WEB PAPER

High-fidelity simulation is superior to case-based discussion in teaching the management of shock

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Abstract

Background: Case-based discussion (CBD) is an established method for active learning in medical education. High-fidelity simulation has emerged as an important new educational technology. There is limited data from direct comparisons of these modalities.

Aims: The primary purpose of this study was to compare the effectiveness of high-fidelity medical simulation with CBD in an undergraduate medical curriculum for shock.

Methods: The subjects were 85 third-year medical students in their required surgery rotation. Scheduling circumstances created two equal groups. One group managed a case of septic shock in simulation and discussed a case of cardiogenic shock, the other group discussed septic shock and experienced cardiogenic shock through simulation. Student comprehension of the assessment and management of shock was then evaluated by oral examination (OE).

Results: Examination scores were superior in all comparisons for the type of shock experienced through simulation. This was true regardless of the shock type. Scores associated with patient evaluation and invasive monitoring, however, showed no difference between groups or in crossover comparison.

Conclusions: In this study, students demonstrated better understanding of shock following simulation than after CBD. The secondary finding was the effectiveness of an OE with just-in-time deployment in curriculum assessment.

Introduction

High-fidelity simulation is coming of age in medical education and now requires critical evaluation. Some 40 years after early implementation in anesthesiology, there has been an explosive growth of medical simulation across specialties, disciplines, and missions. The computer-based, model-driven, full-scale simulator is a technology at the forefront of this movement. This tool has matured precisely at a time when medical educators seek methods of active learning, competency demonstration, and individualized education within constantly evolving curricula. The purported advantages of simulation (SIM) are myriad and include learner-centered educational experiences in an environment free of risk to patients, repeated and controlled exposure to clinically rare events, individualization of learner experiences with standardization of competencies, and team development opportunities. The success of broad and long-standing applications in aviation, the military, and industry are cited in further support of SIM.

Developments thus far are similar to the evolution of other technologies in medical education and clinical practice. The first phase involved adaptation with the development of SIM devices, task training, and crisis resource management by anesthesiologists, often translating practices from aviation (Howard 1992). Next, surveys of learners demonstrated the positive impact of realistic practice on the confidence and attitude toward both routine and rare, high-stakes events. Other specialties and disciplines are now voraciously adopting SIM at essentially all levels of education, training, and practice (Grenvik 2004; de Leng 2006; Gordon 2006; Shukla 2007; Fraser 2009; Fernandez 2010; Schout 2010). The current era is one in which simulation is being evaluated critically, sometimes in comparison with other educational methods, for educational outcomes, learner competencies, and clinical outcomes (McGaghie 2010; Cook 2011). With such investigation, this resource-intensive modality can be used with evidence-based prioritization and cost-effectiveness. In an

Practice points

- Students demonstrated superior understanding of shock after a simulation experience than CBD, regardless of the type of shock.
- An OE, despite just-in-time deployment and minimal faculty development, demonstrated good performance in discriminating group performances after different curricular experiences.
- The OE demonstrated good correlation with learner level, supporting its construct validity.

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effort to continue the critical analysis of simulation education, we report data comparing the comprehension of shock demonstrated by medical students following a SIM experience to that demonstrated after case-based discussion (CBD). Such comparison is particularly relevant now, in an era of frequent curricular reform and widespread simulation implementation.

**Methods**

All described activities occurred during one day of our third-year medical student (MS3) surgical clerkship rotations. The day started with a presentation of the basic presentation, evaluation, and hemodynamic monitoring of shock. Students were then divided alphabetically by last name into two equal groups and attended an airway workshop, a CBD, and a SIM session. The schedule of the student groups is detailed in Table 1, wherein students who attended SIM for cardiogenic session. The schedule of the student groups is detailed in Table 1, wherein students who attended SIM for cardiogenic shock and CBD for septic shock are designated the SIM group, while students undergoing the contraposed assignments are designated SIM

The SIM and CBD sessions were based upon the same cases. The cardiogenic case presented with previous cardiac history, nausea and dyspnea following a complex cholecystectomy. The sepsis case was that of an elderly man with confusion following admission for urinary obstruction. The formats of the two sessions were necessarily different. CBD sessions began with the case presentation. This was followed by group development of (1) differential diagnoses, (2) patient assessment and diagnostic strategy, and (3) both immediate and intermediate management plans. The format of this portion of the experience was that of a problem-based learning discussion (PBLD). As opposed to PBLD, the students did not receive preparation materials, including patient presentation, before the activities. Additionally, the faculty facilitator did complete the session with the review of key elements of shock.

For the SIM experience, the student team was given a brief patient history by their busy supervising house officer. They were then sent to evaluate the patient on the ward with symptoms that had alarmed the floor nurse. At the patient’s bedside, the students were presented initially with non-specific signs and symptoms in a patient whose condition deteriorated over about 10–15 minutes and first required stabilization and then transfer to a critical care unit. After alerting the supervising housestaff and during patient transfer, the students briefly re-conferenced to discuss management. A faculty member facilitated discussion of the group’s working diagnoses and strategy. The group then attended to the patient in the critical care setting during which they further developed and tested their differential diagnoses, implemented immediate management, and planned intermediate management strategies. A final debriefing was then conducted, focusing on key elements of shock recognition, stabilization, assessment, and treatment. Team function was sometimes addressed, but not emphasized. The simulation scenarios were conceived by faculty and implemented by simulation specialists and educators. High-fidelity simulators (ECS simulator, formerly METI®, now CAE® Healthcare, Sarasota, Florida, USA) were utilized in mock ward and ICU bays.

**Data collection**

After completion of all sessions, students underwent an oral examination (OE). The OE was implemented as an attempt to familiarize our students with this type of examination and for program evaluation. Most MS3’s had no experience with OE, yet their final surgery clerkship evaluation would include an OE at the end of the rotation six weeks later. Those students going on to surgery or anesthesiology would eventually undergo specialty OE’s following residency training. As a program evaluation tool, we specifically sought to determine if students could explain key elements of patient evaluation in crises, invasive monitoring, and the pathophysiology and management of shock following educational activities designed for these issues. The students were assured that the OE was “no-stakes” in terms of individual grading for the clerkship.

The OE was based on a written case presented to the student 5–10 minutes before the examination. The presentation (Appendix) included distinct possibilities of hypovolemia, ischemic cardiac dysfunction, and systemic inflammatory response. Anesthesiology faculty and senior residents administered the OE over about 30 minutes. The examiner’s introduction to the scoring instrument (Appendix) was minimal by intent. Examiners were told that a score of 3 represented MS3 level-appropriate understanding or decision-making. Scores of 2 or 1 represented progressively
inferior performance. The expected performance of a new medical school graduate would be scored as 4, while any higher-level performance was scored as 5. A few questions had specific scoring guidelines that superseded this schema. The examiners were asked to complete as many of the questions as possible except for hypovolemic shock. Examiners then reviewed the examination with the student with the discussion of relative strengths and weaknesses as well as key concept reinforcement.

The results of the examination were separated into four topics for analysis. The first six questions regarded initial patient evaluation, prioritization, diagnostics, and differential diagnoses. These questions were termed the evaluation (EVAL) section, as marked in the Appendix. The next five questions, marked as MON, compared the monitoring modalities of central venous pressure and saturation, pulmonary artery catheter and mixed venous saturation, and echocardiography. Four questions each were then devoted to septic shock (SEP) and cardiogenic shock (CRD).

Data treatment
The results from all examinations were entered into statistical software (SPSS 16.0, SPSS of IBM Company, Chicago, IL). Data were entered into a duplicate file and inconsistencies resolved by the review of original scoring sheets. These data were used for graphic representations of group performance during curricula evaluation. Institutional Review Board exemption was granted for the retrospective in-depth analysis of de-identified data. An administrative assistant stripped identifying data and the de-identified and randomly reordered data was provided to author KEL for analysis.

Student scores were calculated both as mean topic scores and as indexed topic scores. The arithmetic mean of all non-missing scores except those for hypovolemia were calculated to determine each student’s ALLavg scores. The scores from each separate section described above were averaged to generate EVALavg, MONavg, SEPavg, and CRDavg. Individually indexed scores were calculated by dividing a student’s mean score for each module by the arithmetic mean of all students for that section to respectively generate EVALi, MONi, SEPi, and CRDi. All calculated values were evaluated by the Kolmogorov–Smirnov (K-S) test. Results indicated that all values were normally distributed, and parametric methods were thus utilized for data analysis.

The data was first analyzed by student’s test to compare individual raw and indexed scores between groups, as summarized in Table 2. Since the circumstances of scheduling provided a crossover assignment pattern, the data was then analyzed to compare the indexed scores of students by paired t-test (Table 3). This analysis was performed for all students, and separately for the two different assignment groups (SIMcardiac versus SIMsepsis), blinded versus non-blinded examiners, and SIM-first versus CBD-first groups, as will be discussed below. The variable names CBD, and SIMi were used to denote the index for the type of shock experienced by

<table>
<thead>
<tr>
<th>Parameter and group</th>
<th>n</th>
<th>Minimum</th>
<th>Maximum</th>
<th>Mean</th>
<th>S D</th>
<th>Difference between groups: r value (95% CI)</th>
</tr>
</thead>
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<tr>
<td>Raw scores</td>
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<td>3.72</td>
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<td>3.73</td>
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<tr>
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<td>4.50</td>
<td>3.48</td>
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<td>3.95</td>
<td>0.64</td>
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<td>CRDavg</td>
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<td>3.58</td>
<td>0.08</td>
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<td>3.88</td>
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<td></td>
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<td>1.00</td>
<td>0.19</td>
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<td>1.01</td>
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<tr>
<td>MONi</td>
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<td>1.39</td>
<td>1.00</td>
<td>0.17</td>
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<td>0.99</td>
<td>0.20</td>
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<td>1.39</td>
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<td>0.19</td>
<td></td>
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<tr>
<td>SEPi</td>
<td>85</td>
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<td>1.35</td>
<td>1.00</td>
<td>0.19</td>
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<td>1.21</td>
<td>0.93</td>
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<tr>
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<td>0.74</td>
<td>1.35</td>
<td>1.06</td>
<td>0.17</td>
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<tr>
<td>CRDi</td>
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<td>0.49</td>
<td>1.40</td>
<td>1.00</td>
<td>0.19</td>
<td></td>
</tr>
<tr>
<td>SIMcardiac</td>
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<td>1.09</td>
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<td></td>
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<td>0.49</td>
<td>1.33</td>
<td>0.92</td>
<td>0.17</td>
<td></td>
</tr>
</tbody>
</table>

Notes: SIM = Simulation, CRD = cardiogenic shock, and SEP = septic shock.
a student through CBD and SIM, respectively. While this analysis was performed for all pair permutations for completeness, Table 3 includes only the EVAL\textsubscript{i}\textendash MON\textsubscript{i}, and SIM/CBD\textsubscript{i} pairings. This is consistent with (1) the expectation that scores for evaluation and monitoring would be similar since they were taught in lecture format and (2) the null hypothesis that there should be no difference between SIM, and CBD, if the sessions were equally effective. (There were, incidentally, no findings of statistical differences for any pairs not shown in Table 3.)

**Results**

Figure 1 presents a graphical representation of SIM\textsubscript{i} versus CBD\textsubscript{i}. A summary of the parameters calculated as described above is presented in Table 2. The results of analysis are summarized in Table 3. No difference was demonstrated between EVAL\textsubscript{i} and MON\textsubscript{i}, in any comparison. In all comparisons, however, of SIM\textsubscript{i} and CBD\textsubscript{i} were statistically different.

**Impact of study design**

There are critical issues of study design that must be considered in the interpretation of our results, and the impact of which we attempted to evaluate as described below.

Conflict of SIM and CBD experiences. A possible confounding factor is that three-fourths of the students experienced CBD before their SIM. This raises the concern that superior performance of the second experience, usually SIM, actually reflected positive impact from the first experience, usually CBD. The second experience could be more productive because of any number of issues such as priming, activation of prior knowledge, cross-knowledge, or situational

![Image](image.png)

**Table 3.** Paired sample t-test of all subjects and various sub-groups as noted.

<table>
<thead>
<tr>
<th>Groups analyzed and number of subjects</th>
<th>Mean</th>
<th>SD</th>
<th>SE Mean</th>
<th>Lower</th>
<th>Upper</th>
<th>t</th>
<th>df</th>
<th>Sig. (2-tailed)</th>
</tr>
</thead>
<tbody>
<tr>
<td>All Students (85)</td>
<td>EVAL\textsubscript{i}\textendash MON\textsubscript{i}</td>
<td>0.0001</td>
<td>0.1503</td>
<td>0.0163</td>
<td>−0.0324</td>
<td>0.0324</td>
<td>0.0000</td>
<td>84</td>
</tr>
<tr>
<td>SIM Cardiac (42)</td>
<td>EVAL\textsubscript{i}\textendash MON\textsubscript{i}</td>
<td>−0.0014</td>
<td>0.1539</td>
<td>0.0237</td>
<td>−0.0494</td>
<td>0.0465</td>
<td>−0.0014</td>
<td>41</td>
</tr>
<tr>
<td>SIM Septic (43)</td>
<td>SIM\textsubscript{i}</td>
<td>0.1519</td>
<td>0.1251</td>
<td>0.0193</td>
<td>0.1129</td>
<td>0.1909</td>
<td>0.1519</td>
<td>41</td>
</tr>
<tr>
<td>SIM First (21)</td>
<td>EVAL\textsubscript{i}\textendash MON\textsubscript{i}</td>
<td>0.1456</td>
<td>0.1457</td>
<td>0.0222</td>
<td>0.1007</td>
<td>0.1904</td>
<td>0.1456</td>
<td>42</td>
</tr>
<tr>
<td>CBD First (64)</td>
<td>EVAL\textsubscript{i}\textendash MON\textsubscript{i}</td>
<td>0.0037</td>
<td>0.1605</td>
<td>0.0201</td>
<td>−0.0363</td>
<td>0.0438</td>
<td>0.0037</td>
<td>63</td>
</tr>
<tr>
<td>Non-Blinded (74)</td>
<td>EVAL\textsubscript{i}\textendash MON\textsubscript{i}</td>
<td>−0.0038</td>
<td>0.1575</td>
<td>0.0183</td>
<td>−0.0403</td>
<td>0.0277</td>
<td>−0.0038</td>
<td>73</td>
</tr>
<tr>
<td>Blinded (11)</td>
<td>SIM\textsubscript{i}\textendash CBD\textsubscript{i}</td>
<td>0.1221</td>
<td>0.1242</td>
<td>0.0144</td>
<td>0.0973</td>
<td>0.1549</td>
<td>0.1261</td>
<td>73</td>
</tr>
</tbody>
</table>

OE instrument. We used a new OE instrument without prior validation. This demands analysis of the instrument’s performance. Face and content validity are reflected in the examination design. The primary purposes of the OE were to prepare students for later high-stakes examination(s) and also for program evaluation. The questions were derived from educational goals and objectives for this medical student experience. The tool was thus not adapted from some prior use but created de novo for this experience.

Construct validity can be approached more quantitatively. Authors KEL and CJS began using the OE as part of their educational activities with both senior medical students and housestaff during the study period. Senior medical students and junior off-service residents were examined during rotations on the anesthesiology service without the benefit of
focused educational experiences immediately preceding the OE. As shown in Figure 2, there is a strong association between the learner level and mean raw score. Pearson's correlation coefficient was found to be 0.71 for ALLavg, 0.65 for EVALavg, 0.61 for MONavg, 0.54 for SEPavg, and 0.66 for CRDavg. These findings were statistically significant with \( p < 0.0001 \) for all parameters.

Discussion

The chief finding of this study is that students demonstrated better understanding of shock following a SIM experience as compared to a CBD experience. The secondary finding is that a new OE tool with just-in-time implementation and without faculty development provided meaningful information for program evaluation. There are both limitations and advantages incurred by our study design. The study is retrospective and there was no randomization of subjects. Conversely, the results reflect normal behavior of faculty and students in small group sessions without self-selection for study participation or likelihood of variously described Hawthorne effects (Holden 2001).

Our data includes results from non-blinded examiners. For this reason, blinded and non-blinded examiner results were separated and analyzed as described above. Blinded examiners found the same pattern of statistically significant differences between SIM and CBD, and non-differences between EVAL and MON, as did the non-blinded examiners. The effect size within the blinded group was much larger, as might be logically surmised from the smaller group size. These findings indicate that non-blinded examiners, at the very least, did not demonstrate bias towards SIM when compared to the patterns of blinded examiners.

Similarly, statistically superior performance was associated with the SIM experience regardless of the order in which the two different methods were experienced, although the difference was less when SIM preceded CBD.

By convention, educational encounters with an effect size of 0.2, 0.5, and 0.8 are considered to respectively have had small, moderate, and large impact in qualitative terms (Colliver 2000). Despite a long-standing and central role in the active learning movement in medical education, PBLD has typically been found to have a small or moderate effect size (Hartling 2010). The effect size of SIM compared to CBD was large in our data (Cohen's \( d \) was 0.68 for septic shock and 0.89 for cardiogenic shock). Because the actual implementation of PBLD in medical education is as variable today as when these inconsistencies were discussed a decade ago (Lloyd-Jones 1998; Smits 2002), it is impossible to compare our CBD to a non-existent “standard” PBLD. If the CBD in this study is considered to be a PBLD, then our study shows large differences between SIM and PBLD, a gold standard of learner centered education. Considering the interactive, small-group nature of the CBD, it would be extreme to consider the experience as only a lecture. But even in this most conservative interpretation, the effect size of SIM is still greater than that demonstrated by PBLD in most studies.
These findings corroborate recent work demonstrating improved educational outcomes following SIM as compared to PBLD, also amongst medical students (Steadman 2006). Further, our study complements prior studies with (1) a large number of subjects, (2) a method of assessment different than one of the educational modalities being investigated, and (3) the enhanced statistical power of a crossover analysis. Critical analysis of SIM’s effectiveness, particularly in comparison to CBD, is timely. The current exponential growth in simulation centers will be associated with wider integration of SIM experiences into medical education. Of note, institutional pioneers of the PBLD movement have recently described the integration of SIM into educational programs (McMahon 2005; Neville 2007; Gordon 2010). However, the extensive equipment, space, time, and faculty resources required by SIM require selective integration. It is therefore imperative that curricula design be based on data regarding areas of proven efficacy (and non-efficacy) for SIM, especially when it can be compared to other active-learning methodologies.

The secondary finding is that a simple OE with just-in-time deployment demonstrated adequate performance for program evaluation. The development of such an instrument is also timely. We live in an era of seemingly constant curricular refinement (Hecker 2009; Maccarrick 2009; Patel 2009; Snelgrove 2009) even as calls are being made for fundamental retooling of medical education (Irby 2010; Prislin 2010; Taylor 2010). Educators will need metrics that rapidly and reliably provide the impact of curriculum changes. The OE utilized in this study appeared to meet these requirements as well as provide an experience identified by faculty as being educationally important for our students in its own right. It is important to note that the OE was not developed as a grading tool for individual students and is not recommended as such.

In summary, our data shows that SIM resulted in markedly superior demonstration of understanding of key clinical

Figure 2. Mean scores for overall OE and individual topics for different learner levels. MS = Medical Student, PGY = Post Graduate Year of housestaff. Examinations by authors CJS and KEL.
concepts than did CBD, and that a simple OE provided meaningful data for curricular evaluation.

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Declaration of interest: The authors report no conflicts of interest. The authors alone are responsible for the content and writing of the article.

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References

Appendix

Student (Last, First) | Examiner (Initials if Anesth) | / / Date (m/d/yy)

Instructions: Please read the case below. You will then spend 10–15 minutes discussing both this case and several general concepts that were addressed during today’s activities.

Your resident asks you to evaluate a patient whose nurse has called from the intensive care unit. The 62-year-old patient is POD #1 from emergency surgery for drainage of an abdominal abscess. She had a partial colon resection two weeks ago for cancer and was admitted yesterday with progressive fevers and malaise, workup resulted in radiographic diagnosis of the abscess. Yesterday’s surgery was uneventful, but the patient’s status has deteriorated today with declining blood pressure, elevated heart rate, and poor urine output. Vital signs are reported as BP 92/60, HR 124. Urine output over the last 4 hours has been a total of 100 ml.

Past medical history includes:

- Coronary artery disease – Right coronary bare metal stent placed two years ago for acute ischemia. Good exercise tolerance (> 5 mets) since and negative stress test three months ago.
- Adenocarcinoma of colon – Discovered by colonoscopy and resected as described with negative nodes and biomarkers.
- Outpatient medications – Metoprolol, lisinopril, and baby aspirin.

### Questions and Scoring

1. **[Vignette interpretation]**
   - From what you know so far, does this patient need urgent evaluation?
   - What do you think are likely etiologies of the described abnormalities?
   - (Prompt if needed, score as noted)

   **Scoring:** 1 2 3 4 5
   - 1-2 for lack of urgency, 4-5 for well articulated concerns
   - 1 pt @ for good diffs. +1 for context (ST changes, antibiotics, etc.). -1 prompting

2. **[Evaluation]**
   - Upon arrival in the ICU, how would you further evaluate the patient? Explain how each step helps you to better understand the situation.
   - (Allow up to four steps)
   - If not already mentioned- what monitoring devices might be particularly helpful in assessing and managing this patient?
   - How would you characterize the relative advantages and disadvantages of a pulmonary artery catheter, central venous pressure, and echocardiography in this situation?

   **Scoring:** 1 2 3 4 5
   - HH&P (incl. urine, etc.)
   - Labs with good logic
   - Radiology / EKG
   - Review of Tx/Rx so far
   - Noninvasive monitors
   - Invasive monitoring
   - Understands PAC +/-
   - Understands CVP +/-
   - Understands Echo +/-

3. **[Treatment]**
   - We will consider each of the following possibilities separately, and I would like you to describe good management for each:

   **Sepsis**
   - Demonstrates understanding of:
     - Sepsis pathophysiology
     - Pharmacology of support
     - Tx (cultures, I&D, antibiotics)

   **Cardiogenic Shock**
   - Demonstrates understanding of:
     - CHF pathophysiology
     - Pharmacology of support
     - Tx (inotropy, vasodilatation, revascularization, support devices, etc.)

   **If time allows (and it probably should NOT):**
   - Demonstrates understanding of:
     - Hypovolemic hemodynamics
     - Hypovolemia pathophysiology
     - Pharmacology role discussion
     - Tx (monitored resuscitation)

   **Not Considered**