Formal Evaluation of the Madigan Endoscopic Sinus Surgery Simulator

submitted to

Simulation and Training Systems Lockheed Martin Tactical Defense Systems

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Abstract

[executive summary here]

Introduction

[overview of simulator evaluation literature, why we need to do this]

Simulator Overview

The Madigan Endoscopic Sinus Surgery (ESS) Simulator uses virtual reality technology, a force feedback (haptic) display, and 3D computer-based anatomy models as a tool to teach a variety of skills needed to perform such surgery.

The heart of the system is a 3D model of the human nasal sinus anatomy derived from the National Library of Medicine's Visible Human Database. Researchers at Ohio Supercomputer Center (OSC) took the photographic cryosections of the male dataset, created segmented surface models of the sinus anatomy and added stochastically generated surface texture to create the model.

This model can be rendered in real time (30 frames per second) on a Silicon Graphics Onyx System and viewed at NTSC resolution on a standard video monitor, thus simulating the view that a surgeon would see of a video endoscopic display.

To interact with the model and perform the simulated surgery, students in training use a pair of 6 degree of freedom input devices developed by Immersion Corporation. One such device represents the video endoscope and the other represents the surgical instrument, such as an injection needle or a forceps.

As the student manipulates the input devices, the simulator tracks the position and orientation of the devices, updates the

positions and orientations of the virtual endoscope and virtual instrument, manipulates the virtual anatomical model accordingly, and displays the resulting virtual endoscopic view on the monitor. In addition, the system tracks the opening of the forceps handle of the instrument input device. All together, the system measures 13 degrees of freedom of the student's input.

The physical input devices are designed to resemble the feel of an endoscope and forceps and are assembled with a latex replica of a human head. The endoscope input device resides outside of the head, while the instrument input device is inserted into the nostril of the latex head and attached to the position tracker inside the head. In addition to displaying the simulated endoscopic view, the system computes the forces that the sinus tissue would apply to the tip of the instrument during surgery and applies the computed force to the instrument input device, via mechanical coupling within the mannequin head.

The proctor's console provides an interface to the run-time system parameters, along with optional radiographic views of the current anatomy and optional performance feedback summaries for the trainees.

Training Aids

Optional 3D graphical overlays are superimposed on the endoscopic view to provide performance aids for the student. These overlays include a path of circular hoops representing the desired endoscope trajectory, bullseye targets representing the desired injection sites, and text labels identifying anatomical feature landmarks.

The system provides voice audio feedback representing the current status of the surgical procedure, as well as a simulated heartbeat which responds to certain user actions.

Training Tasks

Students using the system are instructed to perform a simulated surgical procedure consisting of three subtasks: Navigation, Injection and Dissection. In addition to archiving the frame-byframe position of the devices, the system measures the time required to perform these tasks, as well as the accuracy with which they are performed, to generate an evaluation of the student's overall performance.

Three versions of the procedure were developed corresponding to three skill levels: Novice, Intermediate, and Advanced. The

primary differences among these procedures are the type of geometric model used and the presence of training aids.

The Novice procedure uses a simplified abstract model of the anatomy consisting of several spheres inside an open box. The Intermediate and Advanced procedures use the more complex surface sinus model generated by OSC. The Novice and Intermediate procedures use the training aids described above, whereas the Advanced procedure is performed without benefit of these aids (to more accurately simulate the target procedure).

Evaluation Overview

Evaluation efforts for this project fall into two general categories of activity:

- "formative" evaluation, which attempts to provide design specification input to the development team during the development process, and
- "summative" evaluation, which assesses the success of that effort by formally analyzing the effectiveness of the system.

Each of these activities is discussed in detail below.

Formative Evaluation

Throughout the development of the ESS simulator, the HITL team worked in close collaboration with Dr. Charles Edmond, staff otolaryngologist at Madigan Army Medical Center and principle investigator for the project, to perform ongoing formative evaluation and make design recommendations to the development teams at Lockheed-Martin, the Ohio Supercomputer Center, and Immersion Corporation.

The Methods of Approach for the formative evaluation phase included the following:

- 1) endoscopic video analysis to determine simulator performance requirements
- 2) geometric complexity requirements analysis
- 3) prototype anatomical modeling
- 4) development of spatial awareness aids, interface features and rendering approaches
- 5) development of a prototype simulator with an integrated expert system assistant
- 6) development of a surgical training curriculum to be embedded in simulator

7) survey of medical experts to determine feature and curriculum priorities.

Dr. Edmond provided the core of domain expertise in sinus surgery. After a few weeks of study, the rest of the HITL team had gained a basic familiarity with sinus anatomy and surgical issues. During this time Dr. Edmond gained familiarity with the computer graphics tools available at HITL for use in prototyping simulator design and computer-assisted surgery applications.

Design Requirements Analysis

Frame Rate Requirements

Method: Representative sequences of live videoendoscopic sinus surgery were digitized and manipulated. These sequences were selected to include surgical interaction with tissue as well as to highlight early operative and late operative anatomy. Image degradation due to the scanning process itself was minimal.

We then re-recorded the scanned video at the following frame rates: 30 frames per second, 15 fps, 10 fps, 5 fps, 2 fps, and 1 fps. These recordings were then reviewed by expert surgeons to assess the minimum frame rates required for the sinus surgery simulator.

Results: To the extent possible 30 fps should be maintained. For surgical dissection a minimum of 15 fps (preferably 30 fps) should be maintained; temporary slow downs to 10 fps during deformation and dissection may be tolerable for certain instruments and maneuvers (e.g., side-biter, but not sickle knife). Frame rates lower than the 10fps minimum may cause disorientation in navigation.

Geometric Complexity Requirements

We applied the following guiding principles in assessing the geometric complexity requirements:

- Visual fidelity requirements are task driven.
- The task and sub-task relevant to each individual varies based on level of training and experience.
- Three categories of expertise were defined: novice, intermediate and expert. Tasks and sub-tasks were delineated for each category, and fidelity requirements proposed.
- Accurate anatomic representation (visual realism) and 3-

dimensional spatial awareness are two of the most critical aspects for effective cognitive development. This was felt to be important across all levels of training and experience for endoscopic sinus surgery

• The development of psycho-motor skills necessary for successful endoscopic sinus surgery may require less visual realism and more 3-D spatial awareness.

- Low level visual realism might be less distracting to the novice, and therefore advantageous for both developing and assessing a user's psycho-motor skills.
- Varying degrees of visual fidelity are not only helpful, but necessary to develop the cognitive and psycho-motor skills for sinus surgery.

Methods: To help assess the geometric complexity requirements for the anatomical models we generated a prototype anatomical model using the following technique:

A 3D triangular mesh was generated by texture mapping an actual video endoscopic image onto a flat mesh surface and vertically displacing each vertex of the mesh in proportion to the brightness of texture at that vertex. We then varied the resolution of the mesh and evaluated the quality of the resulting images. This model is a fairly rough approximation, since video brightness does not precisely correspond to geometric distance from the endoscope. Specular highlights in the texture create spike artifacts in model. These were eliminated by painting over the highlights in the texture.

Geometric complexity requirements were determined by visual inspection of the resulting simulation mock-ups by the project domain experts. Evaluation criteria included:

- Ability to identify features
- Subjective evaluation of photo-realism
- Potential for simulator miscues

Results: We concluded that given the performance of the rendering system, frame rate was a critical performance requirement. When the initial versions of the anatomy model were delivered from OSC, the frame rate did not meet our specification. The geometric complexity of the model was then reduced until the specified frame rate was achieved. The geometric complexity requirement was essentially a function of the frame rate requirement and the rendering performance of the hardware and software engines.

The anatomical model developed for the ESS simulator adequately served its function. We noted considerable improvements in quality and efficiency of the model over the duration of our evaluation. Upon each release of a new model, our team would evaluate the model for rendering efficiency and fidelity. Dr. Edmond would suggest modifications to the model and these would be executed in subsequent releases. The addition of textural cues greatly enhanced the apparent geometric detail without sacrificing rendering speed.

Prototype Anatomical Modeling

Prototype anatomical virtual models were developed by Edmond and Oppenheimer using the Alias modeling package. The purpose of these models was to provide guidelines in the design of the patient specific data-driven models produced by OSC, as well as to determine the necessary rendering and interface features for use in the simulator.

Methods: The geometry of these prototypes was based on cadaveral section photographs. In one case, sections were scanned into the computer, and mapped onto parallel image planes. These image planes were then used as templates for drawing surface contours on the planes. These contours were then lofted into surfaces. Additional contour curves were added and edited as needed, based on Dr. Edmonds observations.

In other cases, freeform surface contours were drawn orthogonal to the scanned image plane. The scanned image was then projection texture mapped onto the resulting lofted surface. Although less geometrically accurate, this technique enabled photographic texture maps to be used in the final evaluation model.

Results: These prototype models provided the basis for experimenting with navigation aids, interface features and rendering approaches as described in the next section.

Interface Features and Rendering Approaches

The following design experiments using the prototype anatomical models resulted in recommendations and demonstrations of candidate features to be included in the development of the ESS Simulator:

- Use of texture
- Use of transparent or wireframe surfaces to reveal obscured anatomy
- Use of orientation cues to assist in navigation

- Use of tubular paths of hoops as a navigation aid
- Use of targets as an injection aid
- Use of cross hairs overlaid on instruments to represent orientation, and to assess distance from anatomy
- Use of patient face model
- Anatomical segmentation, and interactive selective segment display
- Displaying endoscope position on CT scans as a navigation aid

• Integration of expert system training aids

Texture mapping

The use of texture mapping offers several benefits. Texture maps can represent detail that would be too expensive to model as polygonal geometry. Photographic texture maps add to the realism of the model.

Texture also provides additional depth cues. If two parallel surfaces overlap from a given perspective view, than they may have the same shading coefficients in the lighting calculation. Without texture maps, the two surfaces would be rendered with the same pixel values and therefore the boundary between them would be indistinguishable. By placing texture maps on the surfaces, the discontinuity between the surfaces can be detected more easily, as a consequence of texture discontinuity.

In conditions of extreme ambient lighting, even non-parallel surfaces may have similar shading. Texture mapping can reveal subtle differences in orientation and distance from observer. In "real life" these shading ambiguities are also a noticible if the surfaces are of uniform texture. Real life objects with even subtle surface textural features are more readily discerned that smooth textureless objects.

Methods: Several texture mapping methods were applied to the models. In some cases an attempt was made to align the texture with the underlying model geometry; such textures included anatomical features. In other cases, the textures were closeup details of tissue type, such as mucosa, and did not include anatomical structures.

Results: On the basis of our experiments, we recommended the use of texture mapping in the target simulator. Since the polygonal resolution of the final models generated from the Visible Human database is as fine as the possible textures derived from the same data, detail texture had to be added algorithmically. Such texture, although not photo-realistic, does add to the realism of the simulation as well as improve the

Transparent and wireframe surfaces

By rendering outer surfaces transparently or in wireframe, the system can display otherwise obscured portions of the anatomy. This allows students to view anatomical landmarks, prior to revealing them through dissection. These landmarks may include anatomy that should not be dissected (such as the lamina, optic nerve, or skullbase). Revealing them may enhance spatial awareness training, thereby preventing a severe surgical error. This feature is not currently implemented in the target simulator.

Orientation cues

By displaying a world-stabilized orthogonal grid or crosshairs, the user can determine the orientation of the endoscope with respect to the patient. This feature is not currently implemented in the target simulator.

Tubular path of hoops

By displaying a tubular surface whose axis is the desired trajectory of the endoscope, the system can provide a navigation tool to the user during surgical simulation. We experimented with two different rendering modes for the tube: (A) a partially transparent surface with alpha-mapped latitudinal rings, and (B) wireframe latitudinal hoops oriented orthogonal to the axial path. The target simulator uses the latter rendering mode (hoops).

Injection targets

Bullseye targets of concentric rings, rendered as flat shaded polygons can be placed on the anatomy at points of injection. These targets provide cues in the novice and intermediate modes of the simulator.

Crosshairs attached to the instrument

By attaching polygonal crosshairs to the tip of the virtual instruments, the system visually represents the relative orientation of the instrument with respect to the endoscope. If a grid texture is mapped onto the crosshairs, the user can assess the distance between the instrument and the anatomy by counting the grid lines between the instrument tip and the intersection between the crosshair and the tissue surface. Grid marks deeper than the tissue surface are obscured and will not be visible. This feature was added to the simulator instrument models but was not used in the evaluation trials.

Patient face model

A polygonal surface representation of the patient face was added

to the virtual model. This face model provides the student with a position and orientation cue, as well as providing additional realism.

Anatomical segmentation, and interactive selective segment display

The prototype anatomical model built in Alias was constructed in segmented pieces, corresponding to recognizable anatomical features. This pre-segmentation allows one to selectively display certain parts of the anatomy as well as to highlight selected anatomical features at run time. In addition, virtual tissue segmentation permits collision detection with procedurally

meaningful objects. The model based on the Visible Human dataset required an editing process to segment the surface anatomy.

Displaying endoscope position on CT scans as a navigation aid

As originally conceived the user interface to the system made use of an auxiliary CT view, in addition to the virtual endoscopic view of the anatomy. The system has the optional capability of updating the auxiliary CT image view with a crosshairs indicator of the current endoscope position. Although not used in the current evaluation configuration of the simulator, this feature may be useful for future trainng protocols.

Prototype simulator with expert system assistant

In parallel with this development project, HIT Lab researcher Mark Billinghurst extended his work with Jesus Savage-Carmona on intelligent multi-modal environments to create a prototype sinus surgery simulator with an integrated expert system assistant. Working with Oppenheimer and Dr. Edmond, Billinghurst developed a system which incorporates knowledge of the endoscopic procedure into a structured rule base, interprets the user's multi-modal inputs (currently voice and virtual endoscope position) and interacts with the user dyadically. While performing the simulated procedure, the user can query the system about anatomy and the specifics of the procedure, asking the system to identify features or demonstrate maneuvers. In turn the system recognizes the user's actions, and can provide vocal and visual feedback, as well as warnings when the user is about to execute a dangerous maneuver.

Although the system architecture is somewhat different, this prototype simulator provided ground work demonstration for certain features of the target surgical simulator, including navigation through the sinus cavity from an endoscopic viewpoint, use of abstract graphic overlays as a navigation aid, and embedding of surgical task sequences into simulator

Iterative Testing

In addition to these design experiments based on prototype modeling, the HIT Lab served as a test site for successive versions of the anatomical model and the simulation system. Having the target hardware platform (SGI onyx) in house made iterative evaluation a viable and useful approach. In particular, the proximity of the lab to Madigan Army Medical Center made it relatively easy for the evaluation team to consult with Dr. Edmond regularly on simulator features and system performance.

[development of haptic emulator]

Evolution of the Training Framework

During the early phases of the project we examined several other surgical simulator systems to look for places to improve the state of the art. What we noticed was that although these systems were making advances in anatomical modeling and user interaction, they were missing a structured educational component. In general these systems simulated a specific surgical domain or task only and were therefore "orphaned" experiences in the educational process.

We concluded that in order to be of significant educational value, one had to not only embed the simulator in the existing academic curriculum, but also develop curriculum within the simulation itself. This would not only serve the educational process but would also facilitate our evaluation of the educational effectiveness of the simulator.

The prevailing paradigm in surgical education is usually summarized as: "See One, Do One, Teach One." Our goal was to improve on this paradigm by taking the "See One, Simulate Many, Do One" approach that had proved effective in the domain of flight training. In order to achieve this we needed to develop a curriculum structure.

Task Analysis

Working with Dr. Edmond, we developed a taxonomy of ESS simulation objectives and simulator performance and interface features. Our objective, which proved far too demanding, was to elicit estimates from experienced ESS surgeons of the desirable system performance requirements for each training task. This approach did, however, provide us with a framework for approaching the issues of curriculum design in this domain.

ESS Domain Expert Survey

Twelve experienced ESS staff surgeons from several leading otolaryngology training programs were surveyed by Dr. Edmond

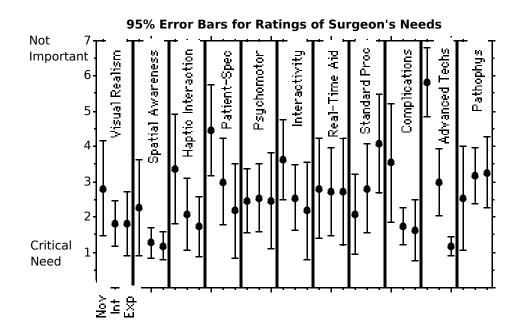
to assess their judgments of the primary simulation requirements for physicians performing ESS procedures. The primary objective of this survey was to determine the curriculum needs and perceived importance of several of the candidate features of the system early in its development.

These domain experts were asked to rate the relative importance (from "critical need" to "not important") for 11 simulator characteristics: visual realism, spatial awareness training, haptic interaction, patient-specific modeling, psychomotor training, real-time interactivity, real-time (intra-

operational) performance aids, standard surgical procedures, complications, advanced techniques, and pathophysiology.

In addition, they were asked to assess the value of these characteristics for target subjects at three levels of ENT experience: "novice" (i.e., junior residents), "intermediate" (senior residents), and "expert" (experienced ENT surgeons).

Results of this survey are summarized in Figure *, which indicates the mean rating for each characeristic for the three target groups, on a 1-7 scale.



As can be seen from this graph, these experts deemed "spatial awareness" the most crucial training need (of those presented in the survey) for all levels of subject experience, although "advanced techniques" were seen as equally critical for the experienced ESS target group. With the exception of "standard procedures" and perhaps "pathophysiology", all requirements were seen as equally important for all target levels or more important with increasing ESS experience.

It is interesting to note that almost all of these characteristics were rated on the "important" end of the scale. The characteristics deemed least important for simulation were "advanced techniques" and "patient-specific modeling" for the

novice group, and "standard procedures" for the advanced group. In addition, "real-time interactivity", "haptic interaction" and training in "complications" were seen as of only moderate importance for ESS novices.

Finally, it should be noted that these survey respondents were overwhelmingly favorable toward the development of an ESS training

simulator. While one or two expressed skepticism about any technological solutions, most respondents noted in their openended comments that this was an interesting and useful endeavor, and that the field had a need for such a system.

Simulator Integration

A considerable amount of effort was spent during the development phase on establishing a "curriculum wrapper" for the simulator. We initially envisioned the simulator as embedded within a multimedia training system which would provide a meaningful context and set the protocol for the trainee.

The development team elected instead to integrate relevant aspects of the emerging ENT curriculum into the simulator itself and to postpone further efforts to develop a total curriculum "package". The reasons for this included:

- development of a full-blown ESS curriculum was beyond the scope of this phase of the project
- it became apparent that a staged protocol approach was necessary to make the simulator effective as a training context
- useful techniques emerged (such as the use of navigation hoops and injection targets) which could be integrated relatively easily

The final product of the formative stage of the project was, in fact, more thoughtful with respect to an integrated ENT curriculum than had originally been anticipated and appears to be more useful as a training tool than other virtual reality medical simulators which have emerged in recent years. Its validity as a procedural simulator and its utility for training are evaluated in the following sections.

Summative Evaluation

After delivery and installation of the system at the HIT Lab by the Lockheed-Martin development team, the evaluation team

began a "shakedown" of the testing protocol on the three targeted subject groups (described below). Midway through this phase the system was relocated to Madigan Army Medical Center for further analysis by staff and resident ENTs.

The primary goals of this phase of the evaluation effort were to

- validate its utility as an ESS training environment,
- assess the usability of the system, and

provide additional iterative feedback to the development team.

The methods of investigation and results of this phase are described below, followed by a set of recommendations for further system development and evaluation derived from these findings. In a planned follow-on phase the primary focus of the evaluation effort will be on assessing the degree and nature of any transfer of this training to the real ESS operating environment (the ultimate objective of the simulator).

Methods

Subjects

Subjects were solicited from three distinct groups: (1) non-MDs with general intelligence and psychomotor abilities roughly comparable to the average otolaryngologist; (2) non-ENT physicians from a variety of specialties, and (3) ENTs with a wide range of ESS experience. We focused on these three groups in order to establish a baseline and asymptote for the evaluation of the efficacy of the simulator in training otolaryngology residents.

The first two groups also allowed us to "shake down" the system and research protocols without sacrificing valuable ENT resident subjects, and provided baseline scores for untrained/unfamiliar subjects. Finally, the non-ENT physician group could provide us with some valuable input about the extensibility of the simulator to non-ENT applications.

Non-MD Group: Twelve volunteers from the University of Washington College of Engineering comprised the non-MD group. These subjects ranged in age from 23 to 54, and included graduate students, professional staff and faculty. All had had some experience with simulation and virtual reality. None had previously used an endoscope or attended medical school.

Non-ENT Physician Group: Eight University of Washington MDs from specialties other than otolaryngology provided

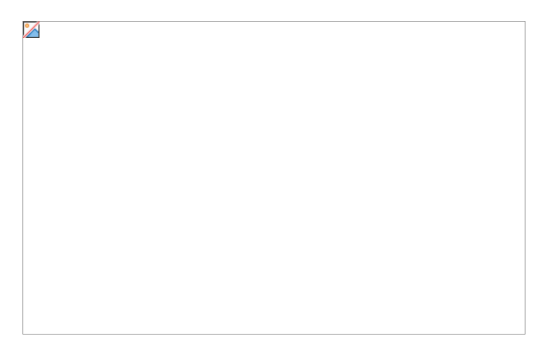
extensive feedback on the simulator design and utility for other medical and surgical tasks: 3 videoendoscopic surgeons, 2 radiologists, 1 neurosurgeon, 1 cardiologist, and 1 anesthesiologist. Four of these also provided us with complete trial performance data.

ENT Staff and Residents: Twelve staff and resident ENTs (1 female, 11 males) from MAMC served as subjects. The subjects in this group ranged in age from 28 to 46, with a mean age of 35.2 years, and a standard deviation of 6.15 years. One was left-handed, 10 right-handed, and the handedness of one was unknown.

ENT experience for this group broke down as follows:

- 4 staff (with an average of five years of training, six years of practice and more than 100 ESS procedures performed)
- 3 R2s (with an average of 1-5 ESS procedures performed or observed)
- 1 R3 (with 6-20 ESS procedures performed or observed)
- 2 R4s (with an average of 21-100 ESS procedures performed or observed)
- 2 R5s (with an average of 21-100 ESS procedures performed or observed).

Eight subjects had had occasional videogame experience, two reported playing videogames once, and two reported never playing videogames. Eleven subjects had had no virtual reality experience, while one reported occasional VR experience. Ten reported having no other simulator experience, while one reported one simulation experience and another reported "occasional" simulation experience. Because of the low incidence of prior experience with these systems, these factors were not evaluated further for this group.



Human Interface Technology Laboratory- 31 Figure * shows the distribution of the number of prior ESS procedures performed by the subjects in this group. The two reporting "1-5" procedures were observational only.

Procedures

All subjects proceeded through a common protocol for one or more sessions, involving:

- general orientation and consent form
- pre-session background questionnaire (see Appendix)
- instructions and orientation to the simulator and tasks
- one or more proctored simulation trials using one of the three training models
- post-session debriefing and questionnaire (see Appendix)

All trials were videotaped for later analysis and "think-aloud" comments. Subjects were free to terminate the session at any time.

MODELS

Three models were constructed to provide for a sequentially more realistic ESS experience. For each model, there was both a right- and left-side version. The anatomy was built for the right nostril and was reflected 180 degrees to simulate the left nostril. The nostril (left or right) with which the subject started was determined by their handedness.

Model 1: The novice/abstract model consisted of only the skin of the face and the entrance to the nasal cavity. A 3D grid pattern replaced the sinus anatomy to provide depth of field during the three tasks: Navigation, Injection and Dissection. Training aids were used to guide the subjects through the task. Navigation training aids consisted of virtual hoops and Injection training aids consisted of virtual targets in space.

During Navigation the subjects maneuvered the endoscope through four sets of virtual hoops. The paths of the hoops represented three passes (sets two and three combined are one pass) commonly taken before the surgery begins to gain familiarity with the patient's anatomy and to allow cleaning of the areas of interest.

Injection consisted of maneuvering both the endoscope and

instrumented forceps within the environment to inject five targets oriented obliquely in space. The instrumented forceps controlled a virtual needle for this task. The placement of the targets in space reflected the common areas of injection of a vasoconstrictor during a Maxillary Antrostomy.

During Dissection the subjects were also required to use both the endoscope and the instrumented forceps. The task consisted of dissecting each of a series of virtual spheres with pre-selected virtual tools. The instrumented forceps represented each of the tools most commonly used in the procedure: freer,

needle, bent needle, sickle knife, microdebrider, suction, straight-biting forceps, up-biting forceps, left-biting forceps and right-biting forceps.

Navigation through the four sets of hoops, injection of the five targets and dissection of each of the spheres was required for a complete score. Digitized voice audio cues were given for each hoop negotiated in Navigation, for the percentage of each target Injected, for percentages completed of each sphere during Dissection, and for final completion of each subtask.

Model 2: The intermediate model was composed of the Navigation and Injection training aids from Model 1, overlaid within a virtual anatomical model of the sinus cavity. Injection and Dissection followed the protocol for a Maxillary Antrostomy: injection of the inferior/anterior Middle Turbinate, superior root of the Middle Turbinate and the lateral nasal, followed by dissection of the Uncinate process, Bulla ethmoidalis and posterior Ethmoid cells. Labels were added for all anatomical structures with which the subject interacted.

Navigation through the four sets of hoops, injection of the five targets, medialization of the Middle Turbinate, dissection of the three anatomical structures, removal of two bone fragments placed in the Uncinate process and removal of three bone fragments placed in the Bulla ethmoidalis were required for a complete score. The widening of the Maxillary Ostium was not included in the procedure due to the inability to dissect enough of the lateral part of the Uncinate process for realistic viewing of the Ostium. Audio cues were given for each hoop negotiated in Navigation, for each target hit in Injection, for each bone fragment removed and for percentages completed for each anatomical structure in Dissection.

Model 3: The advanced model was composed of an anatomical model only. Subjects were expected to perform the three tasks without the training aids and to follow the protocol for a Maxillary Antrostomy. Three polyps were added superior/anterior to the Bulla ethmoidalis.

During Navigation the subject was required to perform the three

passes in the same order as in Models 1 and 2: inferior pass along the floor of the nose to the Nasopharynx, followed by a more superior pass medial to the Middle Turbinate towards the upper aspect of the Nasopharynx and Sphenoid Ostium, then rolling under the Middle Turbinate to inspect the Ostial Meatal Complex, and finally the superior pass medial to the root of the Middle Turbinate towards the Sphenoethmoidal Recess.

During Injection, the subject was cued only by the amount of blanching (whitening) of the virtual tissue as to whether more vasoconstrictor was needed. Dissection followed the protocol for a Maxillary Antrostomy. Navigation through the three passes, injection of the areas of interest, medialization of the Middle Turbinate, dissection of the three anatomical structures, dissection of the three polyps, removal of two bone fragments placed in the Uncinate process and removal of three bone fragments placed in the Bulla ethmoidalis were required for a complete score. Audio cues were given only at the end of each of the Navigation passes, for removal of bone fragments and for percentages completed in Injection and Dissection.

PROCTORING

During each trial, a proctor with knowledge of the procedure was present; proctoring for each of the groups varied according to their familiarity with the task. To assure that all records of the trial and all subject comments were noted, usually a second proctor was present. One proctor would be designated as an instructional proctor who would introduce the subject to the simulator and answer any questions during the trial. The second proctor would manage the forms, records and loading of the trials for the subject.

Non-MD Group

Each subject was initially introduced to the simulator and informed that we were evaluating the simulator as a possible trainer for residents in Otolaryngology. A summary of the reasons for this type of surgery was given along with a brief introduction to the anatomical structures and their locations in the sinus cavity to give the subjects a feel for the dimensions of the area in which they would be working.

Subjects were then introduced to the instrumented endoscope and informed of their ability to rotate the image axially by rotating the shaft of the endoscope. They were then introduced to the instrumented forceps and informed that the forceps would be simulating the virtual needle for Injection and all Dissection tools. The mechanics of the forceps (open and closed) were described as simulating the plunger for the syringe during Injection and opening and closing the jaws of the dissection

tools. They were then shown how the instrumented forceps would be positioned in the opposite nostril until the beginning of Injection.

Subjects were given a brief verbal description of the three tasks for Model 1 (Navigation, Injection and Dissection) and what would be required of them during the trial. They were informed that their introduction to Model 1 would be broken up across the three subtasks. A videotape of Dr. Edmond performing the trial was then started, during which time the subject was allowed to step up to the mannequin and become familiar with the instrumentation, while the proctor described the task in more detail. Subjects *Human Interface Technology Laboratory-* 38 were encouraged to ask any questions and "speak-aloud" during the entire procedure.

Following instructions and video for Navigation, the subject independently performed the Navigation task. The trial was then paused, proctoring and video for Injection was given, and the Injection task was performed. At that time the proctor would pull the instrumented forceps across the Columella Nasi into the nostril being used and place a plug in the original nostril to inhibit re-crossing of the Columella Nasi by the instrumented forceps during the remainder of the trial. After Injection, the trial was then paused, instructions and video for Dissection were provided, and the Dissection task was performed.

Model 1 was the only model where this process was used and it was used only on their initial introduction to the simulator. On subsequent trials of Model 1, they were given verbal proctoring instructions only, and allowed to perform the task. Those who were introduced to Model 2 did not require breaking the trial into subtasks.

Progression to Model 2 was based on performance on Model 1. An average score of 54% was required of the subjects on Model 1 before progressing to Model 2. Since subjects in this group had no familiarity with the procedure or with paranasal sinus anatomy, Model 2 required extensive proctoring instructions on locations and anatomy to dissect during the procedure. No subjects from this group were run through Model 3 because of their inability to achieve adequate proficiency in Model 2 during the time course of the study.

Non-ENT Physician Group

Again, each subject was initially introduced to the simulator and informed that we were evaluating the simulator as a possible trainer for residents in Otolaryngology. They were encouraged to think of ways in which this type of simulator could be used in their own fields. They were then introduced to the virtual endoscope, informed of their ability to rotate the image axially by rotating the shaft of the endoscope, and of the availability of a 30 and 70 degree scope which could be swapped for the zero degree

scope they would initially be given. The optics of the 30 and 70 degree scopes were explained where necessary.

Subjects were then introduced to the instrumented forceps and informed that the instrumented forceps would simulate the virtual needle for Injection and all Dissection tools. The mechanics of the instrumented forceps (open and closed) were described as simulating the plunger for the syringe during Injection and opening and closing the jaws of the dissection tools. They were

then informed of how the instrumented forceps would be positioned in the nostril opposite the one used in the trial, until the beginning of Injection.

The subjects were then given a brief verbal description of the three tasks of Model 1 (Navigation, Injection and Dissection) and what would be required of them during the trial. A video of Dr. Edmond performing the trial was then started, while the proctor continued to describe the subtasks in more detail and what was required for completion of the trial. During this time the subject was allowed to step up to the mannequin and become familiar with the instrumentation. The "blood effects scope" (scope becoming opaque within a set time interval) was shown in the video and reasons for it were described by the proctor, along with how to relieve the problem by wiping the "scope" on the foam pad located on the mannequin's forehead.

The subject was encouraged to ask any questions and "speakaloud" during the procedure. This process was repeated before their first introduction to Model 2, with the appropriate, modelspecific changes. On subsequent trials of Model 1 and Model 2 they were given verbal proctoring instructions only before performing the task.

The subject's familiarity with the procedure determined the need for further instruction by the proctor. On average this group had no familiarity with the procedure, but all had prior introduction to anatomy (during medical school) and had an understanding of the reasons for the procedure. Progression to Model 2 for this group was based on performance on Model 1; an average score of 69% was required on Model 1 before progressing to Model 2. Proctoring for Model 2 required instructions on locations and anatomy to dissect during the procedure. Training aids and video of the procedure provided adequate introduction to the task for Model 1's entirety and for Model 2's Navigation and Injection tasks. No subjects from this group were run through Model 3, due primarily to the limited availability of these MD subjects.

ENT Staff and Residents

Again, each subject was initially introduced to the simulator and informed that we were evaluating it as a possible trainer for residents in Otolaryngology. They were introduced to the virtual endoscope, informed of their ability to rotate the image axially by rotating the shaft of the endoscope, and of the availability of a 30 and 70 degree scope which could be swapped for the zero degree scope they would initially be given.

Subjects were then introduced to the instrumented forceps and informed that they would simulate the virtual needle for Injection and all Dissection tools. The mechanics of the instrumented forceps (open and closed) were described as simulating the plunger for the syringe during Injection and opening and

closing the jaws of the dissection tools. They were then informed of how the instrumented forceps would be positioned in the nostril opposite the one used in the trial until the beginning of Injection.

The subjects were then given a brief verbal description of the three tasks of Model 1 and what would be required of them during the trial. The video of Dr. Edmond performing the trial was then started, while the proctor continued to describe the subtasks in more detail and what was required for completion of the trial. During this time the subject was allowed to step up to the mannequin and become familiar with the instrumentation.

The "blood effects scope" was shown in the video and reasons for it were described by the proctor, along with how to relieve the problem by wiping the scope on the foam pad located on the mannequin's forehead. Subjects were encouraged to ask any questions and "speak-aloud" during the procedure. On subsequent trials they were given verbal proctoring instructions only (with the appropriate, model-specific changes), and allowed to perform the task.

The subject's familiarity with the procedure determined the need for further instruction by the proctor. Typically, the staff Otolaryngologists needed no further instruction on the procedure for the remainder of the session, except for the need to be shown the active areas for dissection of the Uncinate process, Bulla ethmoidalis and Posterior Ethmoid cells within the virtual model. In general, the staff ENT subjects ran through all three model levels.

Similarly, the ENT residents needed no further instruction for Model 1. For Model 2, however, more detailed instructions were sometimes needed during Dissection on what anatomy to dissect and where the active dissection areas were located within the anatomy. In Model 3 more detailed instructions were also needed during Navigation on the order of passes to perform and where the active dissection areas were located within the anatomy. All residents were run through Model 1-right, Model 2right, Model 3-right, Model 3-left and Model 2-left.

USER PERFORMANCE CRITERIA

The scoring algorithm for Version 1.2 takes into consideration two major performance measures for Endoscopic Sinus Surgery: Accuracy/Completeness and Time. Trial scores consisted of the sum of the subtask scores, each calculated as:

*score=accuracy*optimal-time/completed-time.*

The scoring algorithm used for the evaluation of this phase of the project took into account the most important skills needed to perform sinus surgery, as well as the level of difficulty of the model. In an attempt to normalize

across model conditions, the following optimal times (in seconds) were used in this scoring equation:

Navigation: advanced=97, intermediate=97, novice=97 Injection: advanced=90, intermediate=165, novice=90 Dissection: advanced=815, intermediate=635, novice=165

These times were derived from the approximate performance times for the project's lead otolaryngology advisers.

[MORE DETAILS ON SCORING ALGORITHMS HERE]

Results

Non-MD Evaluation

Although not the target domain audience for this simulator, the non-MD group gave us an opportunity to look at the baseline difficulty of the simulator for domain-naive users and to further work out requirements for instructional presentation, proctoring, and pacing of the "curriculum" (after initial "shake down" and protocol development by the project staff). Since the novice model did not require interaction with the complex paranasal sinus anatomy or detailed knowledge of the surgical procedure, it was felt that this group should be able to complete those trials.

The inherent difficulty of the task for inexperienced users is evidenced by subject attrition and by their initial trial scores. Indeed, complete trial data are available for only nine of the 12 subjects in this group. Of the three lost to attrition, one terminated the session due to disorientation and discomfort with visual interface, and two were unable to complete even the navigation task during the time period available for testing. The primary difficulty observed with these three subjects was an inability to adapt to the psychomotor demands of the interface; that is, they were not able to acquire the basic skill necessary to control the positioning of the virtual endoscope within the model.

Of the nine remaining subjects in this group, five had had considerable experience with videogames and three with commercial flight simulators. Thus the trials scores for this group may represent the high end of non-MD users and should not be taken as a sample from the general non-MD population.

Novice model performance

Still, despite attrition of the low-end performers and the unusual prior experience of the remaining subjects, the non-MD group performed significantly worse than the ENT group on their initial novice level trials (as can be seen graphically in a later section).

Unpaired Student's t comparisons with the ENT group revealed significantly

lower overall trial scores (nonMD mean = 45.9, ENT mean = 65.1, t = 2.87, p = .0098),

longer overall trial times (nonMD mean = 1012 sec, ENT mean = 692 sec, t = 2.14, p = .0458),

lower navigation scores (nonMD mean = 47.1, ENT mean = 69.4, t = 2.39, p = .0276),

longer navigation times (nonMD mean = 239 sec, ENT mean = 146 sec, t = 2.52, p = .0209), and

lower injection scores (nonMD mean = 55.3, ENT mean = 81.5, t = 2.46, p = .0235).

Injection times, while not significantly longer, did approach significance (nonMD mean = 257 sec, ENT mean = 117 sec, t = 1.96, p = .0649). Surprisingly, while dissection scores and dissection times were slightly worse for the non-MD group, these differences were not significant. This relatively poor performance by the ENT group on their initial dissection trial is discussed further below.

Practice effect

Six of the 12 subjects in this group had complete trial scores for at least two novice trials. Student's t comparisons of their first and second novice trials showed that their second trial resulted in significantly

higher overall trial scores (trial 1 mean = 46.7, trial 2 mean = 69.5, t = 2.33, p = .0423),

higher injection scores (trial 1 mean = 53.8, trial 2 mean = 86.7, t = 2.22, p = .05),

shorter dissection times (trial 1 mean = 492 sec, trial 2

Human Interface Technology Laboratory- 47 mean = 328 sec, t = 2.39, p = .0382), and

higher dissection scores (trial 1 mean = 35.8, trial 2 mean = 51.8, t = 2.55, p = .0287).

Total trial times, navigation times and scores, and injection times all changed in the expected direction (that is, they improved across trials), but none of these improvements was statistically significant. It appears that, for this nonMD group, the primary challenge may have been mastering the endoscope positioning.

Intermediate model performance

First-time intermediate level scores were acquired for only 3 of the subjects in the non-MD group. Despite this small N, unpaired Student's t comparisons with the ENT group revealed significantly

lower overall trial scores (nonMD mean = 53.7, ENT mean = 76.7, t = 3.53, p = .0041),

longer overall injection times (nonMD mean = 359 sec, ENT mean = 132 sec, t = 6,91, p = .0001), and

lower injection scores (nonMD mean = 47.1, ENT mean = 69.4, t = 2.39, p = .0276).

longer navigation times (nonMD mean = 239 sec, ENT mean = 146 sec, t = 2.52, p = .0209), and

lower injection scores (nonMD mean = 46.0, ENT mean = 94.4, t = 8.09, p = .0001).

Overall trial times, while not significantly longer, did approach significance (nonMD mean = 1532 sec, ENT mean = 1215 sec, t = 1.79, p = .098), as did navigation times (nonMD mean = 199 sec, ENT mean = 138 sec, t = 2.08, p = .0592) and navigation scores (nonMD mean = 48.0, ENT mean = 73.7, t = 1.99, p = .0698). Again, no difference was found between the two groups in initial dissection times and dissection scores on the intermediate model.

The experiential advantage of the ENTs is perhaps most telling in their superior injection task performance. As will be seen below in examining trends within the ENT group, ability to perform efficient injections may by the hallmark of the accomplished surgeon, much more so than the ability to perform efficient dissection (at least at indicated by simulator performance).

Non-ENT Physician Evaluation

The primary objectives of the non-ENT physician evaluations were to provide a medically-trained comparison and "shakedown" group and to explore the perspectives of other medical specialties regarding additional potential applications for this sort of simulation training. In particular, we were interested in applications which might make use of the integrated approach and specific components incorporated into the Madigan ESS simulator.

Novice model performance

Novice performance measures for the non-ENT MD group were very similar to the those for the experienced ENT group (as summarized graphically below in the ENT analysis section). While there were no statistically significant differences between these two groups, it should be noted that the mean differences for all of the performance measures were in the direction we had expected (that is, the means performance measures were consistently better for the ENT group).

This finding may be due to the considerably broad prior medical and interface experience for all of the subjects in this group. Most of the subjects in this group have been involved over the years in developing or testing novel interface devices for medical tasks, and several had previous experience test-driving similar medical simulators. Perhaps the greatest mitigating factor, however, is the extensive experience that two of these four subjects had acquired as videoendoscopic surgeons. This experience provided them with a background comparable to the ENT group in basic endoscopic psychomotor skills, as well as in general procedural knowledge and confidence.

Although there were no statistically significant differences in any performance measures between the non-ENT MDs and the non-MD subjects, it was apparent that they were much more comfortable with the experimental task requirements. This finding (and the nature of the questions asked during the session) suggests the value of a general medical perspective in assuring confidence with this sort of procedural simulator.

Intermediate model performance

Three of the non-ENT MD subjects also tried the intermediate model. Comparison of the non-ENT MDs with the ENTs on their first trial on the intermediate model reveals an interesting finding. While there was no significant difference between the groups in overall score and trial time, the non-ENTs did notably better than the ENTs on the dissection subtask. Dissection scores were significantly better for the non-ENTs (nonENT mean = 84.0, ENT mean = 63.1, t = 2.439, p = .0312), which appears

to be due primarily to their faster times on that task (nonENT mean = 662 sec, ENT mean = 880 sec, t = 1.99, p = .0692), a non-significant but clearly suggestive finding.

Looking at individual scores, we note that this effect is due to the relatively fast dissection times for the two videoendoscopic surgeons. While we would expect that the ENT group would be superior to all other groups on this task, the extensive background in similar procedures appears to have prepared the general surgeons well for this task.

Related Medical and Surgical Applications

Response by the non-ENT physician group was uniformly positive. Each of them suggested additional applications, both within and outside of their particular specialties, that would be amenable to a similar simulation approach. These suggested applications included:

- Neurosurgery: approaching the pituitary gland through the sphenoid sinus
- Anesthesiology: intubation, bronchioscopy

Videoendoscopic surgery: cholecystectomy, bladder, joints, throat, chest procedures

Cardiology: cannulization, thoracoscopic surgery

Gastroenterology: endoscopic procedures

Radiology: flouroscopy, intrusive ultrasound procedures (e.g., transesophageal echocardiography)

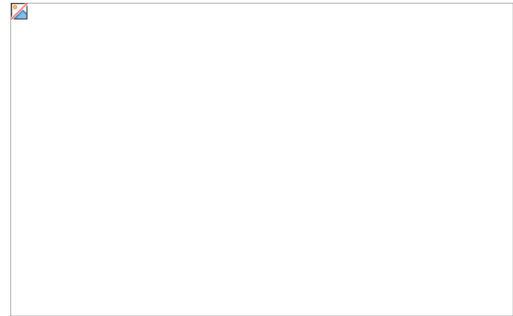
In general, procedures that use endoscopes or probes to explore internal structure and to perform manual procedures should be amenable to this simulation approach. Several of the physicians in this group indicated that they would like to see work proceed in that direction and that there would be support for this development from the leaders within their specialties.

Staff and Resident ENT Evaluations

Simulator performance measures were acquired on 53 separate trials for the 12 ENT subjects between 5/13/97 and 8/12/97 (23 trials run at HITL, and 30 trials run at MAMC). The number of trials acquired for each subject ranged from 2 to 12, as shown in Figure *, with all subjects being tested (minimally) on both the novice and the intermediate model.

Initial Novice Model Performance

Figure * shows the distribution of trial scores on the initial novice trials for the 12 ENT subjects.



To examine the degree to which prior OR experience might contribute to these initial score differences, we calculated the Pearson product-moment correlation coefficients across the primary performance measures (trial time and trial score) and the primary experience measures (age, years of ESS training, and approximate number of actual ESS procedures performed. A matrix of these correlations appears in Table * (critical r = .6021for p < .05, df 9).

	score	ESS perf	years t	age	trialtime
score	1				
ESS perform	.526	1			
years_traini	.38	.923	1		
age	.236	.662	.713	1	
trialtime	966	652	514	33	1

Note: 1 case deleted with missing values.

As can be seen, the best predictor of novice trial time from among the surgical experience measures is the number of actual ESS cases performed (r = -.652), with trial time decreasing significantly with surgical experience. For trial score, the findings are less conclusive, although the best experiential predictor is again the number of ESS cases performed (r = .526). While age is naturally correlated with ESS experience and training, age alone does not appear to be a significant predictor *Human Interface Technology Laboratory-* 54 of simulator performance for these ENT subjects.

In general we see that trial score on the initial novice simulator trial is indeed positively correlated with prior ESS experience. This finding suggests that the simulator provides a valid reflection of the skills acquired in ESS procedures.

Subtask Performance on Initial Novice Trial

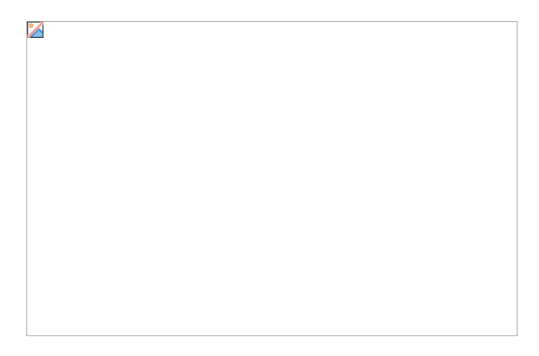
Not surprisingly, overall trial scores are, with the sole exception of injection time, significantly correlated with subtask scores and times, as shown in Table * (critical r = .5760 for p < .05, df 10). Overall trial times are similarly related to subtask performance measures, although by far the strongest predictor of overall trial time is performance on the dissection task (r = .897 for dissection time and -.918 for dissection score).

	score	trialtime	navtime	navscore	injtime	injscore	disstime	disscore
score	1							
trialtime	965	1						
navtime	587	.469	1					
navscore	.68	573	97	1				
injtime	531	.508	109	.092	1			
injscore	.684	667	.016	.033	957	1		
disstime	784	.897	.189	342	.3	476	1	
disscore	.856	918	275	.413	352	.524	964	1
ESS perf	.504	614	245	.227	425	.428	526	.471

As before, prior ESS experience is predictive of overall trial time (r = -.614), and, although its relationship to each of the subtask measures is consistent with simulator validity, none of these correlations are statistically significant.

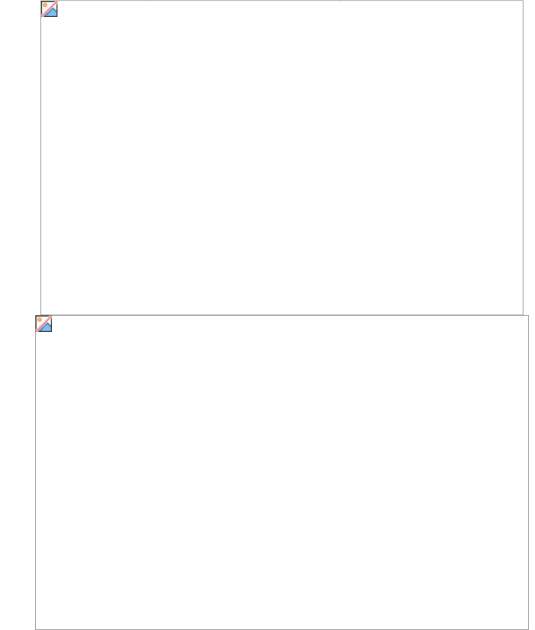
Novice Model Performance Across ENT Groups

Figure * shows the breakdown of first-time novice (abstract) model scores for this group. The linear relationship between overall novice trial score and year of residency is apparent from this mapping. It is interesting, although perhaps not surprising, to note that R5 residents appear to have higher scores on the simulator than do the experienced ENT staff subjects.



Similarly, it can be seen in Figure * that overall trial times for the initial novice trials fall off as expected with year of residency. Both of these findings provide strong evidence for the validity of

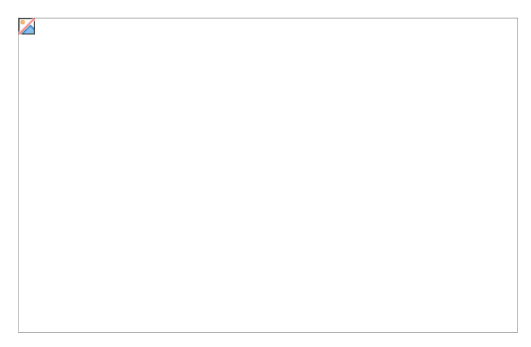
Human Interface Technology Laboratory- 57 the simulator for the ESS task, even when the model is abstract.

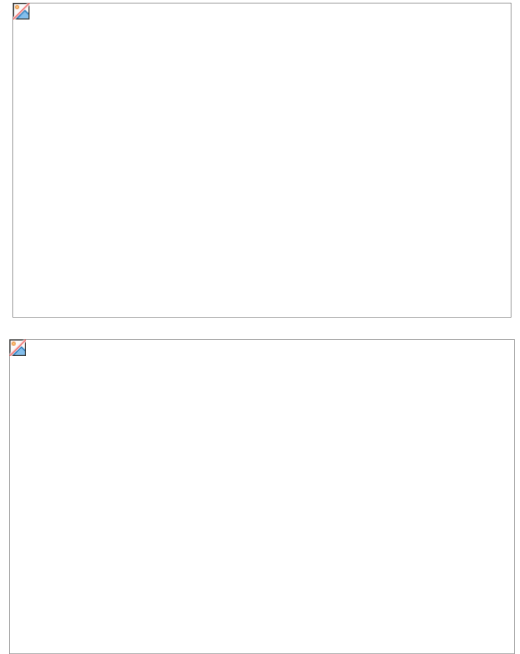


Subtask Times By Year of Residency

Subtask Scores By Year of Residency

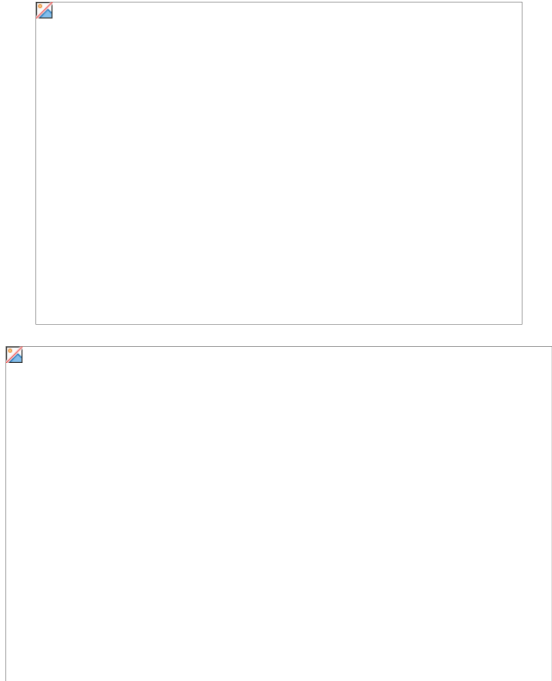
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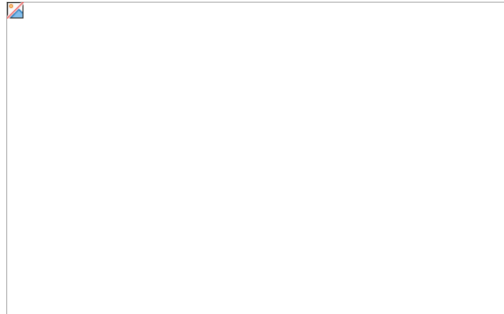
Novice Model Performance Across Subject Groups

Figure * shows the distribution on overall scores on the first novice trial for all three subject groups, while Figure * shows the same thing for overall trial times.



Initial Intermediate Model Performance

Figure * shows the distribution of trial scores on the initial intermediate trials for the 12 ENT subjects (Note that the first intermediate trial for one of these subject was terminated after the navigation phase and is consequently not included here).



Although we might expect a positive relationship between ESS experience and novice model performance, we would expect that OR experience would be especially predictive of performance when the sinus anatomy serves as the model. A matrix of these correlations appears in Table * (critical r = .6319 for p < .05, df 8).

	score	ESS perf	years_t	age	<u>trialtime</u>
score	1				
ESS perform	.555	1			
years_traini	.356	.931	1		
age	.581	.667	.68	1	
trialtime	795	144	014	475	1

In contrast to the initial novice scores, these results indicate that the best predictor of initial intermediate trial time (from among the surgical experience measures) is subject age (r = -.475), but this correlation is not statistically significant. For trial score, the findings are also inconclusive, with the best experiential predictors being subject age (.581) and the number of ESS cases performed (r = .555).

In general we see that initial performance on the intermediate model is indeed positively correlated with prior ESS experience,

but the relationship is not as strong as for initial novice model performance. Table * attempts to shed light on this unexpected finding by looking more closely at experiential

correlates of performance on procedural subtasks for the initial intermediate trial (critical r = .6319 for p < .05, df 8).

	ESS perf	<u>years_t</u>	age	navtime	navscore	injtime	injscore	disstime
ESS perf	1							
years_tr	.931	1						
age	.667	.68	1					
navtime	569	314	386	1				
navscore	.509	.288	.36	971	1			
injtime	869	847	624	.435	413	1		
injscore	.859	.798	.525	476	.446	967	1	
disstime	.199	.229	428	.191	224	275	.362	1
disscore	059	144	.462	354	.381	.186	226	967

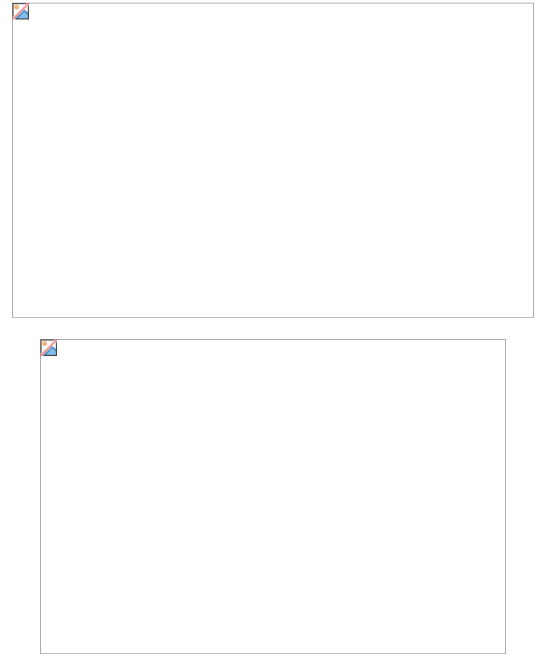
Note: 1 case deleted with missing values.

It appears from Table * that prior ESS experience is indeed a strong predictor of performance on the initial intermediate injection task (r = -.859 for injection time and .859 for injection score) and a slightly weaker predictor of performance on the initial intermediate navigation task (r = -.569 for navigation time and .509 for navigation score).

Performance on the initial intermediate *dissection* task, however, is actually the reverse of our predicted effect: dissection times are slightly longer for more experienced surgeons on their first encounter with the intermediate model, and dissection scores are slightly lower.

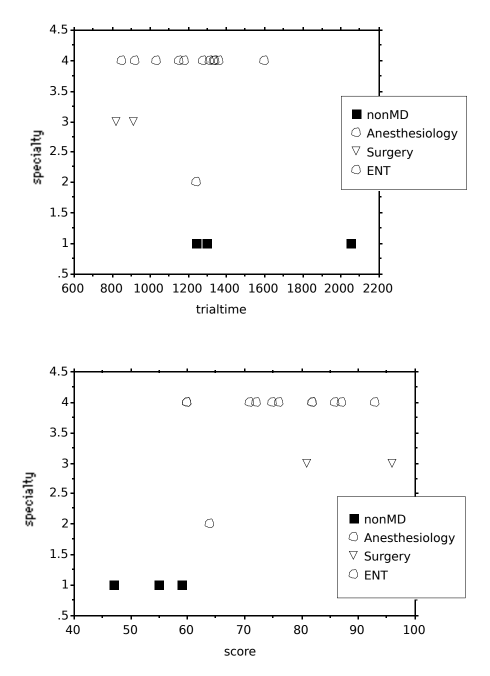
These findings are deserving of further study. One possible explanation might be that there is significant negative transfer to the simulated dissection task due to more extensive experience with the real instruments and real anatomical dissection. Comments from experienced ESS surgeons regarding the difficulty of the dissection task support this notion.

Intermediate Model Performance Across ENT Groups



Intermediate Model Performance Across Subject Groups

Overall times and scores for initial intermediate model trials are broken down by specialty in Figures * and *. As expected, the ENT group performed better on the anatomical model than did the non-MD group. Surprisingly, the other (non- ENT) physicians in this sample had scores that were comparable to the ENT group. (Note that the radiologist from the novice trial comparisons above was not tested on the intermediate or advanced models.)



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While the trial time difference between the ENT and non-MD groups was not statistically significant, the trial score difference was (as summarized in Table *. This finding again suggests that ESS experience is predictive of simulator performance, thus providing evidence for the validity of the system for ESS training.

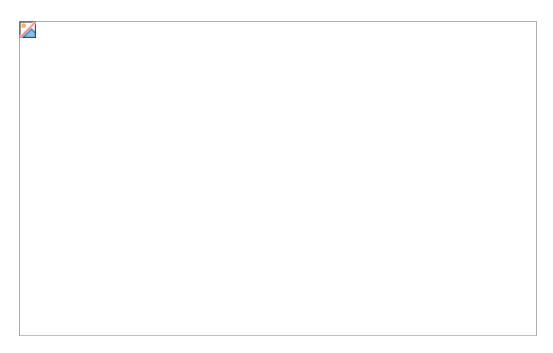
DF:		Unpaired t Val	ue: Prob. (2-tail):	
	12	-3.533	.0041	
Group:	Count:	Mean:	Std. Dev.:	Std. Error:
nonMD	3	53.667	6.11	3.528
ENT	11	76.727	10.631	3.205

Unpaired t-Test X 1: specialty Y 1: score

Initial Advanced Model Performance

Advanced Model Performance Across ENT Groups

Figures * and * show the overall scores and trial times for the first advanced trial by subjects in the ENT group. Note that two of the staff subjects did not attempt the advanced model.



Initial performance on the advanced model revealed a tendency for the more senior residents to perform faster, but perhaps less accurately.

The R3 and the 2 R4s had shorter trial times on their first advanced trial, but the R3s also had lower scores on those trials, indicating that their accuracy was lower.

While overall time and score for staff ENTs did not appear to differ systematically from the resident times and scores on initial advanced model trials, their performance on the *injection* subtask appeared to be superior to the resident group (as shown in Figures * and *). Differential performance was not seen for the navigation and dissection subtasks. Although navigation and dissection may vary with individual care and priorities, the data suggest that injection efficiency may be a hallmark of the experienced sinus surgeon. This finding should be followed up and verified in future evaluation studies.

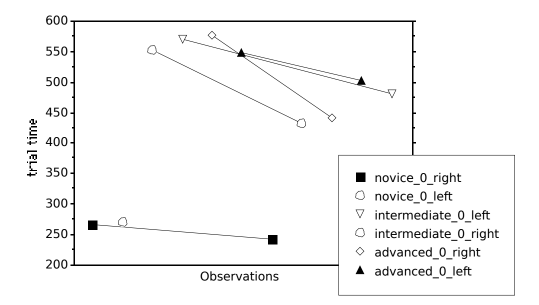
Asymptotic Performance

While extensive learning curve data are not available, repeated measures on the steady-state system (version 1.2) are available for our two primary evaluation proctors (CE, an ESS surgeon, and CA, an engineering student). Given that these two subjects had each tested the simulator in various degrees of development approximately 30-40 times over a 6-8 month period, their later scores may provide an estimate of asymptotic performance on the simulator.

ENT steady-state performance

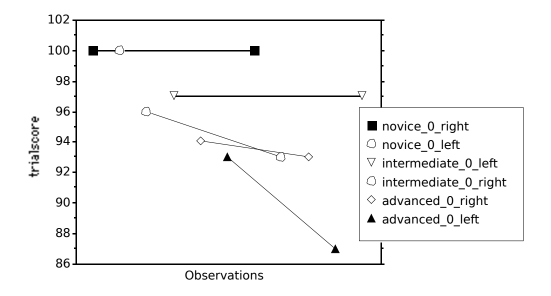
Simulator performance scores were analyzed for the last eleven completed steady-state trials by the experienced ENT proctor. As can be seen in Figure *, steady state trial times for the novice model (mean = 259 seconds, s.d. = 15.9) were lower than those for the intermediate and advanced models (combined mean = 512 seconds, s.d. = 57.1). The similarity between trials times on the intermediate and advanced models may suggest that the two

tasks are comparable for the highly experienced subject and, further, that the intermediate model training aids are no longer necessary for this subject.

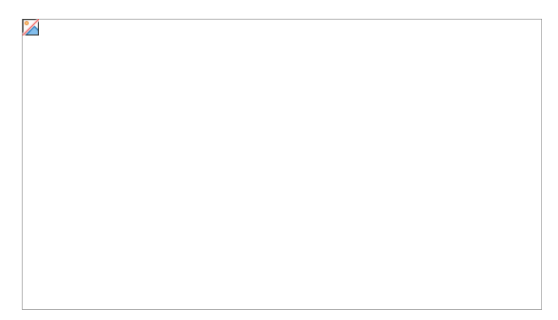


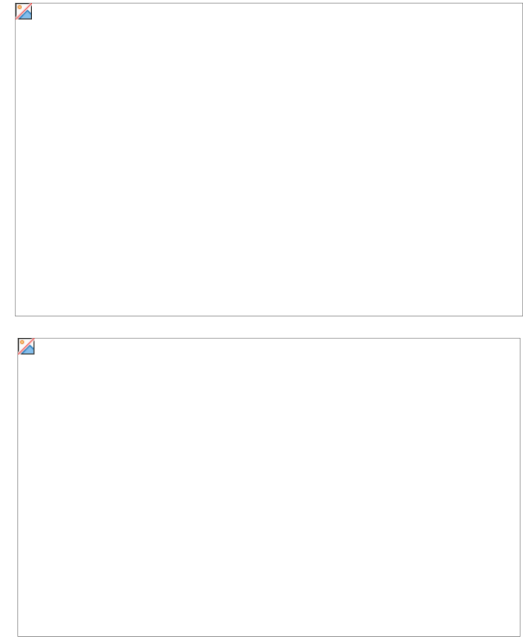
It also appears from this figure that trial times for each model are consistently lower in the last trial than in the previous trial, suggesting that these trials times may, in fact, *not* represent steady state performance. An examination of overall scores for these trials (see Figure *) reveals, however, that for at least some of these conditions the decrease in overall trial time for these later trials came at the cost of trial score. It should be noted that, since this subject achieved the minimum required time on all subtask trials, these trial scores precisely reflect a decrement in performance *accuracy*.

Given the better trial score for the first samples for each model condition, we may tentatively treat the trial times for those first samples as more representative of optimal performance.

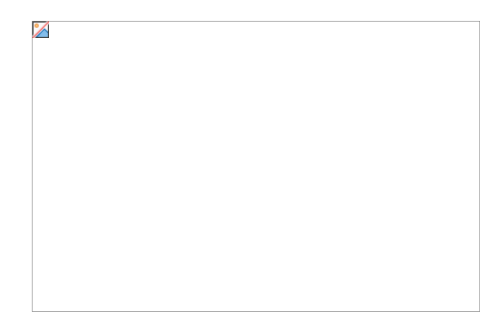


It may also be instructive to look at asymptotic performance for the three subtasks for this highly experienced ENT subject. In Figures *-* we present the subtask times for this same set of trials. As can be seen from these figures, the systematic decrease in overall trial time for intermediate and advanced model (shown in Figure *) appears to be due primarily to a systematic decrease in completion time for the dissection subtask, while navigation and injection times appear to vary more randomly across these temporal samples.

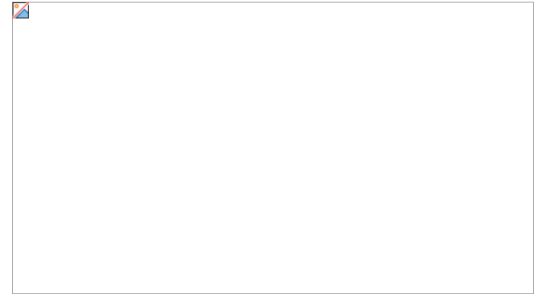




Similarly, examination of Figures *-* suggests that the overall decrease in trial score for the intermediate and advanced models for this experienced subject are due to decreases in navigation score for the advanced model condition and a decrease in dissection score for one of the intermediate models (intermediate-0-right). Injection performance and performance on the navigation and dissection subtasks for the other models appear to indeed be at steady-state asymptotic levels.



%



Collapsing across model over these 11 trials for the expert ENT subject, we observe the Pearson product-moment correlation coefficients shown in Table * among these performance variables (critical r = .6021 for p < .05, df 9).

	<u>trialscore</u>	navtime	navscore	initime	disstime	disscore	<u>trial time</u>
trialscore	1						
navtime	.037	1					
navscore	.856	.119	1				
injtime	718	.384	608	1			
disstime	643	.29	387	.642	1		
disscore	.557	136	.049	416	593	1	
trial time	609	.424	357	.689	.987	581	1

These findings suggest that (for this subject, at least) the best asymptotic predictor of overall trial score is navigation score (r = .856), and that the best predictor of overall trial time is dissection time (r = .987), as suggested above.

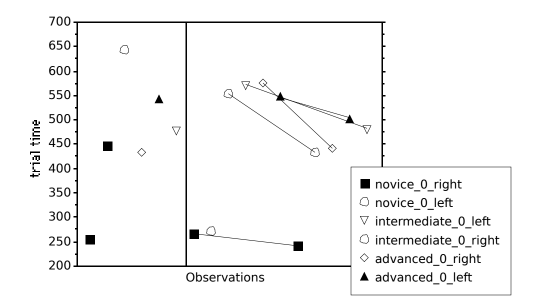
Non-ENT proctor steady-state performance

When we look at the correlations among these same performance variables for the last six trials of the other highly experience proctor (CA), we see the same result, as shoen in Table * (critical r = .8114 for p < .05, df 4). Again, the best predictor of overall trial score is navigation score (r = .924), and the best predictor

Human Interface Technology Laboratory- 78 of overall trial time is dissection time (r = .657), although the latter relationship is not statistically significant.

	<u>trialscore</u>	navtime	navscore	injtime	disstime	disscore	trial time
trialscore	1						
navtime	459	1					
navscore	.924	193	1				
injtime	686	.795	592	1			
disstime	889	.627	664	.657	1		
disscore	.694	805	.37	602	916	1	
trial time	471	.151	27	079	.657	626	1

Indeed, if we plot the trial times and trial scores for this proctor side-by-side with the values for the expert ENT proctor, we see that they are remarkably similar. In Figures * and * we see the trial times and scores, respectively, for the non-ENT proctor on the left-hand side of each graph and the expert ENT proctor on the right-hand side of the each graph. Again, trial times on the novice model are in the 250-second range, while trial times for the other model condition are in the 550-second range for the non-ENT proctor.



These findings suggest that extensive experience with the simulator may afford a non-ENT subject performance values which are comparable with an experienced ENT proctor. Assuming the validation findings discussed above, this bodes well for the training effectiveness of the simulator.

Post-Session Questionnaire

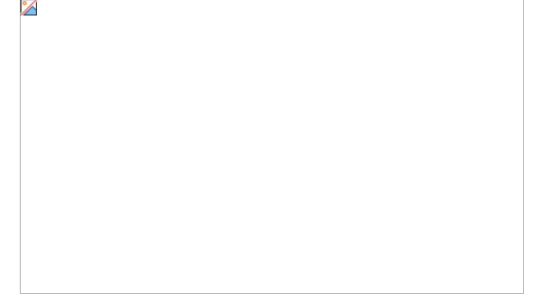
All 12 ENT subjects provide us with post-session questionnaire data. All scale responses indicated here are for 10-point scales anchored at the end points.

Proctor's Instructions: Most subjects found the proctor's instructions "very useful" (6/12) or "adequate" (4/12). The one area singled out for improvement was "endoscope and tool handling".

Simulator Layout: Simulator layout and interaction with the model and instruments was rated as moderately realistic (mean = 6.5 and 6.2, respectively).

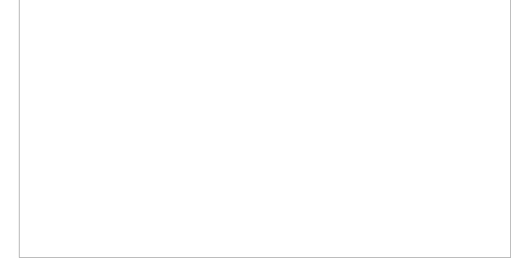
Abstract Model: The assessed training benefit of the abstract model was relatively variable (mean = 5.3, s.d. = 2.6), perhaps reflecting the variability in baseline experience of the subjects. The assessed training value of each of the five "training aids" (hoops, targets, dissecting spheres, voice feedback, and heart

rate) is shown in Figure *. Heart rate was viewed as the least beneficial. Most subjects indicated that the level of difficulty of the abstract model should remain the same (9/12) or increase (3/12).



Intermediate Model: Subjects overwhelmingly indicated that the experience with the Novice Model prepared them for Model 2 (mean = 8.3, s.d. = 1.5). Dissection was judged as more difficult in Model 2 than in Model 1 (mean = 7.1), while navigation and injection were seen as about the same level of difficulty (mean = 5.5 and 5.0, respectively).

Subject ratings of the benefits of the Model 2 training aids are shown in Figure *. As is evident from the non-overlapping notches in the box plots, the heart rate cue was judged to be significantly less beneficial than the hoops, targets or voice feedback. Furthermore, note that the ratings for these training aids are considerably higher for the intermediate model than for the abstract (novice) model. One possible explanation for this surprising finding is that in the novice model the cues were useful for training psychomotor skills and instrument control, while in the intermediate model they were *also* useful for training the specific surgical procedure.



1

Ratings of realism of the anatomical model were also extremely high, with a mean across respondents of 8.0 and no ratings below "6" on the 10-point scale. It is also interesting to note (in examining the correlations among survey responses) that the more realistic the anatomical model was rated by these subject, the easier the Model 2 subtasks were perceived to be (r = -.503, -.477, and -.499, for navigation, injection and dissection, respectively).

Advanced Model: Figure * presents a summary of the distributions of responses by the ENT subjects to all questionnaire items regarding the advanced model. Question 12 ("post12") suggests that the vast majority of these subjects felt that the experience with Model 2 was highly beneficial for performing on Model 3.

1

In rating the difficulty of Model 3 on the three essential tasks ("post13nav", "post13inj", and "post13diss") subjects indicated that Model 3 was moderately more difficult than Model 2 (a surprising finding, which may indicate some confusion about the question). Finally, the level of procedural realism for Model 3 was rated quite high, with a mean of 8.0 on the 10-point scale ranging from "far from reality" to "close approximation".

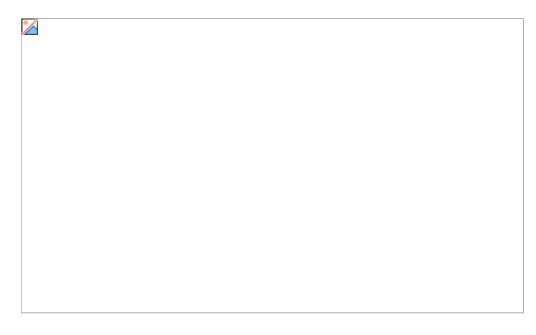
Force Feedback: Seven of the 12 ENT subjects rated the use of the haptic subsystem. When asked whether they preferred the force feedback on or off, 4 preferred it "on", one preferred it "off", and one initially preferred it "off" but later preferred it "on". When asked if they preferred the force display on the forceps or on the endoscope, 6 indicated a preference for forces on the forceps, while 1 indicated a preference for forces on the endoscope. (Note that they only experienced the forces on the forceps, not on the endoscope, so these repsonses were partially speculative.

Ratings of the realism of the force feedback were moderately low, with a mean of 5.1 and a standard deviation of 1.7 on the 10point scale from "far from reality" to "close approximation", although one of the 7 subjects rated it an "8" and none of the subjects gave it a realism rating below "3".

Overall Evaluation: Responses to items regarding the overall evaluation of the simulator are summarized in Figures * through *. Level of difficulty of the simulator for the three subtasks is assessed in Question 18 "post18nav", "post18inj", and "post18diss"); the results suggest that the tasks were perceived as roughly the same level of difficulty as an actual procedure, with the exception of the dissection task, which is perceived as somewhat more difficult on the simulator.

1

In assessing the training value of each model (items "post19m1", "post19m2", and "post19m3") for themselves, there appears to be a linear trend from the novice model to the advanced model, with all seen as valuable but the Advanced model especially so.



In assessing the adequacy of the virtual anatomical model for developing proficiency at the common tasks ("post20nav", "post20inj", and "post20diss" in Figure *), these subjects indicate that the model is quite adequate for training navigation and injection skills, but may be less than adequate for training dissection skills.

1

When asked to indicate what levels of ENT training would benefit from exposure to the simulator, the 10 ENT respondents offered the following distribution of responses, suggesting that in its current implementation, the simulator is best suited for junior residents (as designed):

med student = 2

$$R1 = 6$$

 $R2 = 6$
 $R3 = 4$
 $R4 = 2$
 $R5 = 2$
 $R6 = 0$
 $ALL = 4$

Finally, would further realism enhance training on the common tasks? All 10 respondents indicated "yes", as we would expect.

Proctor Observations

A number of significant insights can be gleaned from subject's open-ended comments, informal observations by the proctor and problems experienced during trials. Following are some observations and recommendations derived from the proctors' experiences.

Haptics

Subjects (most notably the residents and ENT staff) had difficulty with the need to steady their thumb to control the instrumented forceps during injection and dissection. This stabilization while opening and closing the forceps was required to keep the tip of the virtual tool from moving away from the area of interaction. This need for stabilization has been attributed to a lack of realistic haptic feedback.

While the version of the haptics subsytem tested here was somewhat rough, Version 1.3 of the software (not yet formally evaluated) has made significant improvements to the haptics, also providing a much more realistic representation of grasping,

tearing and injecting the virtual tissue. Grasping of the virtual tissue is represented by a "hold" on the tip of the instrumented forceps, after closing the jaw on the tissue, as if holding onto a static object. Tearing of virtual tissue is represented by a resistive force on the tip of the instrumented forceps as it is being pulled away from the "hold" position, with a final "release" after a predetermined distance. Injecting is represented by a "pop" when initially passing the virtual needle through the tissue and a "hold", keeping the instrumented forceps static in the X,Y,Z position, but not the heading, pitch and roll. In addition to these enhancements for stabilization, the suction tool is "pulled" toward the virtual tissue based on its proximity, an initial "jolt" is placed on all tools when initially interacting with the virtual tissue and the feel of the sickle knife simulates cutting paper with a straight razor. These improvements have added a tremendous amount of realism, not only to the injection and dissection tasks, but also to the realism of navigating the instrumented forceps through the sinus cavity. These improvements will be implemented and tested in the next phase of the simulator's development.

Retraction of instrumented forceps

During pilot studies, when subjects wanted to swap tools they were required to retract the instrumented forceps from the sinus cavity (retracting to just posterior to the Columella Nasi) before acquiring the new virtual instrument. This requirement was the major contributor to two problems with the encoders on the instrumented forceps. The first problem took place during the pilot studies; the retraction of the instrumented forceps caused the endoscope shaft to collide with the encoder measuring the roll of the instrumented forceps, which would then cause either fraying or displacement of the cable on the sector of the encoder.

The second problem was interaction of the sector of the encoder and the hard palate of the mannequin during retraction. This interaction forced rotation of the sector, introducing an offset in the initial calibration of the instrumented forceps and therefore an offset in the vector of the virtual tool. The problem was repaired by application of LocTite to the sector and eliminating the requirement that the subjects retract the forceps during instrument swapping. Immersion Corporation is currently upgrading and improving the design of the instrumented forceps encoders for the next phase of the evaluation.

Grabbing bone fragments with jawed tools

Removal of two bone fragments from the Uncinate process and three bone fragments from the Bulla ethmoidalis, which were required for a complete score, was unrealistically, due to the

difficulty in grasping with the jaw of the virtual tool; the fragments could only be grasped at their center. Regardless of skill, subjects were required to learn how to grasp the bone fragments in the simulator and required instruction by the proctor on how to do so. Many of the staff and resident ENT surgeons would initially try to grasp the bone fragments on their edge, as they would in a real surgical environement, but without success. Version 1.3 has corrected this problem by allowing a jawed virtual tool to grasp the bone fragments anywhere along its circumference as well as at its center.

Improvement of the virtual suction tool

During testing, the suction tool's only function was to reduce the volume of the blood spheres inside the anatomy after dissection of part of the tissue. It was brought to our attention that in a surgical environment the ENT surgeon would use the real suction tool not only to remove blood, but also to remove small amounts of tissue and mucous in the Posterior Ethmoid cells, and bone fragments throughout. Version 1.3 allows a small amount of dissection by the virtual suction tool, as well as the ability to grasp and slowly remove bone fragments.

Need to break up Model 1 into separate tasks for the non-MD group

During pilot testing of the non-MD group it was discovered that an initial verbal summary of the task to be performed, without giving a visual example of the task, was overwhelming to the untrained/unfamiliar subject. The need for the subject to memorize each subtask and also to gain an understanding of what was needed to complete the task proved to be too demanding. The original presentation of the material consisted of a videotaped introduction to the simulator (described below) followed by verbal instructions by the proctor, describing each subtask (Navigation, Injection and Dissection) in order and in totality.

The subject was then introduced to the endoscope and instrumented forceps and instructed in their functions for each task. The subject was then asked if there was anything which needed further explanation. We found that the proctoring instructions needed to be repeated multiple times throughout the trial to reinforce what the task entailed. By trial and error, it was finally decided to split each subtask into three independent tasks for their initial trial on the simulator. This breakdown was required for only their initial trial.

Ability to "push through" the virtual anatomy during trials

While performing the three tasks subjects had the ability to push through the virtual anatomy with the shaft of both the virtual endoscope and the virtual tool. The staff ENTs expressed

greatest concern with the virtual *tool* passing through the virtual anatomical structures during Injection and Dissection. A potential, but expensive and cumbersome, solution to this problem would be implementation of haptics with 6 degrees of freedom on the instrumented tool. This solution would be too cumbersome to implement in the current design of the mannequin box, but should be considered for future designs.

The unrealistic ability of the virtual *endoscope* to pass through the anatomy, however, was welcomed by the majority of the staff ENTs. When the virtual scope passes through the anatomy, the image on the monitor currently disappears and the screen turns black until the position of the instrumented endoscope is maneuvered back inside the anatomical model. This solution was seen as helpful because the residents were taught to concentrate on staying within the anatomy for the duration of the procedure without potentially traumatizing real tissue in the operating room. This was seen as one of the many advantages of using the simulator to gain the hand-eye coordination necessary to maneuver the endoscope through the anatomy.

Ability to see Maxillary Ostium for Antrostomy

Although the subjects were told to perform a Maxillary Antrostomy, the antrostomy of the Maxillary Ostium was removed from the requirements during the procedure on the simulator because of the inability to realistically view the Ostium. After dissection of the Uncinate process in an actual procedure, the Maxillary Ostium would be in view with a zero or 30 degree scope and the antrostomy would easily be conducted. After dissecting the Uncinate process in the simulation, however, to view the Ostium required a 70 degree scope, with unrealistic positioning of the scope inside the sinus cavity. To correct this problem, Version 1.3 allows dissection of the Uncinate process more superior/laterally and inferior/laterally than Version 1.2. This added realism will be tested and evaluated in the next phase of the evaluation of the simulator.

Instructional video

An instructional video was made to introduce the subjects to the simulator in order to standardize instructions across subjects. The video was designed to instruct them on the procedure, the anatomical structures inside the sinus cavity, and all available tools at their disposal.

The content structure of the original videotape was as follows:

- Endoscope and Forceps
 - Introduction to the instrumented endoscope
 - How to handle it
 - The need and process for stabilization using both hands
 - Hints on guiding the scope, by use of angles, through the anatomy

• How to eliminate the effects of your natural tremor

• Introduction to transferring the tool across the Columella Nasi

• Instructions on cleaning the scope when it becomes "bloody"

- Hints on alignment of the scope within the plane of the mannequin to keep track of your heading and orientation
- The scope's ability to rotate the image, by axially rotating the shaft of the scope
- Explanation of the 30 and 70 degree scopes and their uses to view around corners
- Introduction to the instrumented forceps

• Limitations of the subject's ability to fully retract the forceps

• Rationale for passing the forceps across the Columella Nasi

- Their rigid connection to the mechanics of the haptics inside the mannequin
- How to swap tools in models 2 and 3
 - Calling out the desired virtual instrument
 - Retracting the forceps to just posterior to the Columella Nasi to receive the desired virtual instrument
- Body Positioning
 - Keeping your body parallel with the mannequin
 - Turning head to see the monitor instead of turning body
- Anatomy
 - Introduction to major anatomical structures on physical "pull away" model
 - Nasal Passage, Septum and Nasopharynx
 - Introduction to sinus cavity anatomical structures on physical "pull away" model
 - Superior, Middle and Inferior Turbinates, Uncinate Process and Bulla ethmoidalis
 - Description of anatomical structures which will be interacted with and removed during procedure
 - Definition and procedure for a Maxillary Antrostomy
 - Description and demonstration of sites for injection during procedure in models 2 and 3
 - Medial Middle Turbinate, root of Uncinate Process and the lateral Nasal wall
 - Description and "pull away" of Dissection tasks for models
 2 and 3
 - Medialization of Middle Turbinate and dissection of Uncinate Process, Bulla ethmoidalis and widening of the Maxillary Ostium
- Procedures
 - Playback of video of Novice model, performed by Dr. Edmond, no voice over to allow the proctor to discuss the trial with the subject
 - Examples of all virtual instruments available to the subject
 - Playback of video of Intermediate model, performed by Dr.

Edmond, no voice over to allow the proctor to discuss the trial with the subject

 Playback of video of Model 3, performed by Dr. Edmond, no voice over to allow the proctor to discuss the trial with the subject

The video proved to be too advanced for the non-MD group, presenting an excessive amount of information for the untrained or unfamiliar subject. Pilot subjects from this group who were shown the video reported being confused as to what was required of them. Specifically, these subjects did not

have enough enough time to comprehend both the simulator interface and the purpose of the procedure being simulated.

Future work is planned in this area to create group-specific videos for each subject group. Shortening the duration of the instructional video will make the testing protocol more efficient, and automating more of the proctoring instructions will permit a single proctor to complete all necessary tasks without difficulty.

Conclusions and Recommendations

Procedural validity

The validity of the simulator for the ESS domain is suggested by a number findings:

- ENT subjects performed better than non-MD subjects on both the novice (abstract) and intermediate (anatomical with aids) models
- initial performance on the novice model was correlated with residency level and degree of prior ESS experience
- patterns of difficulty for asymptotic performance on the simulator seem to match the typical pattern of subtask difficulty in the OR
- subject ratings of the realism of the virtual anatomical model were consistently high on the post-session questionnaire and in open-ended comments
- post-training questionnaire responses confirm that the simulator was generally perceived as valid and useful for ESS training by the ENT subjects

Curriculum design

The ESS simulator takes medical simulation several major steps forward in its evolution. Aside from its technical accomplishments, the integration of a well-thought-out curricular framework allows it to take advantage of virtual reality without sacrificing the benefits of more traditional computer-aided instruction.

Model 1 introduces the student into an abstract environment allowing the student to gain the required hand-eye coordination with the endoscope and the special skills needed to maneuver the instrumented forceps, without requiring them to concentrate on anatomy.

Model 2 introduces the student to the anatomy, but still utilizes the training aids from Model 1. This model gives the student the help of hoops for the initial passes through the anatomy, targets for injection areas and labels on the anatomical structures with which interaction is necessary. The educational advantages of simulation can best be achieved with a model of this kind.

Model 3 introduces the student to a more realistic environment. There are no longer any training aids to guide the student through the procedure. For Navigation of the scope, the student must rely on what was learned when navigating through the hoops in Model 2. For Injection the student must remember where injection of the vasoconstrictor is useful. For dissection, the student has no labels to indicate what anatomy to interact with, and so must rely on what was learned in Model 2 to perform the procedure.

Future Directions for Evaluation

Optimal Instrument/Scope Path Analysis

[Edward's figures...quantify as mean square deviation from lowess regression path for navigation and mean square deviation from path centroids for injection and dissection?]

Evaluation of Improved Haptics

The value of the haptic feedback subsystem to the simulator was assessed only subjectively in the current phase. In general, it was perceived to be only moderately useful, and we suspect that the relatively poor performance on the dissection task, in particular, may ahve been due largely to inadequate feedback about the tissue forces.

As described above, preliminary evaluation of Version 1.3 suggests that the haptic subsystem is greatly improved. The improved system will allow us to evaluate more formally its impact on training trial performance and its contribution to OR preformance. In addition, we will investigate more formally the relative merits of force feedback on the endoscope versus the intrument.

Transfer of Training

Transfer of training to the real operative environment is the primary objective of this procedural simulator, and will be the primary focus of the next phase of our research.

In preparation for that phase, we have collected initial OR videotape data from several first-time ESS procedures by residents for whom we have previous simulator performance data. These tapes will be formally analyzed for subtask proficiency, as rated by their attending staff instructors, to derive a set of systematic measures of performance on the target task. These will then be correlated with prior simulator experience for current and entering residents to determine the training effectiveness of the simulator and to provide guidance for its further development.

Model Enhancements

As noted earlier, the development team elected to focus the first iteration of the ESS simulator (appropriately) on the needs of junior residents. Several enhancements to the library of available anatomical models are currently being integrated which are intended to extend the utility of the system to senior residents and more experienced staff physicians.

Appropriate enhancement implementation will require iterative formative evaluation, as before, and the training utility of each enhancement will need to be assessed. Transfer of training may need to be assessed indirectly in some cases, since real-world incidence of certain conditions of interest may be infrequent under normal conditions.

Criterion Performance Standards

Ideally, subject performance on the simulator will be reliably predictive of OR performance. Achievement of this correpondence will enable us to better establish performance criteria for advancement through the protocol. In a sense, the scoring algorithm represents a theoretical model of surgical proficiency; additional validation studies of the simulator will help to correct and refine those theoretical models.

In addition to examining the learning curve for simulator performance over time, the distribution of trials over time is also

of interest. For our "steady-state" proctors, for example, the temporal distribution of training trials was highly variable, ranging from several trials per day to one trial every 2-3 weeks. Routinely collecting repeated measurements on resident subjects will enable us to explore the optimal number and timing of training trials in the resident's curriculum. Undoubtedly some degree of trial spacing (as opposed to massed practice) will prove most effective, as has been shown in many other domains; of interest is just what that spacing strategy should be for ESS procedure training. Ultimately, we may also be able to establish an equivalence between time on the simulator and time in the OR. Such a correspondence would be useful for residency curriculum development, and perhaps eventually for professional credentialling. Such a system would be of significant benefit for patients, and could greatly extend accessibility to procedural training opportunities.

Summary

This study presents the results of our systematic evaluation of a high-end virtual reality simulator aimed at training a set of skills essential to endoscopic sinus surgery. Our findings suggest that the simulator represents a valid and useful implementation of the target ESS tasks. In addition, the thoughtful integration of an organized curriculum perspective makes this system uniquely valuable among emerging medical simulation systems.

This study also suggests a framework for incorporating systematic evaluation into the process of developing procedural training simulators in the medical domain. Incorporating both formative and summative aspects of evaluation has greatly enhanced the development process and assures the continuing evolution of a usable and effective system.

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Appendix A. Surgical Needs Survey

- **Appendix B.** Trial Matrix Form
- **Appendix C. Pre-Session Survey**
- **Appendix D. Post-Session Survey**

Appendix E. Summary of Questionnaire Comments