source and could cause flammable gases or vapors in the spill area to burn or explode. Contain spill. For larger spills, cover drains and build dikes to prevent entry into sewer systems or bodies of water. Working from around the edges of the spill inward, cover with bentonite, vermiculite, or commercially available inorganic absorbent material. Mix in sufficient absorbent until it appears dry. Collect as much of the spilled material as possible. Clean up residue with an appropriate solvent selected by a qualified and authorized person. Ventilate the area with fresh air. Read and follow safety precautions on the solvent label and MSDS. Collect the resulting residue containing solution. Place in a closed container approved for transportation by appropriate authorities. Dispose of collected material as soon as possible.

In the event of a release of this material, the user should determine if the release qualifies as reportable according to local, state, and federal regulations.

**SECTION 7: HANDLING AND STORAGE**

7.1 HANDLING
Do not eat, drink or smoke when using this product. Wash exposed areas thoroughly with soap and water. Keep away from heat, sparks, open flame, pilot lights and other sources of ignition. Avoid skin contact. Avoid breathing of vapors. Avoid eye contact with vapors, mists, or spray. Keep out of the reach of children. Keep container closed when not in use. Avoid breathing of dust created by cutting, sanding, grinding or machining. For industrial or professional use only. Avoid contact with oxidizing agents. Use general dilution ventilation and/or local exhaust ventilation to control airborne exposures to below Occupational Exposure Limits. If ventilation is not adequate, use respiratory protection equipment.

7.2 STORAGE

SECTION 8: EXPOSURE CONTROLS/PERSONAL PROTECTION

8.1 ENGINEERING CONTROLS
Provide appropriate local exhaust for cutting, grinding, sanding or machining. Use general dilution ventilation and/or local exhaust ventilation to control airborne exposures to below Occupational Exposure Limits and/or control dust, fume, or airborne particles. If ventilation is not adequate, use respiratory protection equipment.

8.2 PERSONAL PROTECTIVE EQUIPMENT (PPE)

8.2.1 Eye/Face Protection
Avoid eye contact.
The following eye protection(s) are recommended: Safety Glasses with side shields, Indirect Vented Goggles.

8.2.2 Skin Protection
Avoid skin contact.
Select and use gloves and/or protective clothing to prevent skin contact based on the results of an exposure assessment. Consult with your glove and/or protective clothing manufacturer for selection of appropriate compatible materials.
Gloves made from the following material(s) are recommended: Butyl Rubber, Nitrile Rubber, Polyethylene, Polyvinyl Alcohol (PVA).

8.2.3 Respiratory Protection
Avoid breathing of vapors. Avoid breathing of dust created by cutting, sanding, grinding or machining.
Select one of the following NIOSH approved respirators based on airborne concentration of contaminants and in accordance with OSHA regulations: Half facepiece or full face air-purifying respirator with formaldehyde cartridges and N95 particulate prefilters, Half facepiece or fullface air-purifying respirator with formaldehyde cartridges and P100 particulate preilters, Half facepiece or fullface air-purifying respirator with formaldehyde cartridges and P95 particulate preilters. Consult the current 3M Respiratory Selection Guide for additional information or call 1-800-243-4630 for 3M technical assistance.

8.2.4 Prevention of Swallowing
Do not eat, drink or smoke when using this product. Wash exposed areas thoroughly with soap and water.

8.3 EXPOSURE GUIDELINES

<table>
<thead>
<tr>
<th>Ingredient</th>
<th>Authority</th>
<th>Type</th>
<th>Limit</th>
<th>Additional Information</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amorphous Silica</td>
<td>CMRG</td>
<td>CEIL</td>
<td>5 mg/m3</td>
<td></td>
</tr>
</tbody>
</table>

SOURCE OF EXPOSURE LIMIT DATA:
ACGIH: American Conference of Governmental Industrial Hygienists
CMRG: Chemical Manufacturer Recommended Guideline
OSHA: Occupational Safety and Health Administration
AIHA: American Industrial Hygiene Association Workplace Environmental Exposure Level (WEEL)

**SECTION 9: PHYSICAL AND CHEMICAL PROPERTIES**

<table>
<thead>
<tr>
<th>Property</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Specific Physical Form</td>
<td>Paste</td>
</tr>
<tr>
<td>Odor, Color, Grade</td>
<td>mild odor, white</td>
</tr>
<tr>
<td>General Physical Form</td>
<td>Liquid</td>
</tr>
<tr>
<td>Autoignition temperature</td>
<td>No Data Available</td>
</tr>
<tr>
<td>Flash Point</td>
<td></td>
</tr>
<tr>
<td>Flammable Limits - LEL</td>
<td>No Data Available</td>
</tr>
<tr>
<td>Flammable Limits - UEL</td>
<td>No Data Available</td>
</tr>
<tr>
<td>Boiling point</td>
<td>&gt;=95 ºF</td>
</tr>
<tr>
<td>Vapor Density</td>
<td>No Data Available</td>
</tr>
<tr>
<td>Vapor Pressure</td>
<td>&lt;=0.1 mmHg</td>
</tr>
<tr>
<td>Specific Gravity</td>
<td>1.063</td>
</tr>
<tr>
<td>Specific Gravity</td>
<td>0.991 [Details: when mixed 10 parts B to 1 part A]</td>
</tr>
<tr>
<td>pH</td>
<td>Not Applicable</td>
</tr>
<tr>
<td>Melting point</td>
<td>Not Applicable</td>
</tr>
<tr>
<td>Solubility in Water</td>
<td>Slight (less than 10%)</td>
</tr>
<tr>
<td>Evaporation rate</td>
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</tr>
<tr>
<td>Volatile Organic Compounds</td>
<td>6.15 % weight [Test Method: tested per EPA method 24A]</td>
</tr>
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<td>Volatile Organic Compounds</td>
<td>4.81 [Test Method: tested per EPA method 24A] [Details: when mixed 10 parts B to 1 part A]</td>
</tr>
<tr>
<td>VOC Less H2O &amp; Exempt Solvents</td>
<td>65 g/l [Test Method: tested per EPA method 24A]</td>
</tr>
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<td>VOC Less H2O &amp; Exempt Solvents</td>
<td>48 g/l [Test Method: tested per EPA method 24A] [Details: when mixed 10 parts B to 1 part A]</td>
</tr>
<tr>
<td>Viscosity</td>
<td>49000 centipoise [@ 73.4 ºF]</td>
</tr>
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</table>

**SECTION 10: STABILITY AND REACTIVITY**

Stability: Stable.

Materials and Conditions to Avoid: Strong acids; Heat; Sparks and/or flames; Strong oxidizing agents

Hazardous Polymerization: Hazardous polymerization will not occur.

**Hazardous Decomposition or By-Products**

<table>
<thead>
<tr>
<th>Substance</th>
<th>Condition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aldehydes</td>
<td>During Combustion</td>
</tr>
<tr>
<td>Carbon monoxide</td>
<td>During Combustion</td>
</tr>
<tr>
<td>Carbon dioxide</td>
<td>During Combustion</td>
</tr>
<tr>
<td>Irritant Vapors or Gases</td>
<td>During Combustion</td>
</tr>
<tr>
<td>Oxides of Nitrogen</td>
<td>During Combustion</td>
</tr>
</tbody>
</table>

**SECTION 11: TOXICOLOGICAL INFORMATION**
Please contact the address listed on the first page of the MSDS for Toxicological Information on this material and/or its components.

**SECTION 12: ECOLOGICAL INFORMATION**

**ECOTOXICOLOGICAL INFORMATION**
Not determined.

**CHEMICAL FATE INFORMATION**
Not determined.

**SECTION 13: DISPOSAL CONSIDERATIONS**

**Waste Disposal Method:** Cure (harden, set, or react) the product according to product instructions.
Dispose of completely cured (or polymerized) wastes in a sanitary landfill.
As a disposal alternative, incinerate uncured product in an industrial or commercial incinerator.

**EPA Hazardous Waste Number (RCRA):** Not regulated

Since regulations vary, consult applicable regulations or authorities before disposal.

**SECTION 14: TRANSPORT INFORMATION**

**ID Number(s):**
62-2886-7530-9, 62-2886-8530-8

Not regulated per U.S. DOT, IATA or IMO.

These transportation classifications are provided as a customer service. As the shipper YOU remain responsible for complying with all applicable laws and regulations, including proper transportation classification and packaging. 3M’s transportation classifications are based on product formulation, packaging, 3M policies and 3M’s understanding of applicable current regulations. 3M does not guarantee the accuracy of this classification information. This information applies only to transportation classification and **not the packaging, labeling, or marking requirements.** The original 3M package is certified for U.S. ground shipment only. If you are shipping by air or ocean, the package may not meet applicable regulatory requirements.

**SECTION 15: REGULATORY INFORMATION**

**US FEDERAL REGULATIONS**
Contact 3M for more information.

**311/312 Hazard Categories:**
Fire Hazard - Yes Pressure Hazard - No Reactivity Hazard - No Immediate Hazard - Yes Delayed Hazard - No
STATE REGULATIONS
Contact 3M for more information.

CHEMICAL INVENTORIES
One or more of the components in this material is not listed on the TSCA inventory, but is approved for specific commercial use(s) under a US EPA low volume exemption (up to 10,000 kg/yr). Research and development production quantities are not included in the 10,000 kg/yr limit.

Contact 3M for more information.

INTERNATIONAL REGULATIONS
Contact 3M for more information.

This MSDS has been prepared to meet the U.S. OSHA Hazard Communication Standard, 29 CFR 1910.1200.

SECTION 16: OTHER INFORMATION

NFPA Hazard Classification

Health: 3  Flammability: 2  Reactivity: 0  Special Hazards: None

National Fire Protection Association (NFPA) hazard ratings are designed for use by emergency response personnel to address the hazards that are presented by short-term, acute exposure to a material under conditions of fire, spill, or similar emergencies. Hazard ratings are primarily based on the inherent physical and toxic properties of the material but also include the toxic properties of combustion or decomposition products that are known to be generated in significant quantities.

Revision Changes:
Copyright was modified.
Section 9: Property description for optional properties was modified.

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3M MSDSs are available at www.3M.com
Material Safety Data Sheet

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SECTION 1: PRODUCT AND COMPANY IDENTIFICATION

PRODUCT NAME: Scotch-Weld(TM) Structural Plastic Adhesive DP-8005 (Part B)
MANUFACTURER: 3M
DIVISION: Industrial Adhesives and Tapes Division
ADDRESS: 3M Center
St. Paul, MN  55144-1000

EMERGENCY PHONE: 1-800-364-3577 or (651) 737-6501 (24 hours)

Issue Date: 05/20/2008
Supercedes Date: 11/09/2007
Document Group: 08-8286-0

Product Use:
Specific Use: part B of 2 part adhesive
Intended Use: Industrial use

SECTION 2: INGREDIENTS

<table>
<thead>
<tr>
<th>Ingredient</th>
<th>C.A.S. No.</th>
<th>% by Wt</th>
</tr>
</thead>
<tbody>
<tr>
<td>Methacrylate</td>
<td>2455-24-5</td>
<td>40 - 60</td>
</tr>
<tr>
<td>2-Ethylhexyl Methacrylate</td>
<td>688-84-6</td>
<td>10 - 30</td>
</tr>
<tr>
<td>Acrylonitrile-Butadiene-Styrene Terpolymer</td>
<td>9003-56-9</td>
<td>10 - 30</td>
</tr>
<tr>
<td>Glass Spheres</td>
<td>68131-74-8</td>
<td>1 - 10</td>
</tr>
<tr>
<td>Impact Modifier</td>
<td>20882-04-6</td>
<td>1 - 3</td>
</tr>
</tbody>
</table>

SECTION 3: HAZARDS IDENTIFICATION

3.1 EMERGENCY OVERVIEW

Specific Physical Form: Paste
Odor, Color, Grade: Translucent, mild acrylic odor
General Physical Form: Liquid
Immediate health, physical, and environmental hazards:

3.2 POTENTIAL HEALTH EFFECTS
Eye Contact:
Moderate Eye Irritation: Signs/symptoms may include redness, swelling, pain, tearing, and blurred or hazy vision.

Vapors released during curing may cause eye irritation. Signs/symptoms may include redness, swelling, pain, tearing, and blurred or hazy vision.

Dust created by cutting, grinding, sanding, or machining may cause eye irritation. Signs/symptoms may include redness, swelling, pain, tearing, and blurred or hazy vision.

Skin Contact:
Moderate Skin Irritation: Signs/symptoms may include localized redness, swelling, itching, and dryness.

Inhalation:
Respiratory Tract Irritation: Signs/symptoms may include cough, sneezing, nasal discharge, headache, hoarseness, and nose and throat pain.

Dust from cutting, grinding, sanding or machining may cause irritation of the respiratory system. Signs/symptoms may include cough, sneezing, nasal discharge, headache, hoarseness, and nose and throat pain.

Ingestion:
Gastrointestinal Irritation: Signs/symptoms may include abdominal pain, stomach upset, nausea, vomiting and diarrhea.

SECTION 4: FIRST AID MEASURES

4.1 FIRST AID PROCEDURES

The following first aid recommendations are based on an assumption that appropriate personal and industrial hygiene practices are followed.

Eye Contact:  Flush eyes with large amounts of water. If signs/symptoms persist, get medical attention.

Skin Contact:  Remove contaminated clothing and shoes. Immediately flush skin with large amounts of water. Get medical attention. Wash contaminated clothing and clean shoes before reuse.

Inhalation:  Remove person to fresh air. If signs/symptoms develop, get medical attention.

If Swallowed: Do not induce vomiting unless instructed to do so by medical personnel. Give victim two glasses of water. Never give anything by mouth to an unconscious person. Get medical attention.

SECTION 5: FIRE FIGHTING MEASURES

5.1 FLAMMABLE PROPERTIES

<table>
<thead>
<tr>
<th>Property</th>
<th>Details</th>
</tr>
</thead>
<tbody>
<tr>
<td>Autoignition temperature</td>
<td>No Data Available</td>
</tr>
<tr>
<td>Flash Point</td>
<td>218 °F [Test Method: SETAFLASH] [Details: SPECIFIC METHOD: ASTM D-3278-96]</td>
</tr>
<tr>
<td>Flammable Limits - LEL</td>
<td>No Data Available</td>
</tr>
</tbody>
</table>
Flammable Limits - UEL: No Data Available
OSHA Flammability Classification: Not Applicable

5.2 EXTINGUISHING MEDIA
Ordinary combustible material. Use fire extinguishers with class A extinguishing agents (e.g., water, foam).

5.3 PROTECTION OF FIRE FIGHTERS
Special Fire Fighting Procedures: Wear full protective equipment (Bunker Gear) and a self-contained breathing apparatus (SCBA).

Unusual Fire and Explosion Hazards: Not applicable.

Note: See STABILITY AND REACTIVITY (SECTION 10) for hazardous combustion and thermal decomposition information.

SECTION 6: ACCIDENTAL RELEASE MEASURES

Accidental Release Measures: Observe precautions from other sections. Call 3M-HELPS line (1-800-364-3577) for more information on handling and managing the spill. Evacuate unprotected and untrained personnel from hazard area. The spill should be cleaned up by qualified personnel. Ventilate the area with fresh air. For large spill, or spills in confined spaces, provide mechanical ventilation to disperse or exhaust vapors, in accordance with good industrial hygiene practice. Warning! A motor could be an ignition source and could cause flammable gases or vapors in the spill area to burn or explode. Contain spill. For larger spills, cover drains and build dikes to prevent entry into sewer systems or bodies of water. Working from around the edges of the spill inward, cover with bentonite, vermiculite, or commercially available inorganic absorbent material. Mix in sufficient absorbent until it appears dry. Collect as much of the spilled material as possible. Clean up residue with an appropriate solvent selected by a qualified and authorized person. Ventilate the area with fresh air. Read and follow safety precautions on the solvent label and MSDS. Collect the resulting residue containing solution. Place in a closed container approved for transportation by appropriate authorities. Dispose of collected material as soon as possible. Cloth or paper contaminated with adhesive should be disposed of in a metal container, covered with water and container sealed.

In the event of a release of this material, the user should determine if the release qualifies as reportable according to local, state, and federal regulations.

SECTION 7: HANDLING AND STORAGE

7.1 HANDLING
Do not eat, drink or smoke when using this product. Wash exposed areas thoroughly with soap and water. Avoid breathing of vapors, mists or spray. Avoid skin contact. Avoid eye contact with vapors, mists, or spray. Keep out of the reach of children. Keep container closed when not in use. Avoid breathing of dust created by cutting, sanding, grinding or machining. For industrial or professional use only.

7.2 STORAGE
Store away from acids. Store away from heat. Store out of direct sunlight.

SECTION 8: EXPOSURE CONTROLS/PERSOINAL PROTECTION

8.1 ENGINEERING CONTROLS
Provide appropriate local exhaust for cutting, grinding, sanding or machining. Use general dilution ventilation and/or local exhaust ventilation to control airborne exposures to below Occupational Exposure Limits and/or control dust, fume, or airborne particles. If
ventilation is not adequate, use respiratory protection equipment.

8.2 PERSONAL PROTECTIVE EQUIPMENT (PPE)

8.2.1 Eye/Face Protection
Avoid eye contact with vapors, mists, or spray.
The following eye protection(s) are recommended: Safety Glasses with side shields, Indirect Vented Goggles.

8.2.2 Skin Protection
Avoid skin contact.
Select and use gloves and/or protective clothing to prevent skin contact based on the results of an exposure assessment. Consult with your glove and/or protective clothing manufacturer for selection of appropriate compatible materials.
Gloves made from the following material(s) are recommended: Butyl Rubber, Nitrile Rubber, Polyethylene, Polyvinyl Alcohol (PVA).

8.2.3 Respiratory Protection
Avoid breathing of vapors, mists or spray. Avoid breathing of vapors created during cure cycle. Avoid breathing of dust created by cutting, sanding, grinding or machining.
Select one of the following NIOSH approved respirators based on airborne concentration of contaminants and in accordance with OSHA regulations: Half facepiece or fullface air-purifying respirator with formaldehyde cartridges and N95 particulate prefilters, Half facepiece or fullface air-purifying respirator with formaldehyde cartridges and P100 particulate prefilters, Half facepiece or fullface air-purifying respirator with formaldehyde cartridges and P95 particulate prefilters. Consult the current 3M Respiratory Selection Guide for additional information or call 1-800-243-4630 for 3M technical assistance.

8.2.4 Prevention of Swallowing
Do not eat, drink or smoke when using this product. Wash exposed areas thoroughly with soap and water.

8.3 EXPOSURE GUIDELINES
None Established

SECTION 9: PHYSICAL AND CHEMICAL PROPERTIES

<table>
<thead>
<tr>
<th>Property</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Specific Physical Form:</td>
<td>Paste</td>
</tr>
<tr>
<td>Odor, Color, Grade:</td>
<td>Translucent, mild acrylic odor</td>
</tr>
<tr>
<td>General Physical Form:</td>
<td>Liquid</td>
</tr>
<tr>
<td>Autoignition temperature</td>
<td>No Data Available</td>
</tr>
<tr>
<td>Flash Point</td>
<td>218 °F [Test Method: SETAFLASH] [Details: SPECIFIC METHOD: ASTM D-3278-96]</td>
</tr>
<tr>
<td>Flammable Limits - LEL</td>
<td>No Data Available</td>
</tr>
<tr>
<td>Flammable Limits - UEL</td>
<td>No Data Available</td>
</tr>
<tr>
<td>Boiling point</td>
<td>&gt;=95 °F</td>
</tr>
<tr>
<td>Vapor Density</td>
<td>No Data Available</td>
</tr>
<tr>
<td>Vapor Pressure</td>
<td>&lt;=0.1 mmHg [@ 20 °C]</td>
</tr>
<tr>
<td>Specific Gravity</td>
<td>0.984 [Ref Std: WATER=1]</td>
</tr>
<tr>
<td>pH</td>
<td>Not Applicable</td>
</tr>
<tr>
<td>Melting point</td>
<td>Not Applicable</td>
</tr>
<tr>
<td>Solubility in Water</td>
<td>Slight (less than 10%)</td>
</tr>
<tr>
<td>Evaporation rate</td>
<td>No Data Available</td>
</tr>
</tbody>
</table>
Hazardous Air Pollutants 0 % weight
Volatile Organic Compounds 39.86 % weight [Test Method: tested per EPA method 24A]
Volatile Organic Compounds 4.81 % weight [Test Method: tested per EPA method 24A] [Details: when mixed 10 parts B to 1 part A]
VOC Less H2O & Exempt Solvents 392 g/l [Test Method: tested per EPA method 24A]
VOC Less H2O & Exempt Solvents 48 g/l [Test Method: tested per EPA method 24A] [Details: when mixed 10 parts B to 1 part A]
Viscosity 25000 centipoise

SECTION 10: STABILITY AND REACTIVITY

Stability: Stable.

Materials and Conditions to Avoid: Strong acids; Heat

Hazardous Polymerization: Hazardous polymerization will not occur.

Hazardous Decomposition or By-Products

<table>
<thead>
<tr>
<th>Substance</th>
<th>Condition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aldehydes</td>
<td>During Combustion</td>
</tr>
<tr>
<td>Carbon monoxide</td>
<td>During Combustion</td>
</tr>
<tr>
<td>Carbon dioxide</td>
<td>During Combustion</td>
</tr>
<tr>
<td>Irritant Vapors or Gases</td>
<td>During Combustion</td>
</tr>
<tr>
<td>Oxides of Nitrogen</td>
<td>During Combustion</td>
</tr>
</tbody>
</table>

SECTION 11: TOXICOLOGICAL INFORMATION

Please contact the address listed on the first page of the MSDS for Toxicological Information on this material and/or its components.

SECTION 12: ECOLOGICAL INFORMATION

ECOTOXICOLOGICAL INFORMATION

Not determined.

CHEMICAL FATE INFORMATION

Not determined.

SECTION 13: DISPOSAL CONSIDERATIONS

Waste Disposal Method: Cure (harden, set, or react) the product according to product instructions.
Dispose of completely cured (or polymerized) wastes in a sanitary landfill.
As a disposal alternative, incinerate uncured product in an industrial or commercial incinerator.
EPA Hazardous Waste Number (RCRA): Not regulated

Since regulations vary, consult applicable regulations or authorities before disposal.

SECTION 14: TRANSPORT INFORMATION

ID Number(s):

Not regulated per U.S. DOT, IATA or IMO.

These transportation classifications are provided as a customer service. As the shipper YOU remain responsible for complying with all applicable laws and regulations, including proper transportation classification and packaging. 3M’s transportation classifications are based on product formulation, packaging, 3M policies and 3M’s understanding of applicable current regulations. 3M does not guarantee the accuracy of this classification information. This information applies only to transportation classification and not the packaging, labeling, or marking requirements. The original 3M package is certified for U.S. ground shipment only. If you are shipping by air or ocean, the package may not meet applicable regulatory requirements.

SECTION 15: REGULATORY INFORMATION

US FEDERAL REGULATIONS
Contact 3M for more information.

311/312 Hazard Categories:
Fire Hazard - Yes  Pressure Hazard - No  Reactivity Hazard - No  Immediate Hazard - Yes  Delayed Hazard - No

Section 313 Toxic Chemicals subject to the reporting requirements of that section and 40 CFR part 372 (EPCRA):

<table>
<thead>
<tr>
<th>Ingredient</th>
<th>C.A.S. No</th>
<th>% by Wt</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glass Spheres (VANADIUM COMPOUNDS)</td>
<td>68131-74-8</td>
<td>1 - 10</td>
</tr>
</tbody>
</table>

STATE REGULATIONS
Contact 3M for more information.

CHEMICAL INVENTORIES
The components of this product are in compliance with the chemical notification requirements of TSCA.

Contact 3M for more information.
INTERNATIONAL REGULATIONS
Contact 3M for more information.

This MSDS has been prepared to meet the U.S. OSHA Hazard Communication Standard, 29 CFR 1910.1200.

SECTION 16: OTHER INFORMATION

NFPA Hazard Classification
Health: 2  Flammability: 1  Reactivity: 0  Special Hazards: None

National Fire Protection Association (NFPA) hazard ratings are designed for use by emergency response personnel to address the hazards that are presented by short-term, acute exposure to a material under conditions of fire, spill, or similar emergencies. Hazard ratings are primarily based on the inherent physical and toxic properties of the material but also include the toxic properties of combustion or decomposition products that are known to be generated in significant quantities.

HMIS Hazard Classification
Health: 2  Flammability: 1  Reactivity: 0  Protection: B

Hazardous Material Identification System (HMIS(r)) hazard ratings are designed to inform employees of chemical hazards in the workplace. These ratings are based on the inherent properties of the material under expected conditions of normal use and are not intended for use in emergency situations. HMIS(r) ratings are to be used with a fully implemented HMIS(r) program. HMIS(r) is a registered mark of the National Paint and Coatings Association (NPCA).

Revision Changes:
Copyright was modified.
Section 9: Property description for optional properties was modified.
Section 2: Ingredient table was modified.

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