Emergency Department (ED) Medical Directives Implementation Kit

Wait Times Strategy
Emergency Department (ED) Medical Directives
ED Medical Directives Working Group
Guidelines
Emergency Department (ED) Prototype Medical Directives

Use these guidelines to understand the ED Prototype Medical Directives Implementation Kit and develop directives in your ED
ED Medical Directives Working Group Members

Dr. Howard Ovens,
Director, Schwartz/Reisman Emergency Centre, Mount Sinai Hospital

Kathy Stevenson,
Director Emergency and Critical Care Services, William Osler Health Centre, Brampton and Etobicoke Sites

Wendy Cheung,
Clinical Nurse Educator, Emergency Services Program, North York General Hospital

Dr. David Dushenski,
Quality Assurance Coordinator, Schwartz/Reisman Emergency Care Centre, Mount Sinai Hospital

Dr. Jeff Eisen,
Medical Director, Emergency Services, Royal Victoria Hospital

Carolyn Farquharson,
Emergency Nursing Unit Administrator, Mount Sinai Hospital

Jane Foster,
Educator/Practice Leader, Emergency Department, Grand River Hospital

Susan Harper,
Educator - Emergency Services, Peterborough Regional Health Centre

Joy McCarron,
Clinical Educator, Emergency Department, Southlake Regional Health Centre

Dr. Naveed Mohammed
Corporate Chief Emergency Medicine, William Osler Health Centre

Dr. Tim Rutledge,
Medical Director, Emergency Services Program, North York General Hospital

Roz Smith,
(Former) Special Advisor to the Assistant Deputy Minister (ADM), Acute Services, Ontario Ministry of Health and Long-Term Care

Heather Stewart,
Consultant, Ontario Hospital Association

Paula May Ponesse,
Consultant, Health Care Policy and Practice
Table of Contents

Purpose .................................................................................................................. 3
Contents ............................................................................................................... 3
Steps for Establishing ED Directives ................................................................. 5
Glossary of Terms ............................................................................................... 6
Frequently Asked Questions (FAQS)
   List of FAQs  .................................................................................................. 8
   FAQs for Physicians (Of Interest to the Team) ............................................. 10
   FAQs for the Entire Team ............................................................................. 14
Appendix A - Sample Policy Considerations for Patients Who Leave Without
Being Seen (LWBS) .......................................................................................... 20

Modules (M)
   • M-1 - Performance Readiness Assessment .............................................. M-1
   • M-2 - Performance Readiness Plan ......................................................... M-2
   • M-3 – Prototype Directives ................................................................. M-3
   • M-4 – Sample Prototype Directive Pre-printed Orders and ED Record M-4
   • M-5 – CEDIS Presenting Complaints Table ........................................... M-5
   • M-6 – Sample Approval Forms ............................................................ M-6

Acknowledgements

The modules contain information and templates from the following sources:


Readers may refer to these resources or other College publications for further information.

The prototype medical directives draw on directives developed by the:

- Child Health Network for the Greater Toronto Area, Toronto
- Grand River Hospital, Kitchener
- Mount Sinai Hospital, Toronto
- North York General Hospital, North York
- Peterborough Regional Health Centre, Peterborough
- Royal Victoria Hospital, Barrie
- Southlake Regional Health Centre, Newmarket
- William Osler Health Centre, Brampton and Etobicoke

This template has been adapted from the Emergency Department Medical Directives Implementation Kit
www.oha.com/edmedicaldirectives
Purpose

This implementation kit has been developed for use by Emergency Department (ED) physicians and staff in response to a recommendation “to maximize the use of medical directives within the emergency department, in order to increase efficiency of patient flow and patient care” made by the OHA/OMA/MOHLTC Physician Hospital Care Committee in its report *Improving Access to Emergency Care: Addressing System Issues* (August 2006), p.10.

Developed by a provincial Expert Panel of ED physicians and clinical staff, the implementation kit provides consensus-based information and tools drawn from current ED practices and directives along with guidelines and templates for establishing medical directives by the College of Physicians and Surgeons of Ontario (CPSO), the College of Nurses of Ontario (CNO), the Federation of Health Regulatory Colleges of Ontario (FHRCO) and other regulatory colleges, thus fulfilling regulatory and legislative expectations for practice.

What the implementation kit and directives are not:

- They are not intended to be a standard of practice or required set of directives, but rather a resource to be used as ED physicians and staff see fit. Modules and content may be adapted, adopted, or referred to in order to augment already existing directives.
- Directives are not a substitute for timely assessment of patients by physicians. Patient-specific orders provided by a physician who has seen and assumed care of a patient remains the gold standard for care. Proper directives authorize designated staff to carry out orders and expedite care before a physician is able to see a patient, but only when it is safe, effective and ethical to do so.
- Properly developed and applied directives should not increase unnecessary tests and procedures administered to patients. Instead, they optimize care by mandating physicians and teams to come to a consensus on practice issues and develop a consistent approach. In addition, time for key tests and interventions can be reduced, improving patient and staff satisfaction.

Contents

The implementation kit has been designed to enable ED physicians and staff to fulfill the following steps necessary for establishing directives:

1. Assure performance readiness (competence) of team members given the specific circumstances in each ED;
2. Assure a medical directive is written as a proper order and suits setting-specific circumstances (patients, competencies, practices);
3. Assure team input and agreement, particularly from co-implementers;
4. Assure approval and sign-off by physicians and hospital administrative authorities; and
5. Assure directives are reviewed for appropriateness as indicated.
The implementation kit contains the following modules (M):

**M-1. Performance Readiness Assessment** - to determine the appropriateness of using directives

**M-2. Performance Readiness Plan** - to establish necessary competencies, including:
   a) Educational Resources – information to assure competencies
   b) Quiz - to assess and demonstrate competencies
   c) Implementer Performance Readiness Forms - to record agreement with directive and achievement of competencies (individual and group)

**M-3. Prototype Directives** – sample to facilitate establishment of setting-specific directives

**M-4. Sample Directive Pre-printed Order Sheets and Emergency Record** – sample options for documenting implementation of orders

**M-5. CEDIS Presenting Complaints Table** – a quick reference summarizing directive orders by presenting complaints

**M-6. Approval Forms**
   a) Physician Approval Form – Staff Physicians
   b) Stakeholder/Administrative Approval Form
   c) Physician Approval Form - Locum, Casual Physicians
   d) Physician Approval Form – New Staff (to enable new staff to sign off)

Electronic versions of the modules are available at: [www.oha.com/edmedicaldirectives](http://www.oha.com/edmedicaldirectives)
Steps for Establishing ED Directives

1. Assure Performance Readiness
   Convene a sponsoring group:
   Consisting of at least the Chief Physician and representatives of administrative, implementing and co-implementing staff to complete:
   - M-1 - Performance Readiness Assessment – assess the appropriateness of using the directives,
   - M-2 - Performance Readiness Plan - establish the competencies to support using the directives
     OR
   - Complete the same due diligence steps using existing setting specific methods

2. Develop Proper Directive
   Complete: Medical directive(s) using the:
   - M-3 - Prototype Directives
   - M-4 - Sample Directive Pre-printed Orders and ED Record
   - Use already existing directives and tools

3. Obtain Team Input and Agreement
   Identify and Consult: Relevant stakeholders, particularly co-implementers using the:
   - M-1 - Performance Readiness Assessment (Section 11) to identify stakeholders and guide consultations
     OR
   - Use already existing processes

4. Obtain Approvals
   Assure Sign-off: By physicians and staff using:
   - M-6 - Approval Forms:
     - Physician (Staff, Chief, Locum, New)
     - Administrative
     OR
   - Use already existing processes

5. Renew Directives at Planned Intervals to Ensure They Remain Appropriate and Maintain Appropriate Corporate Records

Physicians and Staff Begin to Practice Using Medical Directives

This template has been adapted from the Emergency Department Medical Directives Implementation Kit
www.oha.com/edmedicaldirectives
## Glossary of Terms\(^1,2\)

<table>
<thead>
<tr>
<th>Interprofessional Term</th>
<th>Definition</th>
</tr>
</thead>
</table>
| **Authorizing Mechanism**                                 | Device to sanction and enable performance of procedures, treatments, and interventions when such sanctioning is required by law, practice convention or circumstances including:  
  • Controlled acts or restricted procedures authorized to a profession  
  • Orders - direct orders and directives - and delegation  
  • Practice setting mechanisms (e.g. privileges, assignments, role descriptions)                                                                                                                                                                                                 |
| **Order – Direct Order or Medical Directive**             | Direction given by a regulated health professional with legislative ordering authority to permit performance of a procedure. Orders are required by legislation (e.g. the Public Hospitals Act (PHA), 1990; Regulated Health Professionals Act (RHPA), 1991; Healing Arts Radiation and Protection (HARP) Act, 1990 and Laboratory and Specimen Collection Centre Licensing Act (LSCCLA), 1990), and must be provided by the profession authorized to order. There are two types of orders:  
  • *Direct Order* – order given by a physician or authorizer upon assessment of the patient at the time.  
  • *Medical Directive* – advance, written order given by a physician or authorizer prior to his or her direct assessment of the patient that identifies the specific conditions when the order can be implemented, and who is authorized to implement it.  
  Directives may authorize an *implementer* to implement or activate the order and a *co-implementer* to carry it out. For example, a physician’s order for labwork may be implemented by a nurse and carried out by medical laboratory technologist. In this scenario, the nurse is not ordering the labwork, but is implementing the physician’s order, permitting the medical laboratory technologist to carry out the testing in compliance with their legislative requirements for a physician’s order. |
| **Authorizer**                                            | A regulated health professional authorized by legislation to give orders or prescriptions, permitting others to perform procedures or administer diagnostics and therapeutics in accordance with their legislative requirements for orders. In a hospital, those authorized to give orders and prescriptions are physicians, dentists, midwives, and for outpatients, RN(EC)s. |

---


\(^2\) The prototype ED directives do not involve delegation in the RHPA sense, thus delegation is not mentioned in this glossary. For information regarding delegation, consult the references identified in Footnote 1 above, particularly the FHRCO Guide for an interprofessional perspective.
<table>
<thead>
<tr>
<th>Interprofessional Term</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Implementer</strong></td>
<td>Someone authorized to implement and perform procedures or administer diagnostics and therapeutics ordered by a physician, or another health professional authorized to order.</td>
</tr>
</tbody>
</table>
| **Co-implementer**     | Someone authorized to perform procedures/administer diagnostics and therapeutics under physician orders implemented by another, subject to the co-implementer’s:  
  - Understanding that s/he is a designated co-implimentor, and  
  - Agreement with the order at the time.  
  For an example involving co-implementers, see the Medical Directive description above. |
| **Performance Readiness** | Refers to the competencies – i.e. the demonstrated knowledge, skill and judgment – of practitioners to safely, effectively and ethically authorize or perform procedures and manage outcomes given the circumstances in a situation. It is determined by both individual competence and by the conditions within a practice setting that influence ability to practice competently. Assuring performance readiness of both practitioners and settings is a necessary precondition to authorizing and performing any test or procedure. |
| **Circumstances in a Situation** | Refers to the specific attributes in a practice setting that determine what competencies and conditions are necessary for safe, effective, ethical care that fulfills legislative and regulatory requirements. Circumstances in the situation include:  
  - Patient condition and needs,  
  - Team composition, competencies and care delivery processes,  
  - Supervising physician competencies, supervision availability and practice preferences, and  
  - Care setting characteristics such as the availability of supplies and resources and the capacity to develop and maintain delegation and directives. |
| **Performance Readiness Assessment** | A due diligence process undertaken prior to authorizing or performing any procedure to assure performance readiness and thus the appropriateness of authorizing and performing the procedure. See Module 1, Performance Readiness Assessment. |
| **Performance Readiness Plan** | An education and learning plan for use when a more involved, formal competency acquisition program, and/or recordkeeping is desirable. See Module 3, Performance Readiness Plan. |
Frequently Asked Questions (FAQs)³

FAQs for Physicians (Of Interest to the Team)

1. I have read the definition of a medical directive in the glossary of terms and the college references. Can you describe it in terms of everyday ED practice?

2. Why do we need medical directives in an ED?

3. Who needs to be involved in developing an ED directive and who needs to sign off?

4. Why do all staff physicians need to sign directives?

5. As a physician, what can I expect from the hospital, ED administration and from the nurses/staff acting on the directive if I sign off?

6. As a physician, what should I do before I sign off?

7. Can the Chief sign off a directive on behalf of an ED physician (approval by proxy)?

8. What about patients who Leave Without Being Seen (LWBS)?

9. Will the use of directives increase unnecessary tests and procedures?

10. Is this kit saying directives are a substitute for timely assessment of patients by a physician?

11. Why a provincial project on directives?

12. Once we establish directives, how often do we have to routinely renew them, and do they need to be re-approved by everyone?

FAQs for the Entire Team

13. Why do we need orders in an ED?

14. When can the order be in the form of a medical directive?

15. I have read the physician’s FAQ document regarding who to involve in developing, reviewing and signing off directives (#3). Can you describe this in more detail?

16. As a nurse, how do I know when a directive is a proper order and I am fully authorized to implement a procedure pursuant to it?

17. When I implement a directive for a patient, what do I have to document?

18. We place a copy of the directive in the chart when a directive is implemented. Is this necessary?

19. We work in the hospital Pharmacy, the Laboratory and in Diagnostic Imaging. When an order sheet or requisition comes to us, signed off by someone other than a physician, how do we know it is a proper order, and are therefore authorized to carry it out?

20. As an administrative staff member, what do I need to do to assure proper use of directives?

21. What does ‘prototype’ medical directive mean and why have the medical directives in Module 3 been developed as ‘prototypes’?

22. How were the orders chosen for the prototype directives?

23. Why are the prototypes set up by orders and not by presenting complaints? Do we need to change our directives to this format?

24. The CEDIS Presenting Complaint Table (Module 5) makes sense. Why can’t we just use it as the directive?

25. Do we have to use pre-printed orders to document implementation of the directives?

26. Why do the prototypes only apply to the period between when a patient arrives in the ED to first contact with the attending physician?

27. I notice the directives designate nurses as authorized to implement directives. Can other staff members be designated as implementers as well?

28. The nurses in our ED need a more extensive orientation to implement orders for extremity x-rays. What should be included in their orientation?

29. How detailed does the information in a directive need to be?

30. Can we adopt the prototype directives word for word?
FAQs for Physicians (Of Interest to the Team)

1. I have read the definition of a medical directive in the glossary of terms and the college references. Can you describe it in terms of everyday ED practice?

   In an ED, a medical directive is an order that you would usually give verbally or in writing on behalf of a patient, only the order is for all patients meeting specific criteria. It enables nurses or other members of the health care team to anticipate common orders and act on them before the physician has a chance to see the patient.

2. Why do we need medical directives in an ED?

   In most EDs, nurses may perform interventions that, strictly speaking, require a medical directive. Common examples include starting an IV or placing oxygen on a patient. Empowering nurses to act before the physician has seen the patient allows certain basic emergency procedures to be completed more quickly and efficiently, leading to safer patient care and better patient flow. Consider two patients arriving with chest pain, one to an ED with all the directives suggested here and the other to an ED with no directives. The first patient will have the nurses start an IV, apply oxygen, obtain an EKG, send standard bloodwork and will be given ASA by the time the physician sees the patient. The other will only have their clothes removed and will be lying in a stretcher waiting for the physician to arrive. Clearly, directives can make our departments function better. Many EDs would perform some or all of the interventions suggested in the first ED without a formal set of directives; however, it is in everyone’s best interest to bring practice into full compliance with all of the rules and regulations governing patient care.

3. Who needs to be involved in developing an ED directive and who needs to sign off?

   The specific people to develop, review and sign off on directives and the extent to which they become involved may vary depending on the directive and hospital-specific roles and responsibilities. Those who need to sign off include:
   - All physicians whose patients may receive a test or procedure under authority of a directive;
   - Representatives of those implementing the directive;
   - Those with administrative responsibilities for ED practice (e.g. Chief, Manager, Director);
   - Those affected by the directives (e.g. physicians and staff of Diagnostic Imaging and Laboratory Services); and,
   - Corporate committees with responsibility for practice (e.g. Pharmacy and Therapeutics Committee, Medical Advisory Committee. Approval by Medical Advisory Committee may or may not be required, depending on medical policy within the hospital).

4. Why do all staff physicians need to sign directives?

   In the ED, at the time a patient arrives, the nurses can’t always know who will become the attending physician for that patient. For a directive to be implemented safely, a physician will have to assume the care of the patient, including following up on the consequences of the directives such as interpreting results of diagnostic tests or assessing the effect of therapeutic interventions. By signing the directive, physicians indicate that they are aware of the existence of the directive, agree with its intent and are prepared to assume the care of patients for whom directives have been implemented (assuming that they are properly notified of the directives’ implementation). If only some physicians sign off, it would be awkward or impossible for nurses to remember which directives had been signed by which physicians and to anticipate which physicians would be
assuming care of which patients. Thus, directives must represent only those interventions for which a consensus exists and all physicians must approve.

5. As a physician, what can I expect from the hospital, ED administration and from the nurses/staff acting on the directive if I sign off?

It is the responsibility of the hospital to provide a forum for development, review and approval of medical directives, including sign off by the hospital Medical Advisory Committee. It is also the hospital’s responsibility to establish a means by which a record of the directive is appropriately maintained in accordance with applicable recordkeeping standards to ensure that the version that applies for any episode of care is readily accessible, enabling physicians, staff and the hospital to provide appropriate care and fulfill practice expectations when using directives.

The ED leadership team must ensure that the directive is properly written, articulates the patient populations and circumstances the directive is to be used for, and defines how this is to be recorded and reported.

Next, it is the ED leadership team’s responsibility, along with nurses and regulated implementers themselves to ensure that they have the competencies and as necessary, the appropriate training to implement the directive. Competencies and training need to include patient assessment and screening, the indications and contraindications for implementing the directive and knowing how to record and report that the directive has been implemented.

The staff members using the directive in the ED are responsible to ensure that they are competent and practicing according to the criteria documented in the directive. The various health regulatory colleges such as the College of Nurses of Ontario (CNO) and the College of Respiratory Therapists of Ontario (CRTO) have very explicit expectations regarding the use of medical directives.

The hospital and ED leadership teams are responsible for ensuring ongoing quality monitoring and renewal of the directive.

The tools and templates provided in Modules 1 though 6, or hospital-specific ones, can be used to demonstrate fulfillment of these expectations.

6. As a physician, what should I do before I sign off?

You should review the directive to verify that you:

• Agree with its content, that it is an intervention that can be implemented safely and effectively, given the circumstances in your ED as you understand them;
• Are prepared to assume the care of patients who have had an intervention performed as authorized by the directive;
• Know how the staff will document or make you aware that a directive has been implemented so you can assume care appropriately;
• Consider the consequences on patient care if there is no directive, that is, balance the risk of the directive against the risks that may occur in the absence of a directive (usually related to delays in patient care).

If you are a regular staff member in an ED and you are presented with new directives, you should be sure you agree that the directive reflects what you understand to be clinical consensus in your ED. This is your opportunity to provide suggestions on directive content.
If you are a temporary or locum physician, you are likely neither able nor prepared to do such a detailed review. However, you still must familiarize yourself with the existing directives and processes at this ED as part of your orientation, just as you would familiarize yourself with consultant coverage, admitting processes, etc. before starting work.

Any physician who does not agree with the directive should take it up with the Chief of Staff (Chief).

We have provided some resources to ease the administrative and clerical burden on physicians through the use of batch directive sign-off forms and proxy forms, permitting Chiefs to sign off on directives for attendings under specified conditions. These are provided for convenience to be applied appropriately given the wide range of staffing circumstances in Ontario EDs. They do not alter the need for every physician to be aware and supportive of directives in use in their EDs.

For sample prototype approval forms, see Module 6 – Approval Forms.

7. Can the Chief sign off a directive on behalf of an ED physician (approval by proxy)?

Yes, as long as both parties – the Chief and ED physician – can ensure between them that the directive is communicated and agreed upon and that both have agreed in writing to this type of sign-off. When signing off by proxy, the Chief agrees to:

- Approve directives that are agreed to by the physician, and
- Ensure the physician is kept informed of the directive.

The physician agrees to:

- Review the directive to fully understand the conditions under which it will be implemented, including knowing how staff will document or make the physician aware that the directive has been implemented so s/he can assume care appropriately, and
- Assume the care of patients who have had an intervention performed as authorized by the directive.

ED Chiefs and physicians electing to use this option for sign-off must be comfortable with the joint responsibilities incurred. Avenues to address any differences that may arise should be established without compromising care and the accountabilities of staff involved in carrying out the directive. Either the Chief or physician can terminate the proxy at any time. If terminated, the ED physician would need to personally sign off on a directive.

See ‘Agreement for Chief to Approve Directive(s) on Behalf of ED Physician(s)” on p. 2, Module 6 – Approval Forms for a sample method to facilitate sign-off by proxy.

8. What about patients who Leave Without Being Seen (LWBS)?

In theory, a patient who is competent and leaves the ED at any stage of their assessment leaves at their own risk. In practice, if diagnostics ordered under a medical directive return after the patient has left and present important new clinical information that might have influenced their decision, you will want to review this information and determine appropriate action. Each hospital should have a documented process in place to deal with patients who LWBS, and this should include patients who have had a medical directive implemented before they leave.

See Appendix A, p.19 for sample policy considerations.

The prototype directives contain information for how to deal with patients who LWBS (see Module 3 – Prototype Medical Directives – Documentation sections).
9. Will the use of directives increase unnecessary tests and procedures? The CEDIS Presenting Complaints Table (Module 5) seems to indicate this by listing a whole series of tests and procedures to administer based on a presenting complaint.

Properly developed and applied directives should not increase unnecessary tests and procedures administered to patients. Instead, they optimize care by mandating physicians and teams to come to a consensus on practice issues and develop a consistent approach. In addition, time to key tests and interventions can be reduced improving care and satisfaction.

The CEDIS is not a directive, so it cannot authorize unnecessary tests and procedures. Rather, nurses may use it as a quick reference to determine what the full range of ED directives permit, but must still refer to each specific directive to determine exactly when a test or procedure may be performed.

10. Is this kit saying directives are a substitute for timely assessment of patients by a physician?

No, directives are not a substitute for timely assessment of patients by physicians. Patient specific orders provided by a physician who has seen and assumed care of a patient remain the gold standard for patient care. Proper directives authorize designated staff to carry out orders and expedite care before a physician is able to see a patient, but only when it is safe, effective and ethical to do so.

11. Why a provincial project on directives?

There is a lot of confusion in the province about many aspects of directives, including the correct process to follow in establishing directives, the responsibilities of signing physician staff and a lot of variation in directive use from ED to ED. Our project was intended to clarify these issues, assist EDs in developing or adapting their programs to be in full compliance with all governing legislation, and to create a set of directives that represents a broad consensus of physician and nursing experts in the field. We hope this process will lead to some consistency from site to site in directive content, and broaden the use of directives in order to improve patient care and patient flow in Ontario EDs. We do not intend this package to become a standard or required set of directives; rather our work is a resource that each ED can use or adapt as it sees fit.

12. Once we establish directives, how often do we have to routinely renew them, and do they need to be re-approved by everyone?

Directives are routinely reviewed and renewed as often as you deem appropriate, given hospital policy and the circumstances in your ED.

All parties are expected to sign off on renewals and changes. However, some parties may agree that if changes are not substantive, they do not need to sign off every time. For example, the Lab may agree that adding a commonly accepted indication to a lab test may not warrant review and sign-off by their representatives. However, if a change has cost, equipment or specimen processing implications, review and sign-off by the Lab would be warranted.

Regardless of decisions regarding sign-off, all those involved in implementing directives should always be notified of any changes.
FAQs for the Entire Team

13. Why do we need orders in an ED?

In hospitals, orders -- either direct orders or directives -- from a physician are necessary to comply with requirements to permit diagnostic procedures and treatments under the Public Hospital’s Act (PHA), performance of controlled act procedures under the Regulated Health Professions Act (RHPA) and performance of restricted acts under other pieces of health legislation, such as the Healing Arts Radiation and Protection Act (HARP Act) and Laboratory Specimen and Collection Centre Licensing Act (LSCCLA). Failure to have an order in place, when necessary, is a contravention of legislation, subject to penalties for staff and the hospital, including fines, jail terms, findings of professional misconduct and litigation proceedings.

For further information regarding orders, directives and legislative and regulatory expectations, see the CPSO, CNO and FHRCO guidelines along with other regulatory college materials.

14. When can the order be in the form of a medical directive?

In any situation where an order is required, a medical directive may be used when it is appropriate to enable staff to implement procedures prior to a physician’s assessment of the patient at the time. It is appropriate to use a directive when it is in the patient’s best interests and the steps for establishing it can be fulfilled (see Steps for Establishing Directives, Guidelines, p.2).

15. I have read the physician’s FAQ document regarding who to involve in developing, reviewing and signing off directives (#3). Can you describe this in more detail?

Basically, because of the nature of a directive, anyone affected needs to know about it and may be involved in developing, reviewing or signing off on it, depending upon their hospital-specific roles and responsibilities.

Those who may be involved in developing directives include:

- Chief physician or designate and, as desired, physician representatives.
- Representatives of implementing staff (e.g. educators, staff nurses).
- As desired, representatives of co-implementing staff (e.g. pharmacists, medical laboratory technicians, medical radiation technologists), to the extent they would like to be involved. They may wish to simply review it prior to approval.
- ED administrative staff (e.g. manager).

Those affected by implementation of the directive may review it prior to approval to ensure it is appropriate from their perspective. This includes:

- Chief physicians and representatives of co-implementing groups such as Laboratory and Diagnostic Imaging.
- Consulting physicians (representatives), when directives may affect their care of a patient. For example, Internal Medicine or Cardiology consultants may wish to review directives for patients with chest pain. (Note: These prototype directives do not contain orders warranting consulting physician review; however, EDs considering adding orders or conditions would want to keep this point in mind.)
- Corporate administrative staff and committees (e.g. Director of Nursing, affected professional practice leaders, professional practice committees).
Those who have accountabilities for practice as a result of the directives need to sign off including:

- Anyone involved in development;
- Laboratory and Diagnostic Imaging Department (representatives);
- Administrative staff and committees in accordance with hospital-specific organizational structures and roles and responsibilities (e.g. program administrators and committees, relevant professional practice committees); and,
- Corporate committees, e.g. Pharmacy and Therapeutics Committee (for directives involving medications) and Medical Advisory Committee (MAC) (depending on hospital medical policy, MAC may not need to sign off).

The Modules contain forms to facilitate development, review and sign-off by those involved. See:

- Module 1 – Performance Readiness Assessment, Section 14 – for a list to plan who should review and/or sign off on the directives
- Module 2 – Performance Readiness Plan, Performance Readiness Forms – for forms enabling implementers to sign-off
- Module 3 – Prototype Directives, Administrative and Physician Approval sections – to record approvals
- Module 6 – Approval Forms – to obtain approval signatures

16. As a nurse, how do I know when a directive is a proper order and I am fully authorized to implement a procedure pursuant to it?

You would know you are authorized to implement a procedure under a directive when the Steps for Developing ED Directives (p. 2, Guidelines) have been fulfilled such that:

- The directive has been properly constructed – i.e., it contains specific orders, for specific circumstances under specific conditions, including indications and contraindications, designated implementers, provisions for consent, documentation and quality monitoring -- and those affected have provided input and agree it’s appropriate.
- The ED Chief and all attending physicians have signed off, along with locums and casual physicians (who may sign off at the outset of a shift).
- Relevant hospital authorities - e.g. representatives of implementers, lab and diagnostic imaging and administrators (such as ED manager, Pharmacy and Therapeutics Committee, MAC) - have agreed and/or have signed off.
- You are competent and have successfully completed any necessary orientation and training to qualify as a designated implementer.

The modules contain templates and forms to facilitate nurses in determining when they are authorized to implement a procedure under a directive.

17. When I implement a directive for a patient, what do I have to document?

Communication and documentation of directive implementation is critical to providing safe care. When you implement a directive, you must document the following in the designated section of the patient record:

- Specific order,
- Name and number of the medical directive,
- Name of the attending physician (where known),
- Your name and signature, and
Conditions for implementing the directive, implementation and patient response.

It is recommended that you document the specific order in the order section of the chart, and the information pertaining to implementation and patient response in the progress notes, nursing notes, or appropriate form (e.g. medication administration record).

The prototype medical directives contain a documentation section with information on how to document the use of directives (see Module 3).

18. **We place a copy of the directive in the chart when a directive is implemented. Is this necessary?**

Placing the directive in the chart is one way of recording implementation of a directive that readily provides a record of the orders implemented and the conditions that have been assessed to permit implementation. Other options range from writing out the orders and conditions on the order sheet and in the progress notes respectively to developing preprinted forms, or using electronic methods. Each method has its pros and cons. The main features necessary for any method include:

- Ensuring the required information -- medical directive name and number, name and signature of the implemener and where known, the name of the attending physician -- are included. When EDs have more than one attending physician at a time, the directive may be implemented before the specific attending is identified. All attendings sign off on the directive in anticipation of its use, and are responsible for it as their order.
- Ensuring capacity to readily identify the content of the full directive at the time it was implemented. This means retaining a copy of the directive for the same time it is necessary to retain patient records, for a period of at least 10 years or much longer in some cases.

The documentation and communication section of the prototype directives contains information regarding documenting directive implementation (see Module 3). Samples of possible pre-printed orders along with a sample ED record are also available (see Module 4).

19. **We work in the hospital Pharmacy, the Laboratory and in Diagnostic Imaging. When an order sheet or requisition comes to us, signed off by someone other than a physician, how do we know it is a proper order, and are therefore authorized to carry it out?**

A medical directive is proper when the Steps for Developing ED Directives (see Guidelines, p. 2) have been fulfilled. This is indicated when the order sheets, requisitions or requests for consultation specifically refer to the directive in compliance with recommended format identifying the:

- Specific order;
- Name and number of the medical directive;
- Name of the attending physician (where known, when EDs have more than one attending physician at a time, the directive may be implemented before the specific attending is identified); and,
- Name and signature of the nurse or implemener implementing the order/directive

This information may be written out on the order sheet or requisition, or parts of it may be pre-printed. See Module 4 for Sample Pre-printed Orders that may be used with the prototype directives along with a Sample ED Record for recording implementation of directives.

This information on how to properly document directive orders is identified in the Documentation and Communication section of the prototype directives (see Module 3).
20. As an administrative staff member, what do I need to do to assure proper use of directives?

Ensure that the necessary infrastructure and processes exist to enable staff to develop, review and maintain directives and to work with them safely, effectively and ethically, in accordance with regulatory and legislative requirements. This includes enabling physicians and staff to:

- Determine the appropriateness of using a directive;
- Participate in developing and approving directives;
- Readily sign-off;
- Set up and participate in any necessary orientation/training;
- Readily confirm at the point of care that the directives constitute a proper order, with necessary content and approvals;
- Appropriately document directive implementation;
- Maintain the directives on an ongoing basis, with quality monitoring including monitoring and documenting compliance with the directive, and renewal as appropriate, in fulfillment of the steps for establishing ED directives (p. 2, Guidelines).

You also need to agree to and sign off on the directives, addressing your/hospital accountability for ensuring appropriate practices for staff within the hospital.

The Modules contain sample tools and directive prototypes for fulfilling these responsibilities.

21. What does ‘prototype’ medical directive mean and why have the medical directives in Module 3 been developed as ‘prototypes’?

The directives in Module 3 have been developed as prototypes and not as directives proper because each ED must establish its own directives that suit its specific circumstances, such as patient characteristics, team competencies, hospital and physician practice patterns and care delivery models. The prototype directives offer a consensus-based sample to draw upon and adopt to suit each ED’s circumstances.

22. How were the orders chosen for the prototype directives?

The orders were chosen based upon common ED practice patterns, directives and best practices identified by working group members. Each order had to fulfill the following criteria:

- Enhancement of patient flow
- Evidence-based
- Sustainable (practices unlikely to change or require updating)
- High volume (used for frequently encountered presentations)
- Practical (may be implemented by staff at all levels of experience)
- Relevant (applicable across all EDs)

23. Why are the prototypes set up by orders and not by presenting complaints? Do we need to change our directives to this format?

The prototypes have been set up this way to facilitate use, and to foster sharing and evaluation. If you find the format useful, you may wish to adopt it. However, you do not have to change your directives to this format, or ever adopt it if it does not suit your needs.
24. The CEDIS Presenting Complaint Table (Module 5) makes sense. Why can't we just use it as the directive?

The CEDIS Presenting Complaints Table can't be used as the directive on its own because it doesn't contain the indications, contraindications and other information necessary for a proper directive. However, it can be used as a quick reference or prompt once staff is fully familiar with the information in the directives. As well, using the CEDIS presenting complaints framework may offer a means of evaluating the effect of the directives on patient flow.

25. Do we have to use pre-printed orders to document implementation of the directives?

No, you do not have to use pre-printed orders. The Sample Pre-printed Orders corresponding to the prototype directives in Module 4 are presented for your reference and convenience, in the event you would like to use them to develop pre-printed orders for documenting your ED directives.

26. Why do the prototypes only apply to the period between when a patient arrives in the ED to first contact with the attending physician?

The prototypes have been written to apply only to this period to coordinate care and minimize margins for misunderstanding, oversight and error. Once the attending ED physician sees the patient, s/he can provide orders and there is no confusion regarding how orders will be given and who is responsible. In developing your own directives, each ED can decide if this is the preferred approach for coordinating care under directives, or if you prefer to coordinate in another fashion.

27. I notice the directives designate nurses as authorized to implement directives. Can other staff members be designated as implementers as well?

This is to maximize clarity of the information being conveyed, and because all EDs are staffed with nurses. However, any practitioner with the necessary competencies, given the circumstances within an ED may be designated to implement a directive.

28. The nurses in our ED need a more extensive orientation to use a directive for extremity x-rays. What should be included in their orientation?

A more extensive orientation is often warranted to enable nurses to implement a physician's directive order for extremity x-rays safely and effectively. The components of such an orientation would include the following:

- Review of anatomy and physiology of upper and lower extremities
- History and objective assessment of limb injuries (i.e. mechanism of injury, CSM, Ottawa knee, ankle and foot rules)
- Common injury patterns as it relates to limb injuries
- Contraindications for initiation of medical directive (i.e. CSM deficits, open fracture, obvious dislocation, pregnancy, etc.)
- Documentation requirements to initiate medical directive (i.e. limb to be x-rayed, inclusion of brief history with mechanism of injury, CSM status, weight bearing, pregnant, etc.)
The prototype directive for extremity x-rays identifies relevant references and websites (see the Diagnostic Imaging Directive in Module 3 - Prototype Directives), along with Sample Quiz Questions and Answers and a Sample Extremity X-ray Medical Directive Certification Record that identifies competency requirements (see Module 2 - Performance Readiness Plan).

29. How detailed does the information in a directive need to be?

A directive needs to be detailed enough to ensure that anyone implementing it has the information necessary to make exactly the same decision as the ordering physician would make about what to do for whom, and when. Additionally, it must clearly identify how implementation is documented, and how to manage any untoward outcomes so all team members can coordinate to ensure safe, effective care.

30. Can we adopt the prototype directives word for word?

Yes, if they apply word for word to the circumstances in your ED.
Appendix A

Sample Policy Considerations For Patients Who Leave Without Being Seen (LWBS)

- Clarify that the decision to LWBS is the patient’s and that LWBS is actively discouraged by staff.

- Assign a physician to be responsible for LWBSs on each shift.

- Establish a guideline for what warrants patient notification after a patient has left – e.g. abnormal results – and that the decision to notify needs to be made on a case-by-case basis by the assigned physician in light of the information s/he has at hand.

- Identify what the triage/assigned nurse is expected to do, for example:
  - Contact physician immediately, prior to the patient leaving, if the patient’s capacity or condition are a concern
  - Ensure the chart is complete, with the record of results available for review by the physician (may assign this to support staff if appropriate)
  - Participate in patient notification as indicated, once the physician has determined the appropriate action based on his/her review of the chart

- Identify how and where to document.

- Identify whom to notify corporately in the event of any issues.
MODULE 1

Performance Readiness Assessment

ED Prototype Medical Directives Module 1 of 6

Use this tool to help determine if the conditions to support using directives exist in your ED
Table of Contents

Performance Readiness Assessment................................................................. 1
## Performance Readiness Assessment

For Determining the Appropriateness of Establishing Directives, Delegation and Performing Procedures beyond Principal Expectations of Practice

Template from: An Interprofessional Guide on Orders, Directives and Delegation; Federation of Health Regulatory Colleges of Ontario (January 2007)

<table>
<thead>
<tr>
<th>Title/Procedure:</th>
<th>ED Medical Directives</th>
</tr>
</thead>
<tbody>
<tr>
<td>Applicable Authorizing Mechanism:</td>
<td>☑ Delegation ☑ Medical Directive ☑ Direct Order ☐ Unnecessary</td>
</tr>
<tr>
<td>Authorizing Profession:</td>
<td></td>
</tr>
<tr>
<td>Implementing Profession:</td>
<td></td>
</tr>
<tr>
<td>Patient(s):</td>
<td></td>
</tr>
<tr>
<td>Disposition:</td>
<td>☑ Approved ☑ Being forwarded for Approval ☐ Not Approved</td>
</tr>
<tr>
<td>Date:</td>
<td></td>
</tr>
</tbody>
</table>

**Sponsors** *(This section for use in large multi-professional settings)*

<table>
<thead>
<tr>
<th>Representative(s) of Authorizing Profession:</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Representative(s) of Implementing Profession:</td>
<td></td>
</tr>
<tr>
<td>Administrative Representative(s):</td>
<td></td>
</tr>
</tbody>
</table>

- Have all applicable stakeholders been consulted: *(See Section 11 for list)*
  - ☑ Yes ☐ No
- Is a completed Medical Directive or Delegation template attached:
  - ☑ Yes ☑ No ☐ N/A
- Is a completed Performance Readiness Plan attached:
  - ☑ Yes ☑ No ☐ N/A
**Assessment Parameters**

1. **Reason and Specific Benefits of the Directive or Delegation:**
   1.1. Does establishing the directive or delegation address patients’ best interests?  
      - [ ] Yes  
      - [ ] No  
      - [ ] Unsure  
      Comments:

2. **Authorizer:**
   Does the authorizer:
   2.1. Have the scope, authority from their college, competencies and privileges (where applicable) to authorize performance?  
      - [ ] Yes  
      - [ ] No  
      - [ ] Unsure  
   2.2. Have an established or anticipated professional relationship with the patient?  
      - [ ] Yes  
      - [ ] No  
      - [ ] Unsure  
   2.3. Agree the directive applies to all his/her patients who meet the conditions?  
      - [ ] Yes  
      - [ ] No  
      - [ ] Unsure  
   2.4. Have the ability to provide ongoing supervision directly, or are other provisions for appropriate supervision in place?  
      - [ ] Yes  
      - [ ] No  
      - [ ] Unsure  
      Comments:

3. **Implementer:**
   Does the implementer:
   3.1. Have the scope and authority from their own college (where applicable) to perform the procedure(s)  
      - [ ] Yes  
      - [ ] No  
      - [ ] Unsure  
   3.2. Have the baseline competencies to perform the proposed procedure(s) and manage the outcomes given the:
      3.2.1. predictability of the patient’s condition and needs,  
      - [ ] Yes  
      - [ ] No  
      - [ ] Unsure  
      3.2.2. predictability of the procedure and its outcomes, and  
      - [ ] Yes  
      - [ ] No  
      - [ ] Unsure  
      3.2.3. circumstances in the situation including resources and safeguards (such as established standards of practice, written materials, back-up and supervision), and opportunities to attain and maintain competence?  
      Comments:

4. **Consent:**
   4.1. Can informed consent be properly obtained?  
      - [ ] Yes  
      - [ ] No  
      - [ ] Unsure  
      Comments:
### 5. Review and Quality Monitoring Processes:

<table>
<thead>
<tr>
<th>Question</th>
<th>Yes</th>
<th>No</th>
<th>Unsure</th>
</tr>
</thead>
<tbody>
<tr>
<td>5.1. Is there a process in place to ensure a regular review of the directive or delegation?</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>5.2. Is there a process in place to address questions or concerns arising from the directive or delegation?</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Comments:

### 6. Practice Setting Feasibility

<table>
<thead>
<tr>
<th>Question</th>
<th>Yes</th>
<th>No</th>
<th>Unsure</th>
</tr>
</thead>
<tbody>
<tr>
<td>6.1. Are the necessary human and material resources available to support the practice?</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>6.2. Is the practice sustainable? (For example, can new staff readily adopt the practice? If intensive resources are required to support the practice over the longer-term, is this feasible?)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>6.3. Does the practice broadly support effective health care delivery? (For example, if implementers are responsible for implementing the directive or delegation or performing the proposed procedure, will other services only they can provide be disrupted? Will other team members or care delivery systems be negatively impacted? Can these effects be offset?)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>6.4. Can any billing, cost or liability considerations be appropriately managed?</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>6.5. Are there any other situation-specific factors to consider?</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Comments:

### 7. Risk/Benefit Analysis:

<table>
<thead>
<tr>
<th>Question</th>
<th>Yes</th>
<th>No</th>
<th>Unsure</th>
</tr>
</thead>
<tbody>
<tr>
<td>7.1. Do the benefits of proceeding by way of the directive, delegation or practice outweigh the risks?</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Comments:

### 8. Education and Performance Readiness Plan:

<table>
<thead>
<tr>
<th>Question</th>
<th>Yes</th>
<th>No</th>
<th>Unsure</th>
</tr>
</thead>
<tbody>
<tr>
<td>8.1. Is there a plan for enabling implementers to attain the necessary competencies and achieve performance readiness? (Identify a basic plan here, or where the plan is more involved, refer to the Performance Readiness Plan.)</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Comments:
9. Communication Plan:

9.1. Is there a plan for informing stakeholders and for activating the directive, delegation or practice? □ Yes □ No □ Unsure

Comments:

10. References to Support Practice:

10.1. Are there references to support practice? (References may be listed here or attached) □ Yes □ No □ Unsure

Comments:

11. Those Consulted for Input:

11.1. Have all affected stakeholders been consulted? List those consulted in the table below. Add or delete stakeholders to correspond to your practice setting. □ Yes □ No □ Unsure

Comments:

<table>
<thead>
<tr>
<th>Stakeholders Consulted</th>
<th>Names/Positions</th>
<th>Agree?</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Authorizers</td>
<td></td>
<td>□ Yes □ No</td>
</tr>
<tr>
<td>2. Implementers:</td>
<td></td>
<td>□ Yes □ No</td>
</tr>
<tr>
<td>• Implementer(s) or representatives</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Co-implementers (if applicable)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Educators (if applicable)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3. Administrators (List)</td>
<td></td>
<td>□ Yes □ No</td>
</tr>
<tr>
<td>4. Professional Leaders of:</td>
<td></td>
<td>□ Yes □ No</td>
</tr>
<tr>
<td>• Authorizers</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Implementers</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Co-implementers (if applicable)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5. Applicable profession-specific groups/committees of:</td>
<td></td>
<td>□ Yes □ No</td>
</tr>
<tr>
<td>• Co-implementers (if applicable)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6. Program/Corporate Committees (List)</td>
<td></td>
<td>□ Yes □ No</td>
</tr>
<tr>
<td>7. Pharmacy &amp; Therapeutics Committee</td>
<td></td>
<td>□ Yes □ No</td>
</tr>
<tr>
<td>8. Medical Advisory Committee</td>
<td></td>
<td>□ Yes □ No</td>
</tr>
</tbody>
</table>
Use these sample tools to establish the competencies for using a directive (Includes plan, quiz with answers and competency 'certification' forms)
Performance Readiness Plan ................................................................. 1

Prototype ED Directive Quiz
Questions
  Working with Medical Directives ...................................................... 2
  Laboratory Tests and Diagnostic Procedures ..................................... 3
  Extremity X-Rays ............................................................................. 3
  Medications ..................................................................................... 6
  Therapeutic Procedures .................................................................. 8
  Paediatrics ..................................................................................... 9

Answers
  Working with Medical Directives ...................................................... 10
  Laboratory Tests and Diagnostic Procedures ..................................... 11
  Extremity X-Rays ............................................................................. 11
  Medications ..................................................................................... 14
  Therapeutic Procedures .................................................................. 16
  Paediatrics ..................................................................................... 17

Sample Extremity X-Ray Medical Directive Certification Form
(Identifying competency requirements) ............................................... 18

Sample Implementer Performance Readiness (‘Certification’) Forms
  Individual ......................................................................................... 19
  Group .............................................................................................. 20

Acknowledgements:

Performance Readiness Plan Template and Implementer Performance Readiness Implementer Forms from An Interprofessional Guide on the Use of Orders, Directives and Delegation; Federation of Health Regulatory Colleges of Ontario (January 2007)

Sample Quizzes provided by Southlake Regional Health Centre (Laboratory Tests and Diagnostic Procedures), Mount Sinai Hospital (Extremity X-Rays), Grand River Hospital (Medications) and North York General Hospital (Therapeutic Procedures and Paediatrics)

Sample Extremity X-Ray Medical Directive Certification Form provided by North York General Hospital
### Performance Readiness Plan

From: An Interprofessional Guide on the Use of Orders, Directives and Delegation; Federation of Health Regulatory Colleges of Ontario (January 2007)

**Procedure:**

**Date:**

**Plan Endorsed by:**
*(name, position, signature)*

**Designated Educators**
*(if applicable; name, position, signature)*

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>1. Competence and Authority of Educator(s)</strong> (if applicable)</td>
<td>Identify whether any applicable educators have the scope, authority from their College and competencies to perform and teach the procedure.</td>
</tr>
</tbody>
</table>

**Comments:**

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>2. Education Plan</strong></td>
<td>Identify the:</td>
</tr>
</tbody>
</table>

2.1. Knowledge, Skills and Judgment Component (Attach any relevant slides, references and hand-outs).

For relevant information, see:

- Prototype ED Medical Directives Guidelines
- ED Medical Directives established for your ED identifying full clinical criteria and conditions. (Background: Prototype ED Medical Directives, Module 3)
- CEDIS Presenting Complaints Table, for a quick reference of directive orders by presenting complaints. Adapt to suit your ED-specific directives. (Background: Module 5)

2.2. Supervised Practice Component (If any).

2.3. Evaluation of Competence Component (Attach any relevant test materials).

For sample competence evaluation questions, see:


**Comments:**

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>3. Plan for Assuring Ongoing Competence</strong></td>
<td>3.1. Identify the plan for assuring ongoing competence.</td>
</tr>
</tbody>
</table>

**Comments:**

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>4. Practical Arrangements</strong></td>
<td>4.1. Identify the arrangements for delivering the education, both initially and ongoing.</td>
</tr>
</tbody>
</table>

**Comments:**
Prototype ED Directive Quiz

Working with Medical Directives

1. Orders from physicians, dentists, midwives or for outpatients, RN(EC)s are required for treatments and diagnostic tests within a hospital.

   True  False

2. Not having an order when one is required constitutes grounds for professional misconduct and may be subject to a fine or jail term.

   True  False

3. The medical directives are __________________ from ED attending physicians for __________________ or interventions for a specific group of __________________ under specific __________________.

4. A medical directive comes into force as a proper order once:
   a) Staff representatives have had input and agree with the directive.
   b) Relevant stakeholders (e.g. lab and diagnostic imaging) and administrators (e.g. ED manager, MAC) have signed off.
   c) All ED attending physicians have signed off, along with locums and casual physicians who may sign off at the outset of a shift.
   d) All of the above.

5. The CEDIS Presenting Complaints Table can be used as the medical directive.

   Yes  No

6. Only staff members who have successfully completed the ED Medical Directive Orientation Program can implement procedures under authority of a directive, without a physician writing a direct order for the procedure.

   True  False

7. To facilitate care and minimize margin for error due to lack of coordination, the directive applies to the period from __________________ to __________________. After this, the attending will provide direct orders (unless your ED sets up the directive differently, in which case, describe here).

8. Where in the patient record do you document:
   • The directive order, once you decide to implement it? ____________________________
   • Indications for implementation, implementation and patient response ____________________________

9. Documentation and communication of implementation of a directive is critical. Why?

   ____________________________

10. If an untoward or unanticipated outcome arises out of the use of the directive, you would contact the ____________________________ for any matters related to patient care and ____________________________ to follow-up on disposition for the directive.

11. A copy of the directive is available at ____________________________

12. Sign-off can be confirmed by ____________________________
Laboratory Tests and Diagnostic Procedures

1. A medical directive is not required for a nurse to order and perform a capillary blood glucose.
   a) True
   b) False

2. Serum drug screen is ordered under the medical directive for laboratory tests for the following patient presentations:
   a) reported ingestion
   b) altered LOC
   c) current warfarin use
   d) all of the above
   e) a and b

3. Urine for R&M should be sent when POCT urinalysis is positive for:
   a) Leukocytes
   b) Proteins
   c) Blood
   d) Nitrates,
   e) Glucose

Extremity X-rays

Ankle Injury:
A 26 year old female was playing baseball and slid into first base. She caught her foot on the base and the ankle rolled inward. She was not able to walk on it at the time of injury and still can not weight bear. The ankle is swollen and tender over the lateral malleolus, warm to touch, pedal pulse present.

1. Does she meet the Ottawa Ankle Rules criteria for an ankle x-ray? Yes No

Leg Pain
A 67 year old male presents with left ankle swelling and pain. There is no history of traumatic injury to the ankle and the patient is able to walk on it with a limp. There is moderate swelling to the entire lower leg, the foot is warm and the pedal pulse is present.

2. Does he meet the Ottawa Ankle Rules criteria for an ankle x-ray? Yes No
Foot Pain
A 20 year old male presents with right foot pain that began when he kicked in a door. He has pain in the midfoot zone. He has no other injuries and vital signs are within normal limits.

3. Which of the following findings would indicate he needs a foot x-ray?
   a) A hematoma noted over the anterior foot
   b) Tenderness at the lateral malleolus
   c) Tenderness at the base of the 5th metatarsal

Great Toe Injury
A 40-year-old female arrives complaining of pain at the great toe. She caught her toe on a piece of furniture and felt it "snap backwards". The toe is swollen and bruised. She has no other injuries, no medical conditions and vitals are within normal limits.

4. Does she meet the criteria to have an x-ray? Yes No

Ankle Injury
A hospital employee arrives in the emergency department after Occupational Health closed. He twisted his ankle while pushing a stretcher. There is no swelling or deformity, the patient is able to walk on the injured ankle, the foot is warm and the pulse is present. There is no tenderness over the maleolus.

5. Does he meet the criteria to have an x-ray? Yes No

Left Heel Pain
A 50 year old male arrives complaining of pain over the left achilles. He was playing tennis and felt pain and a "pop" at the achilles when he lunged to hit the ball. The foot is warm, pedal pulse is present.

6. Does he meet the criteria to have an x-ray? Yes No

Ankle Injury
A 30 year old female arrives with her left foot angulated/dislocated. She slipped and twisted her ankle on the ice in her driveway while carrying groceries into the house. The foot is pale, cool and the pulse is barely palpable.

7. What is the most appropriate nursing action?
   a) send the patient directly for x-ray
   b) have her assessed by the physician immediately
   c) splint the leg
   d) apply ice
Right Knee Pain
A 19 year old male arrives complaining of right knee pain. He was ice skating and was pushed down. He landed on the ice on his right knee. The knee is swollen and painful.

8. What physical findings indicate that he requires a knee x-ray?
   a) large amount of swelling
   b) large hematoma
   c) patellar tenderness
   d) he is able to flex the knee and weight bear

Patellar Dislocation
A 20 year old with a history of patellar dislocation arrives in the emergency with obvious patellar dislocation. His knee is flexed and supported by a pillow, the patella is clearly dislocated. He is screaming in pain.

9. Should the nurse implement the medical directive to have the knee x-rayed?    Yes                   No

Left Hand Injury
A 22 year old male arrives with a left-hand injury. He got angry and punched a hole in his apartment wall. His hand is swollen and painful, the fingers are pink.

10. Does he require an x-ray?            Yes                   No

11. What injury does the nurse suspect?
   a) boxers fracture base of the 5th metacarpal
   b) fractured distal phalynx
   c) the hand is an unlikely site for fractures

Right Wrist Pain
A 40 year old female arrives complaining of right wrist pain. She fell onto an outstretched wrist while skating. She has no other injuries and vital signs are within normal limits. Her wrist is painful, has an obvious deformity and swelling over the radial surface. The fingers are pink, she has no paraesthesia and can move all fingers.

12. Does she require an x-ray?            Yes                   No

13. What first aid measures can be done for this patient at triage?
   a) ________________________________
   b) ________________________________
   c) ________________________________

14. This same patient should be assessed for a scaphoid fracture. What physical findings would support the decision to order scaphoid views?
   a) ________________________________
   b) ________________________________
   c) ________________________________
Left Hand Pain
A 32 year old female slipped on the sidewalk and landed on an outstretched left hand. The injury occurred four days ago. She complains of arm pain and is not able to straighten the arm. She is not tender at the wrist, but she is tender at the lateral elbow.

15. What is the most appropriate x-ray?
   a) Left elbow
   b) Left elbow and wrist
   c) Left forearm
   d) Left wrist

16. All patients with orthopedic injuries should have the five Ps assessed and documented. What are the five Ps?
   a) _________________________________
   b) _________________________________
   c) _________________________________
   d) _________________________________
   e) _________________________________

17. List the contraindications for implementing orders for extremity x-rays.
   a) ________________________________
   b) ________________________________
   c) ________________________________

Medications

1. Which of the following statements concerning the administration of Acetaminophen or Ibuprofen for adult pain relief are correct:
   a) Indicated for patients triaged as CTAS 3-5 with mild to moderate pain (≤ 7/10 on pain scale)
   b) Pain is related to headache, dental, ear, nose and throat, MSK or skin injuries
   c) The primary clinical contraindication is abdominal pain
   d) A single dose of Acetaminophen 975-1000mg or Ibuprofen 600mg can be administered either po or pr. If the preceding dose has been one of these medications, the other may be administered
   e) all of the above

2. 32 year old Luke Razorblade presents to the ED complaining of a general body aches, sore throat and fever. His temperature at triage is 38.1°C. Given that he has not taken anything for the fever you can administer either Acetaminophen 975-1000mg or Ibuprofen 600mg po.
   a) True
   b) False
3. 66 year old Dennis Denial is triaged as CTAS 2 with chest pain (cardiac features) even though he insists it's just terrible indigestion from his Taco Bell lunch. One of the first drugs you will want to administer will be ASA 160 mg chewable tablet x 1 dose.
   a) True
   b) False

4. A 19 year old female with a history of substance abuse arrives by ambulance with an altered LOC and a blood sugar of 5mmol/L. Once you have established IV access you can administer the following:
   a) Dextrose 50% (25 gms in 50mLs) IV push over 2-3 minutes
   b) Narcan 0.4mg iv push
   c) None of the above

5. In assessing a patient with a full thickness laceration, which of the following statements concerning the medical directive for tetanus immunization are correct:
   a) presents with an open injury to the skin or eye
   b) has completed their primary immunization and hasn't received a booster in 10 years
   c) Td (Tetanus and Diptheria Toxoid) 1.0mLs is given IM in the deltoid muscle
   d) A and B only
   e) All of the above

6. 54 year old Drew Maurier presents to the ED with shortness of breath, B/P 150/92, P 110, RR 24,T 36.4, O2 Sat. 93%. In addition to burning the pack of cigarettes hanging out of his top pocket you want to administer Salbutamol (Ventolin) four to eight puffs or give a wet neb masking (5mg/mL);1 mL in 3 mL saline to assist his SOB. Before you can do this however, the following criteria must be met:
   a) Patient has a history of asthma or COPD
   b) Presents with dysphagia, muffled voice stridor and other symptoms of upper airway compromise
   c) Presents with any of the following symptoms- cough ,presence of respiratory distress, wheeze, tightness or decreased breath sounds on chest auscultation
   d) All of the above
   e) A and C only

7. Any male patient requiring urinary catheterization who is not allergic to amide anaesthetics or has a known urethral trauma or structural abnormality can receive:

   Lidocaine Jelly 2% 200mg/10mLs inserted into the urethra five minutes prior to catheterization to assist in comfort and ease of catheter passage
   a) True
   b) False
8. LET (Lidocaine, Epinephrine and Tetracaine) topical anaesthetic compound can be applied 30 minutes prior to suturing in which of the following situations:
   a) lacerations to the ear, nose, fingers and toes
   b) only lacerations that do not extend beyond the dermis
   c) lacerations that extends beyond the dermis to underlying structures (e.g. bone, cartilage, tendon, etc)
   d) A and C
   e) B only

9. 33 year old Sam Squint arrives from his workplace after splashing a chemical in his left eye and is in a fair amount of discomfort. You have a medical directive to instill two drops of ophthalmic anesthetic eye drops to Sam’s eye. Does he meet the criteria?
   a) Yes
   b) No

Another criteria for instilling these eye drops is in anticipation of Morgan Lens insertion pain caused by foreign body, thermal injury or corneal abrasion.
   a) True
   b) False

### Therapeutic Procedures

1. A saline lock or peripheral IV may be initiated under the following circumstances:
   a) Signs of airway compromise, respiratory distress, shock, dehydration, bleeding, or altered LOC
   b) In anticipation of IV medication administration for pain control, blood and blood product administration, or to provide IV rehydration
   c) History of high risk mechanism of injury, infection in the immunocompromised patients, or overdose
   d) A and B only
   e) All of the above

2. Oxygen therapy may be initiated when the patient presents with signs and symptoms of one or more of the following:
   a) Respiratory distress, SaO2 < 92%, or below established desirable range for the individual patient
   b) Hemodynamic instability
   c) There is evidence of suspected hypoxemia (chest pain, tachycardia, hemorrhage, hypovolemia, sickle cell, altered LOC, Trauma, smoke and/or toxin inhalation
   d) B and C only
   e) All the above
3. Eye irrigation with 1 L NS or Ringer’s Lactate may **NOT** be initiated when…
   a) Penetrating eye trauma, foreign body in the eye, or signs and symptoms of ruptured globe (obvious bleeding)
   b) The patient has flushed prior to arriving in the ED
   c) The patient has not had topical ophthalmic anaesthetic instilled
   d) The patient has suffered a flash burn
   e) All of the above.

**Paediatrics**

1. Salbutamol and Ipratropium Bromide may be administered to a child when...
   a) The child presents with signs and symptoms of upper airway pathology e.g. stridor, drooling, muffled voice, dysphagia
   b) The child presents with audible wheezing, wheezing with retractions, spasmodic cough, dyspnea, tachypnea, or decreased air entry into lung fields on auscultation with a history of reactive airway disease
   c) The child presents with audible wheezing, wheezing with retractions, spasmodic cough, dyspnea, tachypnea, or decreased air entry into lung fields on auscultation with **NO** history of reactive airway disease
   d) A and B only
   e) None of the above

2. A three month old child was given a therapeutic dose of Acetaminophen two hours prior to arriving in the ED. The child’s temperature is 38.5°C. I can administer Ibuprofen as per the mother’s request.
   a) True
   b) False

3. The child presents to the ED with vomiting and diarrhea. The 20 kg child was started on ORT on arrival. The mother advises you the child vomited a moderate amount of emesis and just had a large watery stool while waiting in the room. You should:
   a) Stop giving the ORT as it will precipitate more vomiting
   b) The child should receive 1000 cc of ORT slowly and then be given 200cc ORT for every loose stool
   c) The child should receive 500 cc of ORT slowly and then be given 100 cc ORT for every loose stool
   d) We should start an IV to rehydrate the child
   e) None of the above
Quiz Answers

Working with Medical Directives

1. Orders from physicians, dentists, midwives or for outpatients, RN(EC)s are required for treatments and diagnostic tests within a hospital.
   
   True  False

2. Not having an order when one is required constitutes grounds for professional misconduct and may be subject to a fine, jail term or being assessed damages in a civil proceeding.

   True  False

3. The medical directives are orders from an ED attending physician(s) for procedures, tests or interventions for a specific group of patients under specific conditions.

4. A medical directive comes into force as a proper order once:

   a) It is clear the directive has been properly constructed – i.e. it contains all information necessary to make it a proper order (specific orders, designated implementers, indications and contraindications, consent, documentation and quality monitoring provision), and there has been an assessment of the appropriateness of use, with input and agreement from those affected by it.

   b) ED Chief and all attending physicians have signed off, along with locums and casual ED physicians (who may sign off at the outset of a shift).

   c) Relevant hospital authorities - e.g. representatives of implementers, lab and diagnostic imaging and administrators (such as ED manager, MAC) - have signed off.

   d) All of the above.

5. The CEDIS Presenting Complaints Table can be used as the medical directive.

   Yes  No

Because the table does not contain the indications, contraindications and other information necessary for a proper directive, it cannot be used as a directive on its own. It can be used as a quick reference or prompt once staff are fully familiar with the information in the actual directives.

6. Only staff members who have successfully completed the ED Medical Directive Orientation Program can implement procedures under authority of a directive, without a physician writing a direct order for the procedure.

   True  False

7. If an untoward or unanticipated outcome arises out of the use of the directive, you would contact the attending physician for any matters related to patient care and (Identify contacts for your ED) to follow-up on disposition for the directive.

8. Where in the patient record do you document:

   • The directive order, once you decide to implement it? (Identify where – orders sheets, pre-printed orders, triage record, etc)

   • Indications for implementation, implementation & patient response? (Identify where – progress notes, nursing notes, MAR, other documentation forms, etc)
9. Documentation and communication of implementation of a directive is critical. Why?
   
   Coordination of care, patient safety – an established, reliable means of documenting implementation of a directive enables the physician to assume medical care safely and effectively as well as enabling anyone else involved in care of the patient to coordinate care effectively.

10. If an untoward or unanticipated outcome arises out of the use of the directive, you would contact the attending physician for any matters related to patient care and (Identify contacts for your ED) to follow-up on disposition for the directive.

11. A copy of the directive is available at (Identify where for your ED)

12. Sign-off can be confirmed by: (Identify how for your ED)

   Laboratory Tests and Diagnostic Procedures

1. A medical directive is not required for a nurse to order and perform a capillary blood glucose.
   a) True
   b) False

2. Serum drug screen is ordered under the medical directive for laboratory tests for the following patient presentations:
   a) Reported ingestion
   b) Altered LOC
   c) Current warfarin use
   d) All of the above
   e) A and B

3. Urine for R&M should be sent when POCT urinalysis is positive for:
   Leukocytes
   Proteins
   Blood
   Nitrates,
   Glucose

   Extremity X-rays

   Ankle Injury
   A 26 year old female was playing baseball and slid into first base. She caught her foot on the base and the ankle rolled inward. She was not able to walk on it at the time of injury and still can not weight bear. The ankle is swollen and tender over the lateral malleolus, warm to touch, pedal pulse present.

   1. Does she meet the Ottawa Ankle Rules criteria for an ankle x-ray? Yes No
Leg Pain
A 67 year old male presents with left ankle swelling and pain. There is no history of traumatic injury to the ankle and the patient is able to walk on it with a limp. There is moderate swelling to the entire lower leg, the foot is warm and the pedal pulse is present.

2. Does he meet the Ottawa Ankle Rules criteria for an ankle x-ray? Yes No

Foot Pain
A 20 year old male presents with right foot pain that began when he kicked in a door. He has pain in the midfoot zone. He has no other injuries and vital signs are within normal limits.

3. Which of the following findings would indicate he needs a foot x-ray?
   a) A hematoma noted over the anterior foot
   b) Tenderness at the lateral malleolus
   c) Tenderness at the base of the 5th metatarsal

Great Toe Injury
A 40-year-old female arrives complaining of pain at the great toe. She caught her toe on a piece of furniture and felt it "snap backwards". The toe is swollen and bruised. She has no other injuries, no medical conditions and vitals are within normal limits.

4. Does she meet the criteria to have an x-ray? Yes No

Ankle Injury
A hospital employee arrives in the emergency department after Occupational Health closed. He twisted his ankle while pushing a stretcher. There is no swelling or deformity, the patient is able to walk on the injured ankle, the foot is warm and the pulse is present. There is no tenderness over the maleolus.

5. Does he meet the criteria to have an x-ray? Yes No

Left Heel Pain
A 50 year old male arrives complaining of pain over the left achilles. He was playing tennis and felt pain and a "pop" at the achilles when he lunged to hit the ball. The foot is warm, pedal pulse is present.

6. Does he meet the criteria to have an x-ray? Yes No

Ankle Injury
A 30 year old female arrives with her left foot angulated/dislocated. She slipped and twisted her ankle on the ice in her driveway while carrying groceries into the house. The foot is pale, cool and the pulse is barely palpable.

7. What is the most appropriate nursing action?
   a) Send the patient directly for x-ray
   b) Have her assessed by the physician immediately
   c) Splint the leg
   d) Apply ice
Right Knee Pain
A 19 year old male arrives complaining of right knee pain. He was ice skating and was pushed down. He landed on the ice on his right knee. The knee is swollen and painful.

8. What physical findings indicate that he requires a knee x-ray?
   a) Large amount of swelling
   b) Large hematoma
   c) Patellar tenderness
   d) He is able to flex the knee and weight bear

Patellar Dislocation
A 20 year old with a history of patellar dislocation arrives in the emergency with obvious patellar dislocation. His knee is flexed and supported by a pillow, the patella is clearly dislocated. He is screaming in pain.

9. Should the nurse implement the medical directive to have the knee x-rayed?  Yes  No

Left Hand Injury
A 22 year old male arrives with a left-hand injury. He got angry and punched a hole in his apartment wall. His hand is swollen and painful, the fingers are pink.

10. Does he require an x-ray?  Yes  No

11. What injury does the nurse suspect?
   a) Boxers fracture base of the 5th metacarpal
   b) Fractured distal phalynx
   c) The hand is an unlikely site for fractures

Right Wrist Pain
A 40 year old female arrives complaining of right wrist pain. She fell onto an outstretched wrist while skating. She has no other injuries and vital signs are within normal limits. Her wrist is painful, has an obvious deformity and swelling over the radial surface. The fingers are pink, she has no parasthesia and can move all fingers.

12. Does she require an x-ray?  Yes  No

13. What first aid measures can be done for this patient at triage?
   a) Application of ice
   b) Application of splinting
   c) Application of sling

14. This same patient should be assessed for a scaphoid fracture. What physical findings would support the decision to order scaphoid views?
   a) Tenderness over anatomical snuffbox
   b) Pain over scaphoid area with longitudinal loading of thumb
Left Hand Pain
A 32 year old female slipped on the sidewalk and landed on an outstretched left hand. The injury occurred four days ago. She complains of arm pain and is not able to straighten the arm. She is not tender at the wrist, but she is tender at the lateral elbow.

15. What is the most appropriate x-ray?
   a) **Left elbow**
   b) Left elbow and wrist
   c) Left forearm
   d) Left wrist

16. All patients with orthopedic injuries should have the five Ps assessed and documented. What are the five Ps?
   a) **Pain**
   b) **Pallor**
   c) **Pulselessness**
   d) **Paresthesia**
   e) **Paralysis**

17. List the contraindications for implementing orders for extremity x-rays.
   a) **Deficits in circulation, sensation, movement distal to injury**
   b) Open fracture
   c) Pregnancy

Medications

1. Which of the following statements concerning the administration of Acetaminophen or Ibuprofen for adult pain relief are correct:
   a) Indicated for patients triaged as CTAS 3-5 with mild to moderate pain (≤ 7/10 on pain scale)
   b) Pain is related to headache, dental, ear, nose and throat, MSK or skin injuries
   c) The primary clinical contraindication is abdominal pain
   d) A single dose of Acetaminophen 975-1000mg or Ibuprofen 600mg can be administered either po or pr. If the preceding dose has been one of these medications, the other may be administered
   e) **All of the above**

2. 32 year old Luke Razorblade presents to the ED complaining of a general body aches, sore throat and fever. His temperature at triage is 38.1°C. Given that he has not taken anything for the fever you can administer either Acetaminophen 975-1000mg or Ibuprofen 600mg po.
   a) **True**
   b) False
3. 66 year old Dennis Denial is triaged as CTAS 2 with chest pain (cardiac features) even though he insists it’s just terrible indigestion from his Taco Bell lunch. One of the first drugs you will want to administer will be ASA 160 mg chewable tablet x 1 dose.
   a) True
   b) False

4. A 19 year old female with a history of substance abuse arrives by ambulance with an altered LOC and a blood sugar of 5mmol/L. Once you have established IV access you can administer the following:
   a) Dextrose 50% (25 gms in 50mLs) IV push over 2-3 minutes
   b) Narcan 0.4mg iv push
   c) None of the above

5. In assessing a patient with a full thickness laceration, which of the following statements concerning the medical directive for tetanus immunization are correct:
   a) Presents with an open injury to the skin or eye
   b) Has completed their primary immunization and hasn’t received a booster in 10 years
   c) Td (Tetanus and Diptheria Toxoid) 1.0mLs is given IM in the deltoid muscle
   d) A and B only
   e) All of the above

6. 54 year old Drew Maurier presents to the ED with shortness of breath, B/P 150/92, P 110, RR 24, T 36.4, O2 Sat. 93%. In addition to burning the pack of cigarettes hanging out of his top pocket you want to administer Salbutamol (Ventolin) four to eight puffs or give a wet neb masking (5mg/mL);1 mL in 3 mL saline to assist his SOB. Before you can do this however, the following criteria must be met:
   a) Patient has a history of asthma or COPD
   b) Presents with dysphagia, muffled voice stridor and other symptoms of upper airway compromise
   c) Presents with any of the following symptoms- cough, presence of respiratory distress, wheeze, tightness or decreased breath sounds on chest auscultation
   d) All of the above
   e) A and C only

7. Any male patient requiring urinary catheterization who is not allergic to amide anaesthetics or has a known urethral trauma or structural abnormality can receive:

   Lidocaine Jelly 2% 200mg/10mLs inserted into the urethra five minutes prior to catheterization to assist in comfort and ease of catheter passage
   a) True
   b) False
8. LET (Lidocaine, Epinephrine and Tetracaine) topical anaesthetic compound can be applied 30 minutes prior to suturing in which of the following situations:
   a) Lacerations to the ear, nose, fingers and toes
   b) Only lacerations that do not extend beyond the dermis
   c) Lacerations that extend beyond the dermis to underlying structures (e.g. bone, cartilage, tendon, etc)
   d) A and C
   e) B only

9. 33 year old Sam Squint arrives from his workplace after splashing a chemical in his left eye and is in a fair amount of discomfort. You have a medical directive to instill two drops of ophthalmic anesthetic eye drops to Sam’s eye. Does he meet the criteria?
   a) Yes
   b) No

Another criteria for instilling these eye drops is in anticipation of Morgan Lens insertion pain caused by foreign body, thermal injury or corneal abrasion.
   a) True
   b) False

**Therapeutic Procedures**

1. A saline lock or peripheral IV may be initiated under the following circumstances:
   a) Signs of airway compromise, respiratory distress, shock, dehydration, bleeding, or altered LOC
   b) In anticipation of IV medication administration for pain control, blood and blood product administration, or to provide IV hydration
   c) History of high risk mechanism of injury, infection in the immunocompromised patients, or overdose
   d) A and B only
   e) All of the above

2. Oxygen therapy may be initiated when the patient presents with signs and symptoms of one or more of the following:
   a) Respiratory distress, SaO2 < 92%, or below established desirable range for the individual patient
   b) Hemodynamic instability
   c) There is evidence of suspected hypoxemia (chest pain, tachycardia, hemorrhage, hypovolemia, sickle cell, altered LOC, Trauma, smoke and/or toxin inhalation
   d) B and C only
   e) All the above
3. Eye irrigation with 1 L NS or Ringer’s Lactate may NOT be initiated when…
   a) Penetrating eye trauma, foreign body in the eye, or signs and symptoms of ruptured
globe (obvious bleeding)
   b) The patient has flushed prior to arriving in the ED
   c) The patient has not had topical ophthalmic anaesthetic instilled
   d) The patient has suffered a flash burn
   e) All of the above.

Paediatrics

1. Salbutamol and Ipratropium Bromide may be administered to a child when...
   a) The child presents with signs and symptoms of upper airway pathology e.g. stridor, drooling,
muffled voice, dysphagia
   b) The child presents with audible wheezing, wheezing with retractions, spasmodic
   cough, dyspnea, tachypnea, or decreased air entry into lung fields on auscultation with
   a history of reactive airway disease
   c) The child presents with audible wheezing, wheezing with retractions, spasmodic cough,
dyspnea, tachypnea, or decreased air entry into lung fields on auscultation with NO history of
   reactive airway disease
   d) A and B only
   e) None of the above

2. A three month old child was given a therapeutic dose of Acetaminophen two hours prior to arriving in the
   ED. The child’s temperature is 38.5°C. I can administer Ibuprofen as per the mother’s request.
   a) True
   b) False

3. The child presents to the ED with vomiting and diarrhea. The 20 kg child was started on ORT on arrival.
The mother advises you the child vomited a moderate amount of emesis and just had a large watery stool
while waiting in the room. You should:
   a) Stop giving the ORT as it will precipitate more vomiting
   b) The child should receive 1000 cc of ORT slowly and then be given 200cc ORT for every
   loose stool
   c) The child should receive 500 cc of ORT slowly and then be given 100 cc ORT for every loose
   stool
   d) We should start an IV to rehydrate the child
   e) None of the above
### SAMPLE

Section 1.01 Extremity X-ray Medical Directive

Section 1.02 Certification Record

<table>
<thead>
<tr>
<th>Name: ______________________________</th>
<th>Date Submitted to Clinical Educator: ______________________________</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>Extremity</th>
<th>Meets Expectations</th>
<th>Needs Improvement</th>
<th>Comments</th>
<th>Physician/Designate Signature</th>
</tr>
</thead>
<tbody>
<tr>
<td>Article I.</td>
<td>COMPLETED WRITTEN QUIZ</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Achieved 100% on written quiz</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Article II.</td>
<td>HAND/FINGERS</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Able to identify digits (label)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Orders the appropriate x-ray as per the patient’s injuries</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Article III.</td>
<td>WRIST/FOREARM</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Locates Scaphoid</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Orders the appropriate x-ray as per the patient’s injuries</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Article IV.</td>
<td>ELBOW</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Able to identify the olecranon and radial head</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Orders the appropriate x-ray as per the patient’s injuries</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Article V.</td>
<td>KNEE</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Demonstrates understanding of Ottawa Knee Rules</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Orders the appropriate x-ray as per the patient’s injuries</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Article VI.</td>
<td>ANKLE/ TIB-FIB</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Demonstrates understanding of Ottawa Ankle Rules</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Demonstrates the knowledge of indications for tib/fib x-rays</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Orders the appropriate x-ray as per the patient’s injuries</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Article VII.</td>
<td>FOOT/TOES</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Demonstrates understanding of Ottawa Ankle &amp; Foot Rules</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Orders the appropriate x-ray as per the patient’s injuries</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Article VIII.</td>
<td>DOCUMENTATION</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Documents assessment findings as per the Extremity X-ray Medical Directive Policy</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Implementer Performance Readiness Form - Individual

From: An Interprofessional Guide on Orders, Directives and Delegation; Federation of Health Regulatory Colleges of Ontario (January 2007)

(Name of Implementer)

has demonstrated performance readiness for implementing:

(Name of Directive, Delegation or Practice)

and is authorized to perform the procedure in accordance with the education program (if applicable) and relevant policies and procedures for the period:

and is authorized to teach in the education program: _________________________ (Yes or No)

Implementer ______________________ Signature ______________________ Date ____________

Authorizer or Educator ______________________ Signature ______________________ Date ____________
# Implementer Performance Readiness Form - Group

From: An Interprofessional Guide on Orders, Directives and Delegation; Federation of Health Regulatory Colleges of Ontario (January 2007)

Name and Number of Directive, Delegation or Practice: 

Unit/Area: ___________________________  # of pages: ____________

List Completed by:  
(Authorizer or Educator’s Name, Position, Signature and Initials)

Date Submitted: _________________  For Period: _________________

<table>
<thead>
<tr>
<th>Name of Implementer</th>
<th>Signature</th>
<th>Date</th>
<th>Authorizer or Educator’s Initials</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
MODULE 3

Prototype Directives
ED Prototype Medical Directives Module 3 of 6

Use these prototype directives to establish directives that suit your ED circumstances
### Table of Contents

Prototype ED Directives:

- Laboratory Test and Diagnostic Procedures Directive .......................... 1
- Diagnostic Imaging Directive ........................................................................ 11
- Medications Directive .............................................................................. 17
- Therapeutic Procedures Directive .............................................................. 26
- Paediatric Directive .................................................................................. 33

### Acknowledgements:

Prototype ED Medical Directive - Laboratory Test and Diagnostic Procedures

Title: ED Laboratory Test and Diagnostic Procedures Directive
Number: Assign # in accordance with hospital record-keeping policies

Activation Date: 
Review due by: 

Sponsoring/Contact Person(s)
(name, position, contact particulars):
Typically chief or physician, and nursing staff member most responsible for developing directive

Orders:

<table>
<thead>
<tr>
<th>Appendix Attached:</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
</table>

Title: Laboratory Test and Diagnostic Procedures Order Tables

Orders for laboratory test and diagnostic procedures as identified on the appended order tables:

1. Laboratory Test Order Table (p. 5)
   a) RBW
   b) Liver Profile (LFTs)
   c) Group and Screen
   d) Serum Quantitative HCG
   e) Urine Qualitative HCG
   f) Serum Drug and Alcohol Screen
   g) Cardiac Markers
   h) Serum Coagulopathy
   i) Blood Cultures
   j) Serum lactate
   k) POCT Urinalysis
   l) Urine for R and M

2. Diagnostic Procedures Order Table (p. 9)
   a) 12 Lead ECG
   b) Capillary Blood Glucose

Recipient Patients:

<table>
<thead>
<tr>
<th>Appendix Attached:</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
</table>

Any adult patient (specify age in accordance with hospital policy, e.g. >= 16) presenting to the ED prior to first contact with the attending physician who meets the conditions identified in this directive.

If patients have lab testing and diagnostic procedures administered under this directive but leave without being seen by the attending physician:
- Staff will forward the record of administered tests and procedures to the attending physician for disposition.

Authorized Implementers:

<table>
<thead>
<tr>
<th>Appendix Attached:</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
</table>

All ED nurses and designated staff who have successfully completed the relevant ED Medical Directive orientation program. It is recommended that hospitals maintain a list of authorized implementers as part of the Medical Directive record.

See Module 2 – Performance Readiness Plan for sample certification forms that may be used if desired to maintain a list of authorized implementers. Such forms may be appended to the directive, identifying them in the ‘Appendix.
Indications:

- Lab tests and diagnostic procedures will be administered for the period beginning from when a patient arrives in the ED to first contact with the attending physician, unless the attending physician explicitly orders implementation of the directive beyond that period.
- Prior to implementation of any directive, a patient assessment is completed in accordance with standards of practice and any applicable hospital policy. Allergies and sensitivities must be documented.
- Specific indications are identified in the appended Order Tables.

Definitions for indications used in the table:

1. **Acute Coronary Syndrome (ACS)** – as manifested by discomfort (pressure or pain, radiating or non-radiating, anterior or posterior) from jaw to umbilicus that may include any of the following:
   - Shortness of Breath (SOB)
   - Diaphoresis
   - Pallor
   - Nausea/vomiting
   - Dysrhythmias (palpitations, tachycardia, bradycardia)
   - Syncope
   - Weakness, lightheadedness, pre-syncope
   - Lethargy
2. **Fever** – Temperature $\geq 38^\circ C$
3. **Hypothermia** – Temperature $\leq 36^\circ C$
4. **Hemodynamic instability** – as manifested by one or more of the following signs of shock:
   - Pale
   - Diaphoretic
   - Tachypnic
   - Tachycardia
   - Hypotensive
   - Altered level of Consciousness (LOC)
5. **Immunocompromised** - Patients with one or more of the following:
   - On chemotherapy for cancer,
   - On anti-tnf medication – infliximab (e.g. Remicade),
   - Organ transplant(s),
   - Splenectomy,
   - Hiv,
   - Lupus, rheumatoid arthritis and other chronic inflammatory conditions,
   - Diabetes mellitus,
   - Chronic alcohol abuse,
   - Chronic corticosteroid therapy
6. **Major bleed** – any volume loss that causes hemodynamic instability resulting from possible GI bleed, ruptured aneurysm, ruptured spleen, femur fracture, ectopic pregnancy
7. **Major trauma** - high risk mechanism of injury

Contraindications:
See appended Order Tables.

Consent:

Staff implementing the directive will obtain consent in accordance with the Health Care Consent Act and any relevant hospital policies and procedures (note these).
### Guidelines for Implementing the Order / Procedure:

Blood specimens for diagnostic studies may be collected by:

1. Venipuncture,
2. Saline Lock, or
3. Accessing established Vascular Access Device (not including hemodialysis lines).

For specific guidelines, see appended Order Tables.

### Documentation and Communication:

Implementing staff will document the:

- Specific lab test and diagnostic procedure order in the order section of the patient record, noting the medical directive name and number and signing off the order with implementer’s name and signature as per the attending physician (when attending physician known). Ensure any requisition contains this information.
- Indications, implementation and patient response in accordance with any hospital record-keeping policies (note these)

Append any designated forms used to document implementation of this directive, identifying them in the ‘Appendix Attached’ section directly above. See Module 4 – Sample Prototype Directive Preprinted Orders and ED Record for examples of forms that may be used as desired.

Note: Clear and timely notification, communication and documentation between the nurse and the physician are critical to safe, proper use of a medical directive.

### Review and Quality Monitoring Guidelines:

Staff identifying any untoward or unintended outcomes arising from implementation of orders under this directive, or any issues identified with it will report these to (note to whom) as soon as possible for appropriate disposition. This does not include untoward or unintended outcomes or issues that are possible clinical sequelae regardless of whether a directive or direct order is used.

Additional provisions (e.g. for renewal and re-certification) may be identified here or in Performance Readiness Assessment form (Module 1)

### Administrative Approvals:

Identify the relevant approval committees and staff members, for example: Unit or Program Administrative Staff and Committees, Educators, Professional Leaders, Professional Councils, Medical Advisory Committee etc.; and ensure a signed copy of approvals is maintained as part of the Medical Directive record, either in this section, or in an attached appendix or on designated hospital-specific sign-off forms or see Module 6 – Sample Approval Forms for sign-off forms that may be used as desired.
### Approving Physician(s)/Authorizer(s):

Identify all the relevant approving attending physicians and ensure a signed copy of approvals is maintained with the Medical Directive record.

In addition, ensure there is an accessible and timely means of informing staff of new attendings (e.g. newly privileged physicians, locums, residents).

Signatures may be recorded in this section, either in this section, or in an attached appendix or on designated hospital-specific sign-off forms or see Module 6 – Sample Approval Forms for sign-off forms that may be used as desired.

### References

List any references here, as below, attach appendix, or list references in Performance Readiness Assessment form (Module 1).


Guidelines 2006 for cardiopulmonary and Emergency Cardiovascular care: American Heart Association

## 1. Laboratory Test Order Table

This table cannot be relied upon in the absence of: ED Laboratory Test and Diagnostic Procedures Directive (record directive number)

<table>
<thead>
<tr>
<th>Orders</th>
<th>Indications/Contraindications</th>
<th>Guidelines</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Bloodwork</strong></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
| a) Routine Blood Work – CBC, electrolytes (K, Na, Cl, total CO₂), urea, creatinine & glucose | Signs and symptoms including one or more of the following:  
- Cardiopulmonary condition (chest pain, SOB, weakness, dizziness, looks unwell),  
- Neurologic condition (alteration in neurologic status, decreased LOC, confusion, seizure, suspected stroke)  
- Syncope (> = 65 yrs)  
- Dehydration  
- Major trauma  
- Symptoms of significant blood loss/major bleeding  
- Moderate to severe abdominal pain  
- Signs and symptoms of infection - fever or hypothermia with tachypnea and tachycardia  
- Chemotherapy and immunocompromised patients  
- Chronic, diagnosed disease(s) with change in status or exacerbation in condition  
- Drug or alcohol ingestion | Customize the orders and procedures for collecting this specimen based on hospital-specific lab protocols |
# 1. Laboratory Test Order Table

This table cannot be relied upon in the absence of ED Laboratory Test and Diagnostic Procedures Directive (record directive number)

<table>
<thead>
<tr>
<th>Orders</th>
<th>Indications/Contraindications</th>
<th>Guidelines</th>
</tr>
</thead>
</table>
| b) Liver Profile (LFTs) – Amylase, AST, ALT, T-Bili, Direct Bilirubin, ALP, GGT, Albumin | Signs and symptoms of one or more of the following:  
   - Right upper quadrant or epigastric pain  
   - Blunt abdominal trauma  
   - Acetaminophen overdose (Consider ordering only AST as per hospital policy) | Customize the orders for collecting this specimen based on hospital-specific lab protocols |
| c) Group and Screen | Signs and symptoms of one or more of the following:  
   - Major trauma patients  
   - Patients with actual or potential hemodynamic instability  
   - Pallor, known or suspected anemia  
   - Recent chemotherapy patient who appears anemic  
   - Patients with suspected major bleed  
   - Female patients abdominal pain and/or vaginal bleeding who may be pregnant | Customize the procedures for collecting this specimen based on hospital-specific lab protocols |
| d) Serum Quantitative HCG  
Or  
e) Urine Qualitative HCG as per hospital policy | Women of childbearing age and capacity with signs and symptoms of one or more of the following:  
   - Abdominal pain  
   - Vaginal bleeding  
   - Hemodynamic instability  
   - Syncopal episode  
   - Major trauma patient  
   - Pregnancy suspected by patient or nurse (e.g. late menses, inexplicable weight gain) | Customize the orders for collecting this specimen based on hospital-specific lab protocols |
### 1. Laboratory Test Order Table

This table cannot be relied upon in the absence of: ED Laboratory Test and Diagnostic Procedures Directive (record directive number)

<table>
<thead>
<tr>
<th>Orders</th>
<th>Indications/Contraindications</th>
<th>Guidelines</th>
</tr>
</thead>
</table>
| f) Serum Drug and Alcohol Screen as per hospital policy, e.g.  
  - Acetaminophen  
  - ASA  
  - ETOH  
  - Lithium  
  - Digoxin  
  - Carbamazepine  
  - Phenytoin  
  - Valproic Acid  

Patients with actual or suspected drug ingestion/overdose with one or more of the following:  
  - Reported ingestion  
  - Alterations in mood, behaviour, and/or motor function suggestive of intoxication  
  - Alteration in LOC  
  - New onset seizure  
  - Major trauma  
  - Concern about non-therapeutic drug levels (carbamazepine, phenytoin, lithium, valproic acid)  

Customize the orders for collecting this specimen based on hospital-specific lab protocols. |
| g) (Identify hospital-specific markers) Cardiac Markers: CK, Troponin  

Signs and symptoms of Acute Coronary Syndrome  

Customize the orders for this specimen collection based on hospital-specific lab protocols. |
| h) Serum Coagulopathy: INR/PTT  

Signs and symptoms including one or more of the following:  
  - Acute coronary syndrome  
  - Current warfarin use  
  - Possible pulmonary embolus (SOB, tachycardia, hypoxia, risk factors such as recent surgery, immobility)  
  - Possible stroke/TIA (weakness, numbness, trouble speaking, vision problems, headache, dizziness)  
  - Actual or potential hemodynamic instability  

Customize the procedures for this specimen collection based on hospital-specific lab protocols. |
# 1. Laboratory Test Order Table

This table cannot be relied upon in the absence of: ED Laboratory Test and Diagnostic Procedures Directive (record directive number)

<table>
<thead>
<tr>
<th>Orders</th>
<th>Indications/Contraindications</th>
<th>Guidelines</th>
</tr>
</thead>
</table>
| i) Blood Cultures           | Patients with actual or suspected infection (localized or generalized) with temperature $\geq 38$ or $\leq 36$ who presents with any of the following signs and symptoms:  
• Tachypnea $> 20$ RPM  
• Tachycardia HR $> 90$ BPM  
• Altered LOC  
• Immunocompromised patients  
• Patients with indwelling vascular access devices | Collection standard: 2 sets of cultures 10 minutes apart  
Customize the procedures for this specimen collection based on hospital-specific lab protocols.                                                                                                                      |
| j) Serum Lactate            | Patients with suspected infection who meets SIRS criteria, i.e. looks unwell with 2 or more of the following:  
• Fever (temperature $> 38$ C) or hypothermia (temperature $< 36$ C)  
• Tachypnea $> 20$ RPM  
• Tachycardia HR $> 90$ BPM  
• Chemo or immunocompromised patients | • Reference: SIRS Criteria                                                                                                                                                                                      |
| Urinalysis                  |                                                                                                                                                                                                                               |                                                                                                                                                                                                         |
| k) Point of Care Test (POCT) | Patients presenting with one or more of the following signs and symptoms:  
• Abdominal pain  
• UTI symptoms  
• Patients $\geq 65$ years of age with fever or hypothermia, confusion, delirium, or new onset weakness |                                                                                                                                                                                                         |
### 1. Laboratory Test Order Table

This table cannot be relied upon in the absence of: ED Laboratory Test and Diagnostic Procedures Directive (record directive number)

<table>
<thead>
<tr>
<th>Orders</th>
<th>Indications/Contraindications</th>
<th>Guidelines</th>
</tr>
</thead>
</table>
| I) Urine for R and M | If POCT urinalysis is positive for:  
- Leukocytes  
- Proteins  
- Blood  
- Nitrates  
- Glucose | Follow applicable hospital policy and procedure |

### 2. Diagnostic Procedures Order Table

This table cannot be relied upon in the absence of: ED Laboratory Test and Diagnostic Procedures Directive (record directive number)

<table>
<thead>
<tr>
<th>Orders</th>
<th>Indications/Contraindications</th>
<th>Guidelines</th>
</tr>
</thead>
</table>
| a) 12 Lead ECG  | Signs and symptoms of, or including one or more of the following:  
- Respiratory pathology  
- Cardiac pathology  
- Cerebrovascular accident  
- Metabolic imbalance  
- Trauma to chest wall  
- Major or multiple trauma  
- Possible toxic ingestion  
- Possible adverse reaction to medications or drugs  
- Weakness or lightheadedness in those > 65  
- Electrical injury | Follow applicable hospital policy and procedure |
## 2. Diagnostic Procedures Order Table

This table cannot be relied upon in the absence of: ED Laboratory Test and Diagnostic Procedures Directive (record directive number)

<table>
<thead>
<tr>
<th>Orders</th>
<th>Indications/Contraindications</th>
<th>Guidelines</th>
</tr>
</thead>
</table>
| b) Capillary Blood Glucose  | To determine baseline glucose status in patients with a diagnosis of diabetes mellitus and for signs and symptoms suggestive of hypoglycemia or hyperglycemia, including one or more of the following:  
  - Altered LOC  
  - Confusion, agitation, behavioural changes  
  - Recent or active seizure  
  - Suspicion of alcohol ingestion  
  - Syncopal event  
  - Listless, lethargic, fatigued | Follow applicable hospital policy and procedure                                                                                                                        |
Prototype ED Medical Directive - Diagnostic Imaging

**Title:** ED Diagnostic Imaging Directive  
**Number:** Assign # in accordance with hospital record-keeping policies

**Activation Date:**  
**Review due by:**

**Sponsoring/Contact Person(s):** Typically chief or physician, and nursing staff member most responsible for developing directive

### Orders:

**Appendix Attached:** Yes  No  
**Title:** Extremity Plain Film Radiography X-ray Order Table

Orders for diagnostic imaging as identified on the appended order table:

1. **Extremity Plain Film Radiography X-ray Order Table (p. 15)**
   - a) Foot x-ray
   - b) Ankle x-ray
   - c) Knee x-ray
   - d) Tibia/Fibula x-ray
   - e) Hand x-ray
   - f) Finger(s) x-ray
   - g) Scaphoid x-ray
   - h) Wrist x-ray
   - i) Forearm x-ray
   - j) Elbow x-ray

**Recipient Patients:**

Any adult patient (specify age in accordance with hospital policy, e.g. >/= 16) presenting to the ED prior to first contact with the attending physician who meets the conditions identified in this directive.

If patients have plain film radiography studies administered under this directive but leave without being seen by the attending physician:

- Staff will forward the record of studies administered to the attending physician for disposition.

**Authorized Implementers:**

All ED nurses and designated staff who have successfully completed the relevant ED Medical Directive orientation program. It is recommended that hospitals maintain a list of authorized implementers as part of the Medical Directive record.

See **Module 2** – Performance Readiness Plan for sample certification forms that may be used if desired to maintain a list of authorized implementers. Such forms may be appended to the directive, identifying them in the 'Appendix Attached' section directly above, or maintain them in accordance with hospital record-keeping policies.
### Indications:
- Plain film radiography x-ray orders will be implemented for the period beginning when a patient arrives in the ED to first contact with the attending physician, unless the attending physician explicitly orders implementation of this directive beyond that period.
- Plain film radiography x-rays will only be ordered under this directive if patients have received trauma to the affected area, and have pain referable to that area.
- Prior to implementation of any directive, a patient assessment is completed in accordance with standards of practice and any applicable hospital policy. Allergies and sensitivities must be documented.
- Prior to a study being requisitioned, ensure:
  - Neurological and orthopedic status are assessed and documented
  - Any jewellery or constricting clothing is atraumatically removed

Specific indications are identified in the appended Order Table.

### Contraindications:
1. Known or suspected pregnancy.
2. Unstable patient – physician to be contacted immediately.
3. Signs or symptoms of neurovascular compromise in the affected limb – physician to be contacted immediately.
4. Open fractures will be assessed by the emergency physician directly.
5. Patient is intoxicated or has other distracting injuries and is unable to follow direction, maintain motor control or is uncooperative

See appended Order Table for further specific contraindications.

### Consent:
- Staff implementing the directive will obtain consent in accordance with the *Health Care Consent Act* and any relevant hospital policies and procedures (note these).

### Guidelines for Implementing the Order / Procedure:
See appended Order Table.
### Documentation and Communication:

Implementing staff document the:

- Specific x-ray order in the order section of the patient record, noting the medical directive name and number, and signing off the order with implementer’s name and signature as per the attending physician (when attending physician known). Ensure any requisition contains this information.

- Indications, implementation and patient response in accordance with any hospital record-keeping policies (note these)

Append any designated forms used to document implementation of this directive, identifying them in the ‘Appendix Attached’ section directly above. See Module 4 – Sample Prototype Directive Preprinted Orders and ED Record for examples of forms that may be used as desired.

Note: Clear and timely notification, communication and documentation between the nurse and the physician are critical to safe, proper use of a medical directive.

### Review and Quality Monitoring Guidelines:

Staff identifying any untoward or unintended outcomes arising from implementation of orders under this directive, or any issues identified with it will report these to (note to whom) as soon as possible for appropriate disposition. This does not include untoward or unintended outcomes or issues that are possible clinical sequelae regardless of whether a directive or direct order is used.

Additional provisions (e.g. for renewal and re-certification) may be identified here or in Performance Readiness Assessment form (Module 1)

### Administrative Approvals:

Identify the relevant approval committees and staff members, for example: Unit or Program Administrative Staff and Committees, Educators, Professional Leaders, Professional Councils, Medical Advisory Committee etc.; and ensure a signed copy of approvals is maintained as part of the Medical Directive record.

Signatures may be recorded either in this section, or in an attached appendix, or on designated hospital-specific sign-off forms, or see Module 6 – Sample Approval Forms for sign-off forms that may be used as desired.

### Approving Physician(s)/Authorizer(s):

Identify all the relevant approving attending physicians and ensure a signed copy of approvals is maintained with the Medical Directive record.

In addition, ensure there is an accessible and timely means of informing staff of new attendings (e.g. newly privileged physicians, locums, residents).

Signatures may be recorded either in this section, or in an attached appendix, or on designated hospital-specific sign-off forms, or see Module 6 – Sample Approval Forms for sign-off forms that may be used as desired.
References

List any references here as below and attach appendix, or list references in Performance Readiness Assessment form (Module 1).


2. Stiell IG et al, “Decision Rules for the Use of Radiography in Acute Ankle Injuries - Refinement and Prospective Validation”, JAMA 1993 269(9); 1127-1132.


11. Stiell IG et al “Prospective Validation fo a Decision Rule for the Use of radiography in Acute Knee Injuries” JAMA Feb 1996 275(8) 611-615.


## Extremity Plain Film Radiography X-Ray Orders Table

This table cannot be relied upon in the absence of ED Diagnostic Imaging Directive (record directive number)

<table>
<thead>
<tr>
<th>Orders</th>
<th>Indications</th>
<th>Contraindications</th>
<th>Guidelines</th>
</tr>
</thead>
<tbody>
<tr>
<td>a) Foot x-ray</td>
<td>Pain in the midfoot zone and one or more of the following findings:</td>
<td>1. Known or suspected pregnancy</td>
<td>See Ottawa Ankle Rules Protocol (record version and maintain as part of medical directive record)</td>
</tr>
<tr>
<td></td>
<td>1. Bone tenderness at the base of the 5th metatarsal</td>
<td>2. Unstable patient – physician to be contacted immediately</td>
<td></td>
</tr>
<tr>
<td></td>
<td>2. Bone tenderness at the navicular</td>
<td>3. Signs or symptoms of neurovascular compromise in the affected limb physician to be contacted immediately</td>
<td></td>
</tr>
<tr>
<td></td>
<td>3. Inability to bear weight both immediately and in the emergency department</td>
<td>4. Open fractures will be assessed by the emergency physician directly</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>5. Patient is intoxicated or has other distracting injuries and is unable to follow direction, maintain motor control or is uncooperative.</td>
<td></td>
</tr>
<tr>
<td>b) Ankle x-ray</td>
<td>Upon palpation of the entire distal 6 cm of the fibula and tibia to assess the malleolar zone, pain in the malleolar zone and one or more of the following:</td>
<td>• As above</td>
<td></td>
</tr>
<tr>
<td></td>
<td>1. Bone tenderness at the posterior edge or tip of the lateral malleolus</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>2. Bone tenderness at the posterior edge or the tip of the medial malleolus</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>3. Inability to bear weight both immediately and in the emergency department</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
## 1. Extremity Plain Film Radiography X-Ray Orders Table

This table cannot be relied upon in the absence of ED Diagnostic Imaging Directive (record directive number).

<table>
<thead>
<tr>
<th>Orders (Specify exam name and views that correspond to hospital-specific policy)</th>
<th>Indications</th>
<th>Contraindications</th>
<th>Guidelines</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>c) Knee x-ray</strong></td>
<td>Knee injury patients with one or more of these findings: 1. Isolated tenderness of the patella (that is, no bone tenderness of the knee other than the patella) 2. Tenderness at the head of the fibula 3. Inability to flex to 90 degrees 4. Inability to bear weight both immediately and in the emergency department (4 steps; unable to transfer weight twice onto each lower limb regardless of limping)</td>
<td>• As above</td>
<td>See Ottawa Knee Rules Protocol (record version and maintain as part of medical directive record)</td>
</tr>
<tr>
<td><strong>d) Tibia/Fibula x-ray</strong></td>
<td>Deformity and or swelling in the area</td>
<td>• As above</td>
<td></td>
</tr>
<tr>
<td><strong>e) Hand x-ray</strong></td>
<td>Deformity and or swelling in the area, impaired range of motion and localized bony tenderness</td>
<td>• As above</td>
<td></td>
</tr>
<tr>
<td><strong>f) Finger(s) x-ray</strong></td>
<td>Isolated finger injury distal to the MCP joint</td>
<td>• As above</td>
<td></td>
</tr>
<tr>
<td><strong>g) Scaphoid x-ray</strong></td>
<td>Pain at the anatomic “snuff box” Pain on axial compression of thumb</td>
<td>• As above</td>
<td></td>
</tr>
<tr>
<td><strong>h) Wrist x-ray</strong></td>
<td>Deformity and swelling in the area, impaired range of motion and localized bony tenderness</td>
<td>• As above</td>
<td></td>
</tr>
<tr>
<td><strong>i) Forearm x-ray</strong></td>
<td>Deformity and swelling in the area, localized bony tenderness</td>
<td>• As above</td>
<td></td>
</tr>
<tr>
<td><strong>j) Elbow x-ray</strong></td>
<td>Deformity and or joint effusion or swelling in the area, impaired range of motion and localized bony tenderness</td>
<td>• As above</td>
<td></td>
</tr>
</tbody>
</table>
**Prototype ED Medical Directive - Medications**

<table>
<thead>
<tr>
<th>Title:</th>
<th>ED Medication Directive</th>
<th>Number:</th>
<th>Assign # in accordance with hospital record-keeping policies</th>
</tr>
</thead>
<tbody>
<tr>
<td>Activation Date:</td>
<td></td>
<td>Review due by:</td>
<td></td>
</tr>
<tr>
<td>Sponsoring/Contact Person(s):</td>
<td>Typically chief or physician, and nursing staff member most responsible for developing directive</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Orders:**

- Appendix Attached: ☑ Yes ☐ No
- Title: Medication Order Table

Orders as identified on the appended order table:

1. **Medication Order Table (p. 21)**
   - a) Acetaminophen - Pain
   - b) Ibuprofen - Pain
   - c) Acetaminophen – Fever
   - d) Ibuprofen - Fever
   - e) Acetylsalicylic Acid (ASA) – Acute Coronary Syndrome
   - f) Dextrose 50% IV direct injection
   - g) Tetanus and Diptheria Toxoid
   - h) Salbutamol
   - i) Lidocaine Jelly 2%
   - j) Lidocaine, Epinephrine and Tetracaine Compound (LET)
   - k) Ophthalnic Anaesthetic Eye Drops

**Recipient Patients:**

- Appendix Attached: ☑ Yes ☐ No
- Title: 

Any adult patient (specify age in accordance with hospital policy, e.g. >/= 16) presenting to the ED prior to first contact with the attending physician who meets the conditions identified in this directive.

If patients have medications administered under this directive but leave before being seen by the attending physician:
- Staff will forward the patient record, including the record of administered medications to the attending physician for disposition.

**Authorized Implementers:**

- Appendix Attached: ☑ Yes ☐ No
- Title: 

All ED nurses and designated staff who have successfully completed the relevant ED Medical Directive orientation program. It is recommended that hospitals maintain a list of authorized implementers as part of the Medical Directive record.

See **Module 2 – Performance Readiness Plan** for sample certification forms that may be used if desired to maintain a list of authorized implementers. Such forms may be appended to the directive, identifying them in the ‘Appendix Attached’ section directly above, or maintain them in accordance with hospital record-keeping policies.
Indications:

- Medications will be administered from the period beginning from when a patient arrives in the ED to first contact with the attending physician, unless the attending physician specifically orders implementation of the directive beyond that period.
- Prior to implementation of any directive, a patient assessment is completed in accordance with standards of practice and any applicable hospital policy. Allergies and sensitivities must be documented.
- Specific indications are identified in the appended Order Table.

Definitions for indications used in the table:

1. **Acute Coronary Syndrome (ACS)** – as manifested by discomfort (pressure or pain, radiating or non-radiating, anterior or posterior) from jaw to umbilicus that may include any of the following:
   - Shortness of Breath (SOB)
   - Diaphoresis
   - Pallor
   - Nausea/vomiting
   - Dysrhythmias (palpitations, tachycardia, bradycardia)
   - Syncope
   - Weakness, lightheadedness, pre-syncope
   - Lethargy

2. **Fever** – Temperature ≥ 38°C

3. **Hypothermia** – Temperature ≤ 36°C

4. **Hemodynamic instability** – as manifested by one or more of the following signs of shock:
   - Tachypnic
   - Tachycardia
   - Hypotensive
   - Altered level of consciousness (loc)
   - Pale
   - Diaphoretic

5. **Immunocompromised** - Patients with one or more of the following:
   - On chemotherapy for cancer,
   - On anti-tfn medication – infliximab (e.g. Remicade)
   - Organ transplant(s)
   - Splenectomy
   - Hiv
   - Lupus, rheumatoid arthritis and other chronic inflammatory conditions
   - Diabetes mellitus
   - Chronic alcohol abuse
   - Chronic corticosteroid therapy

6. **Major bleed** – any volume loss that causes hemodynamic instability resulting from possible GI bleed, ruptured aneurysm, ruptured spleen, femur fracture, or ectopic pregnancy

7. **Major trauma** - high risk mechanism of injury

Contraindications:
See appended Order Table.
Module 3: Prototype Directives

Consent:
Appendix Attached: ☐ Yes ☒ No
Title:

Staff implementing the directive will obtain consent in accordance with the Health Care Consent Act and any relevant hospital policies and procedures (Note these).

Guidelines for Implementing the Order / Procedure:
Appendix Attached: ☒ Yes ☐ No
Title: Medication Order Table

See appended Order Table.

Documentation and Communication:
Appendix Attached: ☐ Yes ☒ No
Title:

Implementing staff will document the:

• Medication order (including name of medication, dose, route, time of administration) in the order section of the patient record, noting the medical directive name and number, signing off the order as per the attending physician (when attending physician known)

• Indications, administration, and patient response in accordance with any hospital record-keeping policies (note these)

Append any designated forms used to document implementation of this directive, identifying them in the 'Appendix Attached' section directly above. See Module 4 – Sample Prototype Directive Pre-printed Orders and ED Record for examples of forms that may be used as desired.

Note: Clear and timely notification, communication and documentation between the nurse and the physician are critical to safe, proper use of a medical directive.

Review and Quality Monitoring Guidelines:
Appendix Attached: ☐ Yes ☒ No
Title:

Staff identifying any untoward or unintended outcomes arising from implementation of orders under this directive, or any issues identified with it will report these to (note to whom) as soon as possible for appropriate disposition. This does not include untoward or unintended outcomes or issues that are possible clinical sequelae regardless of whether a directive or direct order is used.

Additional provisions (e.g. for renewal and re-certification) may be identified here or in Performance Readiness Assessment form (Module 1).

Administrative Approvals:
Appendix Attached: ☐ Yes ☒ No
Title:

Identify the relevant approval committees and staff members, for example: Unit or Program Administrative Staff and Committees, Educators, Professional Leaders, Professional Councils, Medical Advisory Committee etc., and ensure a signed copy of approvals is maintained as part of the Medical Directive record.

Signatures may be recorded either in this section, or in an attached appendix, or on designated hospital-specific sign-off forms, or see Module 6 – Sample Approval Forms for sign-off forms that may be used as desired.
### Approving Physician(s)/Authorizer(s):

<table>
<thead>
<tr>
<th>Appendix Attached:</th>
<th>Yes</th>
<th>No</th>
<th>Title:</th>
</tr>
</thead>
</table>

Identify all the relevant approving attending physicians and ensure a signed copy of approvals is maintained with the Medical Directive record.

In addition, ensure there is an accessible and timely means of informing staff of new attendings (e.g. newly privileged physicians, locums, residents).

Signatures may be recorded either in this section, or in an attached appendix, or on designated hospital-specific sign-off forms, or see Module 6 – Sample Approval Forms for sign-off forms that may be used as desired.

### References

<table>
<thead>
<tr>
<th>Appendix Attached:</th>
<th>Yes</th>
<th>No</th>
<th>Title:</th>
</tr>
</thead>
</table>

List any references here as below and attach appendix or list references in Performance Readiness Assessment form Module 1.

- Guidelines 2006 for cardiopulmonary and Emergency Cardiovascular care: American Heart Association
- Compendium of Pharmaceuticals and Specialties (CPS) 2007
- Lacy,C: Armstrong,L;Lance, L Drug Information Handbook 2004
- Jovey, Roman Managing Pain: The Canadian Healthcare Professional Reference: The Canadian Pain Society 2002
# Appendix

Medication Order Table

<table>
<thead>
<tr>
<th>Orders</th>
<th>Indications</th>
<th>Contraindications</th>
<th>Guidelines</th>
</tr>
</thead>
</table>
| a) Acetaminophen (975-1000 mg – as per formulary) PO or PR (if unable to take PO) x 1 dose - Pain or As per patient choice, or hospital formulary | Patients triaged as CTAS 3, 4 or 5 with mild-moderate pain (=/< 7/10 on pain scale) as follows:  
  - Headache pain  
  - Dental pain  
  - Ear, nose and/or throat pain  
  - Musculoskeletal pain  
  - Skin pain | Acetaminophen:  
  - Abdominal pain  
  - Allergy or sensitivity to acetaminophen,  
  - Ingestion of acetaminophen in last 4 hours  
  - Hepatitis, liver disease, intoxicated patients | Reassess patient and document patient response within 30-60 minutes of administration or as indicated |
| b) Ibuprofen 600 mg PO x 1 - Pain   |                                                                                  | Ibuprofen:  
  - Abdominal pain  
  - History of liver or renal disease  
  - Allergy or sensitivity to Ibuprofen or Non-Steroidal Anti-Inflammatory Drugs (NSAIDs)  
  - History of Inflammatory Bowel Disease, Peptic ulcer, GI bleed  
  - Ingestion of ibuprofen within last 6 hours  
  - Nursing and pregnancy |                                                                                  |                                                                                  |

If a preceding dose of one of these medications has been taken by the patient, then the other may be administered as indicated under this directive.

**Note:** Do not give medications by mouth in cases where patient is obviously going to require surgery and consult physician for pain management.
### 1. Medication Order Table

This table cannot be relied upon in the absence of ED Medication Directive (record directive number)

<table>
<thead>
<tr>
<th>Orders</th>
<th>Indications</th>
<th>Contraindications</th>
<th>Guidelines</th>
</tr>
</thead>
</table>
| c) Acetaminophen (975-1000 mg – as per formulary) PO or PR (if unable to take PO) x 1 dose - Fever Or As per patient choice, or hospital formulary | • Patients with fever

**Note:** Do not give PO meds in cases where patient is obviously going to require surgery and consult physician for fever management.

| d) Ibuprofen 600 mg PO x 1 - Fever                                      | • As above for Acetaminophen and Ibuprofen – Pain     | • Allergy to ASA or NSAIDs
• Ingestion of ASA in past 4 hours
• Active upper GI bleed | Reassess patient and document patient response within 30-60 minutes of administration or as indicated. |
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>e) Acetylsalicylic Acid (ASA) chewable 160 mg x1 dose</td>
<td>• Symptoms of acute coronary syndrome</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
| f) Dextrose 50% (25 gms in 50 mLs) IV over 2-3 minutes                 | • Patient who has an altered LOC with blood sugar \(\leq 4\) mmol/L
• See ED Therapeutic Procedure Medical Directive for order to initiate venous access | • None where indications present | • Patient and blood glucose reassessment and documentation as indicated |
| g) Tetanus and Diptheria Toxoid (Td) 0.5 mLs IM, deltoid muscle or Identify compound, dosage and administration in accordance with hospital formulary/policy | • Any adult patient presenting with any open injury to the skin or eye who has completed primary immunization and has not had a booster in 10 yrs | • Allergy or sensitivity to Td
• Incomplete primary immunization | May consider monitoring for adverse reactions after administration in accordance with hospital policies |
## 1. Medication Order Table

This table cannot be relied upon in the absence of ED Medication Directive (record directive number)

<table>
<thead>
<tr>
<th>Orders</th>
<th>Indications</th>
<th>Contraindications</th>
<th>Guidelines</th>
</tr>
</thead>
<tbody>
<tr>
<td>h) As per hospital formulary/policy:</td>
<td>Patients presenting with SOB and a history of asthma or Chronic Obstructive Pulmonary Disease (COPD) with one or more of the following symptoms:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Salbutamol (Ventolin) 100 mcg/puff; 4-8 puffs by metered dose inhaler x 1</td>
<td>• Cough</td>
<td>• Allergy or sensitivity to salbutamol or adrenergic amines (salmeteral, terbutaline, albuterol, fomerterol)</td>
<td>• Reassess and document patient response and vital signs within 15 minutes following administration or as indicated</td>
</tr>
<tr>
<td>Or</td>
<td>• Presence of respiratory distress, wheeze, tightness or decreased breath sounds during chest auscultation</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Salbutamol (Ventolin) 5 mg/mL; 1 mL in 3 mL of saline via wet nebulizer mask over 10 minutes x1</td>
<td>Carry out febrile respiratory illness screening on all patients. If screen is positive, isolate patient prior to using nebulizer.</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td><strong>Note:</strong> For patients with severe respiratory distress, give medication and notify physician STAT</td>
<td></td>
<td></td>
</tr>
<tr>
<td>i) Lidocaine Jelly 2%, 200 mg/10mLs approximately 5 minutes prior to male urinary catheter insertion. May repeat x1 to max of 400 mgs.</td>
<td>Any male requiring urinary catheterization who is not allergic to amide anaesthetics</td>
<td>• Allergy to lidocaine or other amine-type topical anaesthetics (e.g. articaine, bupivacaine, prilocaine, mepivacaine, ropivacaine)</td>
<td></td>
</tr>
<tr>
<td>j) Lidocaine, Epinephrine and Tetracaine Compound (LET) as topical anaesthetic applied to cover lacerations 30 minutes prior to suturing. (Identify compound, dosage and application technique in accordance with hospital formulary)</td>
<td>Lacerations requiring suturing that do not extend beyond dermis</td>
<td>• Urethral trauma or known structural abnormality</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Laceration to ear, nose, fingers, toes, penis, no mucous membranes</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Significant injury to underlying structure (bone, cartilage, tendons, nerve and vessels) beneath the dermis</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Known allergy or hypersensitivity to any compound components</td>
<td></td>
</tr>
</tbody>
</table>
### 1. Medication Order Table

This table cannot be relied upon in the absence of ED Medication Directive (record directive number)

<table>
<thead>
<tr>
<th>Orders</th>
<th>Indications</th>
<th>Contraindications</th>
<th>Guidelines</th>
</tr>
</thead>
</table>
| k) Ophthalmic Anaesthetic Eye Drops to affected eye x 1 | • In anticipation of Morgan Lens insertion  
• For pain caused by any of the following:  
  • Foreign body  
  • Chemical splash  
  • Thermal injury  
  • Corneal abrasion | • Penetrating injury  
• Known allergy to ester-type agents (amethocaine, benzocaine, proparacaine, chloroprocaine, cocaine, procaine)  
• Known allergy or sensitivity to other eye drop ingredients | |
Prototype ED Medical Directive - Therapeutic Procedures

Title: ED Therapeutic Procedures Directive
Number: Assign # in accordance with hospital record-keeping policies

Activation Date: Review due by:

Sponsoring/Contact Person(s)
(name, position, contact particulars): Typically chief or physician, and nursing staff member most responsible for developing directive

Orders:

Orders for therapeutic procedures as identified on the appended:

1. Therapeutic Procedures Order Table (p. 29)
   a) Saline Lock Insertion,
   b) Peripheral IV Access with NS 30 mLs/hour,
   c) Accessing Established Vascular Access Device with NS 30 mLs/hour,
   d) Initiate and Titrate Oxygen by mask or nasal prongs to maintain \( \text{SaO}_2 \geq 95\% \),
   e) Urinary Catheter Insertion,
   f) Eye irrigation with 1 L NS with or without Insertion of Morgan Lens, or
   g) Spinal Backboard Removal.

Recipient Patients:

Any adult patient (specify age in accordance with hospital policy, e.g. \( \geq 16 \)) presenting to the ED prior to first contact with the attending physician who meets the conditions identified in this directive.

If patients have therapeutic procedures administered under this directive but leave before being seen by the attending physician:

- Staff will forward the patient record, including the record of administered procedures to the attending physician for disposition.

Authorized Implementers:

All ED nurses and designated staff who have successfully completed the relevant ED Medical Directive orientation program. It is recommended that hospitals maintain a list of authorized implementers as part of the Medical Directive record.

See Module 2 – Performance Readiness Plan for sample certification forms to maintain a list of authorized implementers. Such forms may be appended to the directive, identified in the ‘Appendix Attached’ section directly above and maintained in accordance with hospital record-keeping policies.
### Indications:

- Therapeutic procedures will be administered for the period beginning from when a patient arrives in the ED to first contact with the attending physician, unless the attending physician orders implementation of the directive beyond that period.
- Prior to implementation of any directive, a patient assessment is completed in accordance with standards of practice and any applicable hospital policy. Allergies and sensitivities must be documented.
- Specific indications are identified in the appended Order Table.

#### Definitions for indications used in the table:

1. **Acute Coronary Syndrome (ACS)** – as manifested by discomfort (pressure or pain, radiating or non-radiating, anterior or posterior) from jaw to umbilicus that may include any of the following:
   - SOB
   - Diaphoresis
   - Pallor
   - Nausea/vomiting
   - Dysrhythmias (palpitations, tachycardia, bradycardia)
   - Syncope
   - Weakness, lightheadedness, pre-syncope
   - Lethargy

2. **Fever** – Temperature >/= 38 C

3. **Hypothermia** – Temperature </= 36 C

4. **Hemodynamic instability** – as manifested by one or more of the following:
   - Pale
   - Diaphoretic
   - Tachypneic
   - Tachycardia,
   - Hypotensive
   - Altered LOC

5. **Immunocompromised** - Patients with one or more of the following:
   - On chemotherapy for cancer,
   - On anti-TNF medication – Infliximab (e.g. Remicade)
   - Organ transplant(s)
   - Splenectomy
   - HIV
   - Lupus, rheumatoid arthritis and other chronic inflammatory conditions
   - Diabetes mellitus
   - Chronic alcohol abuse
   - Chronic corticosteroid therapy

6. **Major bleed** – any volume loss that causes hemodynamic instability resulting from possible GI bleed, ruptured aneurysm, ruptured spleen, femur fracture or ectopic pregnancy

7. **Major trauma** - high risk mechanism of injury

### Contraindications:

See appended Order Table.
### Consent:

Staff implementing the directive will obtain consent in accordance with the *Health Care Consent Act* and any relevant hospital policies and procedures (note these).

### Guidelines for Implementing the Order / Procedure:

See appended Order Table.

### Documentation and Communication:

Implementing staff will document the:

- Specific orders in the order section of the patient record, noting the medical directive name and number, and signing off the order with the implementer’s name and signature as per the attending physician (when attending physician known)
- Indications, implementation and patient response in accordance with hospital record-keeping policies (note these)

Append any designated forms used to document implementation of this directive, identifying them in the ‘Appendix Attached’ section directly above. See Module 4 – Sample Prototype Directive Pre-printed Orders and ED Record for examples of forms that may be used as desired.

**NB:** Clear and timely notification, communication and documentation between the nurse and the physician are critical to safe, proper use of a medical directive.

### Review and Quality Monitoring Guidelines:

Staff identifying any untoward or unintended outcomes arising from implementation of orders under this directive, or any issues identified with it will report these to (note to whom) as soon as possible for appropriate disposition. This does not include untoward or unintended outcomes or issues that are possible clinical sequelae regardless of whether a directive or direct order is used.

Additional provisions (e.g. for renewal and re-certification) may be identified here or in Performance Readiness Assessment form (Module 1).

### Administrative Approvals:

Identify the relevant approval committees and staff members, for example: Unit or Program Administrative Staff and Committees, Educators, Professional Leaders, Professional Councils, Medical Advisory Committee etc., and ensure a signed copy of approvals is maintained as part of the Medical Directive record.

Signatures may be recorded either in this section, or in an attached appendix, or on designated hospital-specific sign-off forms, or see Module 6 – Sample Approval Forms for sign-off forms that may be used as desired.
Identify all the relevant approving attending physicians and ensure a signed copy of approvals is maintained with the Medical Directive record.

In addition, ensure there is an accessible and timely means of informing staff of new attendings (e.g. newly privileged physicians, locums, residents).

Signatures may be recorded either in this section, or in an attached appendix, or on designated hospital-specific sign-off forms, or see Module 6 – Sample Approval Forms for sign-off forms that may be used as desired.

List any references here as below and attach appendix, or list references in Performance Readiness Assessment form (Module 1).


AHA Guidelines, 2007 http://stroke.ahajournals.org/cgi/content/abstract/30/10/2033
### Title and Number of Directive:
**ED Therapeutic Procedures Directive (record directive number)**

### Appendix
**Therapeutic Procedures Order Table**

<table>
<thead>
<tr>
<th>Order</th>
<th>Indications</th>
<th>Contraindications</th>
<th>Guidelines</th>
</tr>
</thead>
</table>
| a) Saline Lock Insertion, or | Signs and symptoms of one or more of the following actual or potential:  
  - Airway compromise  
  - Respiratory distress  
  - Hemodynamic instability / shock, dehydration bleeding  
  - Altered LOC  
  - Severe pain  
  - High risk mechanism of injury  
  - Infection in immunocompromised patients where venous access is not already established  
  - Ingestion of substances or toxins that have or may result in any of the above signs and symptoms  
In anticipation of one or more of the following:  
  - IV medication administration for pain, symptom control and/or/treatment,  
  - Blood product administration,  
  - To provide IV rehydration where dehydration present and oral intake compromised,  
  - To provide fluid resuscitation to improve hemodynamic status |  
  - Hemodialysis lines are not to be accessed for the purposes of this medical directive | Follow applicable hospital policy and procedure |
| b) Peripheral IV Access with NS 30 mLs/hour, or | | | |
| c) Accessing Established Vascular Access Device with NS 30 mLs/hour | | | |

This template has been adapted from the Emergency Department Medical Directives Implementation Kit [www.oha.com/edmedicaldirectives](http://www.oha.com/edmedicaldirectives)
## 1. Therapeutic Procedures Order Table

This table cannot be relied upon in the absence of ED Therapeutic Procedures Directive (record directive number).

| 1. a) Initiate and Titrate Oxygen by mask or nasal prongs to maintain $\text{SaO}_2 \geq 95\%$: | Signs and symptoms of one or more of the following actual or potential:  
- Respiratory distress,  
- $\text{SaO}_2 \leq 94\%$, or below established desirable range for the individual patient  
- Hemodynamic instability  
- Evidence of suspected hypoxemia (chest pain, tachycardia, hemorrhage, hypovolemia, sickle cell, altered LOC, Trauma, smoke and/or toxin inhalation)  
If signs and symptoms and/or $\text{SaO}_2$ levels do not improve promptly, notify physician | **Note:** For possible CVA patients, $\text{O}_2$ administration should only be considered when $\text{SaO}_2 \leq 90\%$ | Follow applicable hospital policy and procedure |
|---|---|---|---|
| 1. b) Urinary Catheter Insertion:  
1. a) Straight in and out  
   Or  
2. ii) Indwelling urinary catheter  
   Or  
3. iii) 3-way catheter | • For urine specimen collection in immobilized patients where unable to collect by other means  
   Or  
   • To assess and monitor fluid balance in patients with actual or potential signs and symptoms of hemodynamic compromise, and/or  
   • Inability to void or difficulty voiding;  
   Or  
   • Moderate to gross hematuria and inability to void or difficulty voiding  
   • Prior to insertion in males, administer Lidocaine Jelly (See ED Medication Medical Directive for applicable order). | • Pelvic trauma  
• Recent urethral surgery or bladder reconstruction  
• Known strictures or previous difficult catheterizations | Consider using coude tip for all male patients  
Follow applicable hospital policy and procedure |
### 1. Therapeutic Procedures Order Table

This table cannot be relied upon in the absence of ED Therapeutic Procedures Directive (record directive number)

<table>
<thead>
<tr>
<th>c) Eye irrigation with 1 L NS or Ringer’s Lactate as per hospital policy with or without insertion of Morgan Lens</th>
<th>d) Spinal Backboard Removal</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Chemical splash to the eye</td>
<td>• Alert and oriented</td>
</tr>
<tr>
<td>• Irrigate eye even if patient has flushed prior to arriving at ED</td>
<td>• Prior to removal, ensure:</td>
</tr>
<tr>
<td>• Prior to irrigation, remove any powdered chemical by dry wiping</td>
<td>o Team presence - all removals from backboards must be done in a safe team fashion as per standards of practice</td>
</tr>
<tr>
<td>• Prior to insertion of Morgan Lens, topical ophthalmic anaesthetic must be administered (See ED Medication Medical Directive for applicable order)</td>
<td>o Physician presence for any patients suspected of severe injuries - these patients should have the number of logrolls and transfers minimized, making physician presence desirable unless physician directs otherwise</td>
</tr>
<tr>
<td></td>
<td>• Focal neurological deficits including parasthesia or loss of sensation in extremities</td>
</tr>
<tr>
<td></td>
<td>• Vomiting</td>
</tr>
<tr>
<td></td>
<td>• Altered LOC</td>
</tr>
<tr>
<td></td>
<td>• Inability to follow direction or verbally communicate symptoms</td>
</tr>
<tr>
<td></td>
<td>• Suspicion of or ingestion of drugs or alcohol rendering patient unable to cooperate or communicate effectively</td>
</tr>
<tr>
<td></td>
<td>• High risk mechanism of injury as defined by the following:</td>
</tr>
<tr>
<td></td>
<td>o Fall from elevation &gt; 3 ft (1 metre/5 stairs)</td>
</tr>
<tr>
<td></td>
<td>o Axial load to head</td>
</tr>
<tr>
<td></td>
<td>o MVC high speed (&gt;100km/hr), roll over or ejection</td>
</tr>
<tr>
<td></td>
<td>o Motorcycle or motorized recreational vehicle collision</td>
</tr>
</tbody>
</table>

Note: No patient should be left indefinitely on a backboard; if contraindications exist or nurses are uncomfortable, suggest summoning the ER physician to assess/assist with removal even if definitive medical assessment is delayed until later. | • Follow applicable hospital policy and procedure |

Follow applicable hospital policy and procedure
Prototype ED Medical Directive - Paediatrics

Title: ED Paediatric Directive  
Number: Assign # in accordance with hospital record-keeping policies

Activation Date:  
Review due by:  

Sponsoring/Contact Person(s)  
(name, position, contact particulars): Typically chief or physician, and expert nursing staff member most responsible for developing directive

Orders:  
Appendix Attached:  
Yes  
No

Title: Paediatric Order Table

Paediatric orders as identified on the appended order table:

1. Paediatric Order Table (p. 35)
   a) Salbutamol and Ipratropium Bromide by Nebulizer
   b) Salbutamol and Ipratropium by Unit Dose Pre-mixed Nebules
   c) Medication by MDI spacer as per hospital policy and practices
   d) Oxygen Administration
   e) Acetaminophen - Pain
   f) Ibuprofen - Pain
   g) Acetaminophen - Fever
   h) Ibuprofen - Fever
   i) Emla®
   j) Oral Rehydration Solution

Recipient Patients:  
Appendix Attached:  
No  
No

Title:

Any child (specify age in accordance with hospital policy, e.g. \(<\leq 15\)) presenting to the ED who meets the conditions identified in this directive.

If a child has therapeutic procedures administered under this directive but leaves before being seen by the attending physician:
• Staff will forward the patient record, including the record of administered procedures to the attending physician for disposition.

Authorized Implementers:  
Appendix Attached:  
No  
No

Title:

All ED nurses and designated staff who have successfully completed the relevant ED Medical Directive orientation program. It is recommended that hospitals maintain a list of authorized implementers as part of the Medical Directive record.

See Module 2 – Performance Readiness Plan for sample certification forms that may be used if desired to maintain a list of authorized implementers. Such forms may be appended to the directive, identifying them in the ‘Appendix Attached’ section directly above or maintained in accordance with hospital record-keeping policies.
Indications:

- Therapeutics will be administered for the period beginning from when a patient arrives in the ED to first contact with the attending physician, unless the attending physician orders implementation of the directive beyond that period. **Exception:** Emla® or Ametop® may be administered throughout a child’s visit.
- Prior to implementation of any directive, a patient assessment is completed in accordance with standards of practice and any applicable hospital policy. Allergies and sensitivities must be documented.
- Specific indications are identified in the appended Order Table.

**Definitions** of Indications used in the order table:

1. **Respiratory distress** is characterized by:
   - Tachypnea
   - Dyspnea reported
   - Nasal flaring
   - Wheezing
   - Indrawing
   - Spasmodic cough
   - Decreased air entry to lung fields on auscultation
   - Cyanosis

2. **Hypoxia** - cyanosis, pallor, decreased level of consciousness, anxiety, or restlessness

3. **Immunocompromised** - Patients with one or more of the following:
   - on chemotherapy for cancer
   - on anti-TNF medication – Infliximab (e.g. Remicade)
   - organ transplant(s)
   - splenectomy
   - HIV
   - lupus, rheumatoid arthritis and other chronic inflammatory conditions
   - diabetes mellitus
   - chronic corticosteroid therapy

Contraindications:

See appended Order Table.

Consent:

Staff implementing the directive will obtain consent in accordance with the **Health Care Consent Act** and any relevant hospital policies and procedures (note these).

Guidelines for Implementing the Order / Procedure:

See appended Order Table.
**Documentation and Communication:**

Implementing staff document the:
- Specific order in the order section of the patient record, noting the medical directive name and number, signing off the order as per the attending physician (when attending physician known)
- Indications, implementation and patient response in accordance with any hospital record-keeping policies (note these)

Append any designated forms used to document implementation of this directive, identifying them in the ‘Appendix Attached’ section directly above. See Module 4 – Sample Prototype Directive Preprinted Orders and ED Record for examples of forms that may be used as desired.

NB: Clear and timely notification, communication and documentation between the nurse and the physician are critical to safe, proper use of a medical directive.

**Review and Quality Monitoring Guidelines:**

Staff identifying any untoward or unintended outcomes arising from implementation of orders under this directive, or any issues identified with it will report these to (note to whom) as soon as possible for appropriate disposition. This does not include untoward or unintended outcomes or issues that are possible clinical sequelae regardless of whether a directive or direct order is used.

Additional provisions (e.g. for renewal and re-certification) may be identified here or in Performance Readiness Assessment form (Module 1).

**Administrative Approvals:**

Identify the relevant approval committees and staff members, for example: Unit or Program Administrative Staff and Committees, Educators, Professional Leaders, Professional Councils, Medical Advisory Committee etc., and ensure a signed copy of approvals is maintained as part of the Medical Directive record. See Appendix for sample form.

Signatures may be recorded either in this section, or in an attached appendix, or on designated hospital-specific sign-off forms, or see Module 6 – Sample Approval Forms for sign-off forms that may be used as desired.

**Approving Physician(s)/Authorizer(s):**

Identify all the relevant approving attending physicians and ensure a signed copy of approvals is maintained with the Medical Directive record. See Appendix for sample form.

In addition, ensure there is an accessible and timely means of informing staff of new attendings (e.g. newly privileged physicians, locums, residents).

Signatures may be recorded either in this section, or in an attached appendix, or on designated hospital-specific sign-off forms, or see Module 6 – Sample Approval Forms for sign-off forms that may be used as desired.

**References**

List any references here as below and attach appendix or list references in Performance Readiness Assessment form (Module 1).

Emergency Medical Directives, Child Health Network for the Greater Toronto Area, December 2002.
## Appendix
### Paediatric Order Table

<table>
<thead>
<tr>
<th>Order</th>
<th>Indications</th>
<th>Contraindications</th>
<th>Guidelines</th>
</tr>
</thead>
<tbody>
<tr>
<td>a) By nebulizer</td>
<td>Child has a history of reactive airway disease (asthma or bronchiolitis) and presents with moderate to severe respiratory distress characterized by one or more of the following:</td>
<td>One or more of the following:</td>
<td>Post-Administration Assessment:</td>
</tr>
<tr>
<td>Salbutamol (5 mg/mL) x 1</td>
<td>• Audible wheezing</td>
<td>• Child presents with sudden onset after a choking episode with a suggestion of foreign body aspiration</td>
<td>• Vital signs, including capillary refill</td>
</tr>
<tr>
<td>• &lt;6.7 kg = 0.2 mL (1 mg)</td>
<td>• Wheezing with retractions</td>
<td>• Child presents with signs and symptoms of upper airway pathology e.g. stridor, drooling, muffled voice, dysphagia</td>
<td>• Oxygen saturation level</td>
</tr>
<tr>
<td>• 6.7 – 33 kg = 0.03 mL/kg (maximum 1 mL {5 mg})</td>
<td>• Spasmodic cough</td>
<td>• There is a known allergy to either of the medications (salbutamol or ipratropium bromide)</td>
<td>• Peak flow</td>
</tr>
<tr>
<td>• 33 kg = 1 mL (5 mg)</td>
<td>• Dyspnea</td>
<td></td>
<td>• Respiratory assessment</td>
</tr>
<tr>
<td>AND</td>
<td>• Tachypnea</td>
<td></td>
<td>• Observe patient’s response to the inhalation and notify physician.</td>
</tr>
<tr>
<td>Ipratropium Bromide (250mcg/mL)</td>
<td>• Decreased air entry into lung fields on auscultation</td>
<td></td>
<td>• Document post-treatment assessment as per organizational guidelines (e.g. including vital signs, including capillary refill, oxygen saturation level, peak flow, respiratory assessment)</td>
</tr>
<tr>
<td>• 250 mcg (1mL) OR</td>
<td>Prior to initiating the medical directive, the nurse will assess the patient by obtaining and documenting the following:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>b) For Unit Dose Pre-mixed Nebules</td>
<td>• Complete set of vital signs, including capillary refill</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Salbutamol</td>
<td>• Respiratory assessment, including chest auscultation, respiratory effort, presence of wheezing, etc.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>(5 mg/mL) x 1</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Up to 10 kg = 0.25 mL</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>• 10-20 kg = 0.5 mL</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>• &gt; 20 kg = 1.0mL</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AND</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ipratropium Bromide</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
### 1. Paediatric Order Table

This table cannot be relied upon in the absence of ED Paediatric Directive (record directive number)

**Based on: Emergency Medical Directives, Child Health Network for the Greater Toronto Area, December 2002.**

<table>
<thead>
<tr>
<th>Order</th>
<th>Indications</th>
<th>Contraindications</th>
<th>Guidelines</th>
</tr>
</thead>
</table>
| (250mcg/mL) x1  
  • 250 mcg  
  OR  
  c) If Using MDI spacer, administer 1 dose as per hospital policy & practices | • Peak flow (in accordance with hospital protocols, if age and ability appropriate [usually over 6 years of age])  
• Oxygen saturation  
• Weight in kilograms  
• History of presenting illness, current medication history and past medical history  
• Assessment should be documented prior to initiating the directive  
• Allergy history | | • If there is no improvement or deterioration in the patient’s condition, notify the physician immediately and initiate oxygen as outlined below in c). |
| d) Oxygen Administration  
Initiate and Titrate Oxygen via blow by nasal prongs or mask to maintain SaO₂ ≥ 95%:  
Signs and symptoms including one or more of the following actual or potential:  
• Respiratory distress  
• SaO₂ < 95%, or below established desirable range for the individual patient  
• Hemodynamic instability / shock  
• Evidence of suspected hypoxemia (tachycardia, hemorrhage, hypovolemia, sickle cell, altered LOC, trauma, or smoke and/or toxin inhalation)  
**Note:** Notify physician immediately of any child who requires O₂ to maintain SaO₂ ≥ 95% | • No contraindications where indications present | • There are few contraindications for oxygen administration in children. Premature infants less than 34 weeks gestation are prone to eye damage. This should never affect the decision to provide short-term emergency oxygen to infants and children of any age. |
1. Paediatric Order Table

This table cannot be relied upon in the absence of ED Paediatric Directive (record directive number)
Based on: Emergency Medical Directives, Child Health Network for the Greater Toronto Area, December 2002.

<table>
<thead>
<tr>
<th>Order</th>
<th>Indications</th>
<th>Contraindications</th>
<th>Guidelines</th>
</tr>
</thead>
<tbody>
<tr>
<td>e) Acetaminophen 15-20 mg/kg PO/PR x 1, maximum 975 mg</td>
<td>Patients triaged as P-CTAS 3,4 or 5 with mild to moderate pain using a developmentally appropriate pain assessment tool with negligible risk of surgical origin and one or more of the following signs and symptoms: • Headache • Dental (e.g. toothache) • Ear, nose and/or throat pain (e.g. possible otitis media) • Musculoskeletal (e.g. minor sprains, possible non-displaced closed fracture) • Skin injuries (e.g. minor burns)</td>
<td>Note: Children who are immunocompromised should never receive any medication by the rectal route unless specifically ordered by a physician. • Child has a known allergy to acetaminophen or ibuprofen • Child has received a therapeutic dose of acetaminophen, ibuprofen or other analgesic within the past 4 hours (acetaminophen) or 6 hours (ibuprofen). If sub-therapeutic dose has been given, calculate the difference and give the remainder of the recommended dose</td>
<td>Post-Administration Assessment: • Reassessment of patient response including vital signs, capillary refill and pain using the same developmentally appropriate pain assessment tool within 30-60 minutes or as indicated</td>
</tr>
</tbody>
</table>

| | | | |
| f) Ibuprofen 5-10 mg/kg PO x 1 to maximum of 40mg/kg/24hrs | If a preceding dose of one of these medications has been taken by the patient, then the other may be administered as indicated under this directive | | |
| | | | |
| | | | |
| | | | |
| | | | |
| | | | |

Form of medication (suspension, tablet, chewable tablet or suppository) should be based on the developmental stage and preference of the child and/or caregiver

Patients triaged as P-CTAS 3,4 or 5 with mild to moderate pain using a developmentally appropriate pain assessment tool with negligible risk of surgical origin and one or more of the following signs and symptoms:

- Headache
- Dental (e.g. toothache)
- Ear, nose and/or throat pain (e.g. possible otitis media)
- Musculoskeletal (e.g. minor sprains, possible non-displaced closed fracture)
- Skin injuries (e.g. minor burns)

Prior to administering the analgesic, the following assessments should be done and documented:

- Vital signs including capillary refill
- Weight in kilograms
- Pain assessment using a developmentally appropriate pain assessment tool
- History of presenting illness, past medical history and recent medication history
- History of allergies

Post-Administration Assessment:

- Reassessment of patient response including vital signs, capillary refill and pain using the same developmentally appropriate pain assessment tool within 30-60 minutes or as indicated
### 1. Paediatric Order Table

This table cannot be relied upon in the absence of ED Paediatric Directive (record directive number)

Based on: Emergency Medical Directives, Child Health Network for the Greater Toronto Area, December 2002.

<table>
<thead>
<tr>
<th>Order</th>
<th>Indications</th>
<th>Contraindications</th>
<th>Guidelines</th>
</tr>
</thead>
</table>
| g) Acetaminophen 15 –20 mg/kg PO/PR x 1, maximum 975 mg | • Child presents at the ED with a temperature of >38.0°C when measured by any route  
• Child must:  
  • Be alert and have an intact gag reflex (if the oral route is to be used)  
  • Be greater than 3 months of age * | Note: Children who are immunocompromised should never receive any medication by the rectal route nor should a temperature be taken by the rectal route unless specifically ordered by a physician.  
• Child has an allergy to acetaminophen or ibuprofen  
• Infant is less than 3 months of age  
• A therapeutic dose of acetaminophen has been given within the past 4 hours or a therapeutic dose of ibuprofen has been given within the past 6 hours. If a sub-therapeutic dose has been given, calculate the difference between the inadequate dose and the therapeutic dose and administer that amount. | Post-administration assessment includes:  
• Vital signs including capillary refill  
• Temperature 30 minutes after medication administration |
| OR  
As per patient choice or hospital formulary:  
| h) Ibuprofen X 1 - Fever | • < 6 months - 5 mg/kg PO  
• 6-12 yrs, temp <39°C – 5 mg/kg PO  
• 6-12 yrs, temp >39°C – 10 mg/kg PO | Note: Infants under 3 months of age, who present with a fever, are classified as Triage Level 2 (P-CTAS) and should be seen by a physician within 15 minutes.  
Prior to initiating the administration of the antipyretic, complete the following assessment:  
• Weight in kilograms  
• Vital signs including capillary refill  
• History of antipyretic therapy (adequacy of dose, response) and other current medication history  
• History of presenting illness and past medical history  
• History of allergy to medication |  
| If a preceding dose of one of these medications has been taken by the patient, then the other may be administered as indicated under this directive |  
| • Form of medication (suspension, tablet, chewable tablet or suppository) should be based on the developmental stage and preference of the child and/or caregiver | |  

*Note: Children who are immunocompromised should never receive any medication by the rectal route nor should a temperature be taken by the rectal route unless specifically ordered by a physician.*
1. Paediatric Order Table

This table cannot be relied upon in the absence of ED Paediatric Directive (record directive number)
Based on: Emergency Medical Directives, Child Health Network for the Greater Toronto Area, December 2002.

<table>
<thead>
<tr>
<th>Order</th>
<th>Indications</th>
<th>Contraindications</th>
<th>Guidelines</th>
</tr>
</thead>
</table>
| i) Emla® (Eutectic Mixture of Lidocaine and Prilocaine) or Ametop® (Amethocaine) | Any child who is a candidate for venous or capillary blood sampling, IM or SC injection, IV initiation, subcutaneous implanted port access (or other venous access device) or lumbar puncture who has intact skin | • Child has a known allergy to the active ingredient in the medication(s) - Emla® (Eutectic Mixture of Lidocaine and Prilocaine) or Ametop® (Amethocaine)  
• The child’s skin is not intact, i.e. broken or lacerated skin  
• There is active eczema or skin rashes | • Cover medicated skin with an occlusive dressing (e.g., Op site®, Tegaderm®)  
• The area will be anaesthetized after 30 minutes (Ametop®) or 60 minutes (Emla®) and effects will last up to 2 hours  
• Prior to blood sampling or IV insertion, wipe medication off with a dry gauze or tissue  
• Perform an aseptic skin preparation prior to needle insertion  
• Ensure all sites are cleansed and dressings removed prior to discharge from the emergency department |

• Oral rehydration solution (ORT) 45-60 mmol/L sodium (Pedialyte™, Lytren™, or Gastrolyte™ or as per hospital formulary):  
  • 50 mg/kg over 4 hours for mild dehydration, or  
  • Child must have vomiting and/or diarrhea and have signs of mild or moderate dehydration as described in the table “Clinical Signs of Dehydration” below.  
  • Prior to initiating the oral rehydration therapy, the following assessment should be done:  
  • Vital signs including blood pressure and | • Do not implement directive and notify physician immediately if:  
  • Child appears extremely ill, lethargic or has altered perfusion  
  • Child has bilious or bloody vomiting | Administration:  
• For mild dehydration, offer a total of 50mLs/kg ORT by age appropriate method (feeding cup, medication cup, syringe, or regular cup) starting with small sips of 5 mL at a time every 1-2 minutes. Bottle-
1. Paediatric Order Table

This table cannot be relied upon in the absence of ED Paediatric Directive (record directive number).

**Based on: Emergency Medical Directives, Child Health Network for the Greater Toronto Area, December 2002.**

<table>
<thead>
<tr>
<th>Order</th>
<th>Indications</th>
<th>Contraindications</th>
<th>Guidelines</th>
</tr>
</thead>
<tbody>
<tr>
<td>• 100 mg/kg over 4 hours for moderate dehydration, and&lt;br&gt;• For each subsequent stool, add ORT 10 mg/kg to 4 hr total, and for any subsequent emesis, add ORT volume for volume to 4 hr total</td>
<td>capillary refill&lt;br&gt;• Weight in kilograms&lt;br&gt;• Level of consciousness&lt;br&gt;• Level of dehydration (see table below)&lt;br&gt;• Urine output (can be estimate e.g. number of wet diapers over past 6-8 hours and judgment of small, medium or large volume of urine by parents estimate)&lt;br&gt;• History of oral intake and the number of stools</td>
<td>• Child has vomiting alone (no diarrhea) and has signs associated with neurologic/toxicologic etiology&lt;br&gt;Clear fluids such as fruit juices, soft drinks, popsicles or sports drinks are not appropriate and should not be used for oral rehydration, unless it is the only fluid the child will accept and parents insist.&lt;br&gt;• For moderate dehydration, offer a total of 100 mL/kg of ORT, plus replacement of ongoing losses (stool and emesis, see below) over a four hour period&lt;br&gt;• For each subsequent stool, add 10 mL/kg of ORT to the four hour total&lt;br&gt;• Emesis should be replaced on a volume-for-volume basis added to the four-hour total&lt;br&gt;• Continue to offer ORT even if vomiting continues</td>
<td>feeding can be used but fluid should be offered in small amounts to avoid further vomiting resulting from drinking a large amount of fluid. The fluid should be offered over a 4 hour period. In addition, ongoing losses (stool and emesis, see below) should be replaced by adding to the 50 mL/kg four hour intake total&lt;br&gt;• If the patient refuses to drink the solution, may substitute Pediapops™ (frozen form of oral rehydration fluid).&lt;br&gt;• May add sugar-free flavouring powder (Crystal Light™ with Aspartame™) to disguise taste. Permission should be obtained from the parent/guardian prior to giving aspartame-containing fluids.&lt;br&gt;Note: This directive may only be used for mild-moderate dehydration. If child has severe dehydration, notify physician immediately.</td>
</tr>
</tbody>
</table>

For a summary of orders, see table “Summary of ORT” below.
1. Paediatric Order Table

This table cannot be relied upon in the absence of ED Paediatric Directive (record directive number).

Based on: Emergency Medical Directives, Child Health Network for the Greater Toronto Area, December 2002.

<table>
<thead>
<tr>
<th>Order</th>
<th>Indications</th>
<th>Contraindications</th>
<th>Guidelines</th>
</tr>
</thead>
</table>

- If infant is breastfed, breastfeeding may be continued in addition to the ORT. The duration of each breastfeeding episode should be kept brief to avoid large amounts of breast milk being ingested at a time, which may induce vomiting.

- Parents may administer ORT. Explain the process clearly to the parents and ask them to record amounts taken.

- Patients unable to increase oral intake within 1 – 2 hours should have an IV started. Notify physician.

Post–directive Patient Assessment:

- Observe the infant/child’s response to ORT at least every 30 minutes, including frequency of vomiting and stooling.
## 1. Paediatric Order Table

This table cannot be relied upon in the absence of ED Paediatric Directive (record directive number)

**Based on: Emergency Medical Directives, Child Health Network for the Greater Toronto Area, December 2002.**

<table>
<thead>
<tr>
<th>Order</th>
<th>Indications</th>
<th>Contraindications</th>
<th>Guidelines</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Check vital signs including capillary refill and colour at least every 30 minutes and more frequently based on assessment</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Monitor level of consciousness/ alertness</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Urine and stool output (number of wet diapers and volume of fluid). It may be necessary to weigh diapers to determine urine/fluid volume output</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Notify physician immediately if deterioration in condition is observed</td>
</tr>
</tbody>
</table>
### ORT Table 1 - Summary of ORT

<table>
<thead>
<tr>
<th>Assessment</th>
<th>Mild Dehydration</th>
<th>Moderate Dehydration</th>
</tr>
</thead>
<tbody>
<tr>
<td>Over first 4 hours</td>
<td>50 mL/kg</td>
<td>100 mL/kg</td>
</tr>
<tr>
<td></td>
<td>Replace each stool loss at 10 mL/kg</td>
<td>Replace each stool loss at 10 mL/kg</td>
</tr>
<tr>
<td></td>
<td>Replace emesis losses at volume for volume</td>
<td>Replace emesis losses at volume for volume</td>
</tr>
<tr>
<td>Reassess intake and patient response every 30 minutes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>If intake poor or condition deteriorates, notify physician immediately</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

### ORT Table 2 - Clinical Signs of Dehydration

<table>
<thead>
<tr>
<th>Weight loss</th>
<th>Mild 3-5%</th>
<th>Moderate 6-10%</th>
<th>Severe 9-15%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vital Signs</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Heart rate</td>
<td>Slight↑</td>
<td>Increased</td>
<td>Markedly increased</td>
</tr>
<tr>
<td>Respiratory rate</td>
<td>Normal</td>
<td>Normal</td>
<td>Tachypnea</td>
</tr>
<tr>
<td>Blood pressure</td>
<td>Normal</td>
<td>Normal</td>
<td>Decreased</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Skin</th>
<th>Mild &lt;2 seconds</th>
<th>Moderate 2-3 seconds</th>
<th>Severe &gt;3 seconds</th>
</tr>
</thead>
<tbody>
<tr>
<td>Capillary refill (abdomen)</td>
<td>&lt;2 seconds</td>
<td>2-3 seconds</td>
<td>&gt;3 seconds</td>
</tr>
<tr>
<td>Elasticity</td>
<td>Normal</td>
<td>Decreased</td>
<td>Increased(tenting)</td>
</tr>
<tr>
<td>Anterior fontanel (&lt;18 months of age)</td>
<td>Normal</td>
<td>Depressed</td>
<td>Depressed</td>
</tr>
<tr>
<td>Mucous membranes</td>
<td>Normal</td>
<td>Dry</td>
<td>Dry</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>CNS</th>
<th>Mild Normal</th>
<th>Moderate Alerted</th>
<th>Severe Depressed</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mental status</td>
<td>Normal</td>
<td>Alerted</td>
<td>Depressed</td>
</tr>
<tr>
<td>Decreased muscle tone</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Eyes</th>
<th>Mild Normal/absent</th>
<th>Moderate Absent</th>
<th>Severe Absent</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tearing</td>
<td>Normal</td>
<td>Absent</td>
<td>Absent</td>
</tr>
<tr>
<td>Appearance</td>
<td>Normal</td>
<td>Sunken</td>
<td>Sunken</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Urine</th>
<th>Mild Small</th>
<th>Moderate Oliguria</th>
<th>Severe Oliguria/anuria</th>
</tr>
</thead>
<tbody>
<tr>
<td>Volume</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
MODULE 4

Sample Directive Pre-Printed Order Forms & ED Record

ED Prototype Medical Directives Module 4 of 6

Use these forms as samples of how to document implementation of directives
## Table of Contents

Sample Pre-Printed Orders:

- Laboratory Testing and Diagnostic Procedures ........................................ 1
- Diagnostic Imaging .................................................................................. 2
- Medications ........................................................................................... 3
- Therapeutic Procedures ......................................................................... 4
- Paediatrics ............................................................................................ 5

Sample ED Record .................................................................................... 6
### (Insert Hospital Name)

**Medical Directive Pre-Printed Orders**

**ED Medical Directive #**

**Labwork and Diagnostic Procedures**

For Use in Area(s):

---

**Sample**

**Patient Identification Here**

---

**DO NOT** use form if expiry date (yyyy/mm) has passed. May only be filled in by staff authorized to implement directive.

<table>
<thead>
<tr>
<th>Activate</th>
<th>Orders</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yes</td>
<td>Initial “yes” or “no” column. If “no”, stroke out orders.</td>
</tr>
<tr>
<td>No</td>
<td></td>
</tr>
</tbody>
</table>

- **Routine Blood Work** – CBC, electrolytes (K, NA, Cl, total CO₂), urea, creatinine & glucose
- **Liver Profile (LFTs)** – Amylase, AST, ALT, T-Bili, Direct Bili, ALP, GGT, Albumin
- **Group & Screen**
- **Serum Quantitative HCG**
- **Serum Drug & Alcohol Screen:**
  - Acetaminophen,
  - ASA
  - ETOH
  - Benzodiazepines
  - Lithium
  - Digoxin
  - Tegretol
  - Dilantin
- **Cardiac Markers:** CK, Troponin (Identify hospital-specific markers)
- **Serum Coagulopathy:** INR/PTT
- **Blood Cultures**
- **Serum Lactate**
- **Point of Care Test (POCT) Urinalysis**
- **Urine for R&M**
- **12 Lead ECG**
- **Capillary Blood Glucose**

Filled in by: ____________________________ / ____________________________

Signature, Designation & Printed Name       Initials  / (yyyy/mm/dd) (hh:mm)

---

*Form #* ("Approval Date")

Original: Health Record; Copy: Pharmacy

*Place in Orders Section of Health Record*

---

This template has been adapted from the Emergency Department Medical Directives Implementation Kit

www.oha.com/edmedicaldirectives
(Insert Hospital Name)
Medical Directive Pre-Printed Orders

ED Medical Directive #
Diagnostic Imaging
For Use in Area(s):

Sample
Patient
Identification
Here

DO NOT use form if expiry date (yyyy/mm) has passed.
May only be filled in by staff authorized to implement directive.

<table>
<thead>
<tr>
<th>Activate</th>
<th>Orders</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yes</td>
<td>Initial “yes” or “no” column. If “no”, stroke out orders.</td>
</tr>
<tr>
<td>No</td>
<td></td>
</tr>
</tbody>
</table>

- Foot x-ray (Specify exam name/views that correspond to hospital-specific policies and procedures)
- Ankle x-ray (‘)
- Knee x-ray (“)
- Tibia/Fibula x-ray (“)
- Hand x-ray (“)
- Finger(s) x-ray (“)
- Scaphoid view x-ray (“)
- Wrist x-ray (‘)
- Forearm x-ray (“)
- Elbow x-ray (“)

Filled in by: ____________________________/______________________________
Signature, Designation & Printed Name                      Initials    ____________________________/________________________
(yyyy/mm/dd) (hh:mm)

“Form #” (‘Approval Date”)
Original: Health Record; Copy: Pharmacy
Place in Orders Section of Health Record
**Sample**

**Patient Identification Here**

---

**(Insert Hospital Name)**

**Medical Directive Pre-Printed Orders**

**ED Medical Directive #**

**Medications**

**For Use in Area(s):**

---

**DO NOT** use form if expiry date (yyyy/mm) has passed.

May only be filled in by staff authorized to implement directive.

<table>
<thead>
<tr>
<th>Activate Orders</th>
<th>Initial “yes” or “no” column. If “no”, stroke out orders.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>☐</td>
<td>☐</td>
</tr>
</tbody>
</table>

Filled in by: __________________________/_____________________________  ________  ___________/________

Signature, Designation & Printed Name  Initials  (yyyy/mm/dd) (hh:mm)

---

"Form #" ("Approval Date")

Original: Health Record; Copy: Pharmacy

*Place in Orders Section of Health Record*
**Sample Medical Directive Pre-Printed Orders**

**ED Medical Directive #**
**Therapeutic Procedures**
**For Use in Area(s):**

**Patient Identification Here**

**DO NOT use form if expiry date (yyyy/mm) has passed. May only be filled in by staff authorized to implement directive.**

<table>
<thead>
<tr>
<th>Activate Orders</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Yes</strong></td>
<td><strong>No</strong></td>
</tr>
</tbody>
</table>

- Initial “yes” or “no” column. If “no”, stroke out orders.

- **Saline Lock Insertion**
- **Peripheral IV Access with NS 30 mLs/hour**
- **Accessing Established Vascular Access Device with NS 30 mLs/hour**

- **Initiate and Titrte Oxygen by mask or nasal prongs to maintain SaO₂ ≥ 95%:**

- **Straight In and Out Urinary Catheter Insertion**
- **Indwelling Urinary Catheter Insertion**
- **3 Way Urinary Catheter Insertion**

- **Eye irrigation with 1 L NS or Ringer’s Lactate as per hospital policy –**
  - with insertion of Morgan Lens
  - without insertion of Morgan Lens

**Filled in by:**

---

Signature, Designation & Printed Name

Initials

(yy/mm/dd) (hh:mm)

**"Form #" ("Approval Date")**

Original: Health Record; Copy: Pharmacy

*Place in Orders Section of Health Record*
### Medical Directive Pre-Printed Orders

**ED Medical Directive #**

Paediatrics

**For Use in Area(s):**

**Sample Patient Identification Here**

---

**DO NOT use form if expiry date (yyyy/mm) has passed.**

**May only be filled in by staff authorized to implement directive.**

<table>
<thead>
<tr>
<th>Activate</th>
<th>Orders</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Yes</strong></td>
<td><strong>Initial “yes” or “no” column. If “no”, stroke out orders.</strong></td>
</tr>
<tr>
<td><strong>No</strong></td>
<td></td>
</tr>
</tbody>
</table>

- **By nebulizer;** Salbutamol (5 mg/ml) x1
  - <6.7 kg = 0.2 ml (1 mg) _____________ mgs
  - 6.7 – 33 kg = 0.03 mL/kg (maximum 1mL {5 mg}) _____________ mgs
  - 33 kg = 1 ml (5 mg) _____________ mgs
  
  **AND**
  
  Ipratropium Bromide (250mcg/ml) **250 mcg (1mL)**

  **OR**

- **For Unit Dose Pre-mixed Nebules:** Salbutamol (5 mg/ml) x1
  - Up to 10 kg = 0.25 ml
  - 10-20 kg = 0.5 ml
  - > 20 kg = 1.0ml

  **AND**

  Ipratropium Bromide (250mcg/ml), 250 mcg x1

  **OR**

  If Using MDI spacer, administer as/hospital policy & practices

- **Initiate and Titrate Oxygen via blow by nasal prongs or mask to maintain SaO2 ≥ 95%:**
  - Acetaminophen 15-20 mg/kg PO/PR x1, max dose 975 mg) - Pain
    - **And/Or**
      - **As/patient choice or hospital formulary;** Ibuprofen 5-10 mg/kg PO x1 maximum of 40mg/kg/24hrs – Pain
  
  - Acetaminophen 15-20 mg/kg PO/PR x1, max dose 975 mg - Fever
    - **And/Or**
      - **As/patient choice or hospital formulary;** Ibuprofen - Fever
        - < 6 months - 5 mg/kg PO _________________ mgs.
        - 6-12 mos, temp <39°C – 5 mg/kg PO _________________ mgs.
        - 6-12 mos, temp ≥39°C – 10 mg/kg PO _________________ mgs.

- **Emla® or Ametop® 1 – 2 gm (~a 25-cent piece) topically to intact skin pre-procedure for pain prevention**

- **Oral rehydration solution 45-60 mmol/L sodium (Pedalyte™, Lytren™, or Gastrolyte™ or as/hospital formulary):**
  - 50 mL/kg over 4 hours for mild dehydration, or
  - 100 mL/kg over 4 hours for moderate dehydration, and
  - For each subsequent stool, add ORT 10 mL/kg to 4 hr total, and for any subsequent emesis, add ORT volume for volume to 4 hr total

- **See MAR (or identify applicable record) for amount administered**

---

**Filled in by:**

__________________________________/______________________________  ________      ___________/________

**Signature, Designation & Printed Name**  **Initials**  (yyyy/mm/dd) (hh:mm)

---

*This template has been adapted from the Emergency Department Medical Directives Implementation Kit*

[www.oha.com/edmedicaldirectives](http://www.oha.com/edmedicaldirectives)
### Emergency Triage Record

**From: Mount Sinai Hospital**

<table>
<thead>
<tr>
<th>Date:</th>
<th>Triage Time:</th>
<th>Room #:</th>
</tr>
</thead>
</table>

**CEDIS Complaint**

#### Canadian Triage and Acuity Scale (CTAS)
- 1 – Resuscitative
- 2 – Emergent
- 3 – Urgent
- 4 – Less urgent
- 5 – Non urgent

**Subjective Assessment**

<table>
<thead>
<tr>
<th>Respiratory Assessment</th>
<th>Haemodynamic Assessment</th>
<th>Glasgow coma Scale (GCS) Score</th>
<th>Pain Score</th>
</tr>
</thead>
<tbody>
<tr>
<td>□ No Distress</td>
<td>□ Skin well perfused</td>
<td>Eye opening</td>
<td>/10</td>
</tr>
<tr>
<td>□ Mild</td>
<td>□ Skin not well perfused</td>
<td>□ 1. No eye opening</td>
<td></td>
</tr>
<tr>
<td>□ Moderate</td>
<td></td>
<td>□ 2. Open in response to pain to limbs</td>
<td></td>
</tr>
<tr>
<td>□ Severe</td>
<td></td>
<td>□ 3. Eyes open in response to speech</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>□ 4. Spontaneous eye opening</td>
<td></td>
</tr>
</tbody>
</table>

**Radial Pulse Quality**

- □ Strong
- □ Weak
- □ Regular
- □ Irregular

**Best Verbal Response**

- □ T. Intubated
- □ 1. No response to pain
- □ 2. Extensor posturing to pain
- □ 3. Inappropriate Speech
- □ 4. Confused Conversation
- □ 5. Orientated Response

**Best Motor Response**

- □ 1. No response to pain
- □ 2. Extensor posturing to pain
- □ 3. Abnormal flexor response to pain
- □ 4. Withdraws to pain
- □ 5. Localizing response to pain
- □ 6. Obey commands

<table>
<thead>
<tr>
<th>Time (HH.MM)</th>
<th>RR</th>
<th>SpO2</th>
<th>HR</th>
<th>BP</th>
<th>GCS</th>
<th>Pain</th>
<th>Temp</th>
<th>Triage Reassessment and Interventions</th>
<th>RN</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Time (HH.MM)</th>
<th>Orders/Dir (Dir=Medical Directive)</th>
<th>Time (HH.MM)</th>
<th>RN</th>
<th>Time (HH.MM)</th>
<th>Order/Dir</th>
<th>Time (HH.MM)</th>
<th>RN</th>
<th>Urine Drip</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>CEC, Lactate, U. Cr, Glucose</td>
<td>Leukocytes</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>GSP</td>
<td>Nitrates</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>INR</td>
<td>Protein</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>TTP, GPT, Trop T</td>
<td>Blood</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>UFi*</td>
<td>Ketones</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Serum-ESTH</td>
<td>Glucose</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Serum-AKA &amp; Asat</td>
<td>Urine Micro</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Blood</td>
<td>Creatinine</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Cultures</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>ECG</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Repeat ECG and Cardiac Markers</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Tetanus Status**

- □ Not known
- □ < 10 years
- □ > 10 years

<table>
<thead>
<tr>
<th>Time (HH.MM)</th>
<th>Orders/Dir (Dir=Medical Directive)</th>
<th>Time (HH.MM)</th>
<th>RN</th>
<th>Time (HH.MM)</th>
<th>Order/Dir</th>
<th>Time (HH.MM)</th>
<th>RN</th>
<th>Urine Drip</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>CEC, Lactate, U. Cr, Glucose</td>
<td>Leukocytes</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>GSP</td>
<td>Nitrates</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>INR</td>
<td>Protein</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>TTP, GPT, Trop T</td>
<td>Blood</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>UFi*</td>
<td>Ketones</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Serum-ESTH</td>
<td>Glucose</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Serum-AKA &amp; Asat</td>
<td>Urine Micro</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Blood</td>
<td>Creatinine</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Cultures</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>ECG</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Repeat ECG and Cardiac Markers</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Emergency Record**

- **History and Chief Complaint**
- **Emergency Record**
- **History and Chief Complaint**
- **Emergency Record**
- **History and Chief Complaint**

**X-Ray:**

- **ECG:**
- **Fecal Occult Blood:**

**Staff Physician Notes**

**Discharge Time**

**Discharge Disposition**

**Prescriptions**

**Discharge Instructions**

- □ Instructed to return if symptoms worsen
- □ Other instructions:

**Diagnosis**

**Follow-up**

- □ Family MD
- □ Other:

<table>
<thead>
<tr>
<th>Time (HH.MM)</th>
<th>Orders/Dir (Dir=Medical Directive)</th>
<th>Time (HH.MM)</th>
<th>RN</th>
<th>Time (HH.MM)</th>
<th>Order/Dir</th>
<th>Time (HH.MM)</th>
<th>RN</th>
<th>Urine Drip</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>CEC, Lactate, U. Cr, Glucose</td>
<td>Leukocytes</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>GSP</td>
<td>Nitrates</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>INR</td>
<td>Protein</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>TTP, GPT, Trop T</td>
<td>Blood</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>UFi*</td>
<td>Ketones</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Serum-ESTH</td>
<td>Glucose</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Serum-AKA &amp; Asat</td>
<td>Urine Micro</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Blood</td>
<td>Creatinine</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Cultures</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>ECG</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Repeat ECG and Cardiac Markers</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Consultation 1**

<table>
<thead>
<tr>
<th>Time (HH.MM)</th>
<th>Orders/Dir (Dir=Medical Directive)</th>
<th>Time (HH.MM)</th>
<th>RN</th>
<th>Time (HH.MM)</th>
<th>Order/Dir</th>
<th>Time (HH.MM)</th>
<th>RN</th>
<th>Urine Drip</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>CEC, Lactate, U. Cr, Glucose</td>
<td>Leukocytes</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>GSP</td>
<td>Nitrates</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>INR</td>
<td>Protein</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>TTP, GPT, Trop T</td>
<td>Blood</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>UFi*</td>
<td>Ketones</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Serum-ESTH</td>
<td>Glucose</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Serum-AKA &amp; Asat</td>
<td>Urine Micro</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Blood</td>
<td>Creatinine</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Cultures</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>ECG</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Repeat ECG and Cardiac Markers</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Consultation 2**

<table>
<thead>
<tr>
<th>Time (HH.MM)</th>
<th>Orders/Dir (Dir=Medical Directive)</th>
<th>Time (HH.MM)</th>
<th>RN</th>
<th>Time (HH.MM)</th>
<th>Order/Dir</th>
<th>Time (HH.MM)</th>
<th>RN</th>
<th>Urine Drip</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>CEC, Lactate, U. Cr, Glucose</td>
<td>Leukocytes</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>GSP</td>
<td>Nitrates</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>INR</td>
<td>Protein</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>TTP, GPT, Trop T</td>
<td>Blood</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>UFi*</td>
<td>Ketones</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Serum-ESTH</td>
<td>Glucose</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Serum-AKA &amp; Asat</td>
<td>Urine Micro</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Blood</td>
<td>Creatinine</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Cultures</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>ECG</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Repeat ECG and Cardiac Markers</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Date: ___________________ Time: _____________ YYYY MM DD HH.MM</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Room: _____________________________________</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Presenting Health Problem**: 

**Health History**: 

**Smoker**: □ Yes □ No

- **Social Situation** Language Spoken ____________ 
- □ Interpretation Services contacted 
- □ Lives with family/friend □ Lives alone □ Has CCAC □ Other
- Clothing/Belongings/Own Medications given to:
  - □ with patient □ Family □ Security □ Other:________
- **Airway** □ Patent □ Compromised
  - □ Oral Airway #________ □ Nasal Airway #________
  - □ Suctioned □ Intubation ET#________ Level at lip =____ cm
- **Breathing** □ No respiratory distress □ Mild distress
  - □ Moderate distress □ Severe distress
- **Breath Sounds** □ Equal, clear bilaterally □ Crackles □ Wheezes □ Other
- **Circulation** Skin colour_____ Skin temp_____ Diaphoresis______
- **Radial Pulse** □ Strong □ Regular □ Weak □ Irregular
  - □ Well hydrated □ Mild dehydration
  - □ Moderate dehydration □ Severe dehydration
- **Neurologic** □ Awake, alert and oriented □ Altered LOC [see Neuro Vital Sign Record, page 3]

**Affix Patient Label**

**Initial Assessment by: ________________________, RN**

<table>
<thead>
<tr>
<th>Interventions</th>
<th>Time</th>
<th>Intravenous Initiation</th>
<th>Gauge</th>
<th>Intravenous Location</th>
<th>Initials</th>
</tr>
</thead>
<tbody>
<tr>
<td>Site #1</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Site #1</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Site #1</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Intravenous Intake**

<table>
<thead>
<tr>
<th>Time</th>
<th>Site #</th>
<th>Solution</th>
<th>Volume</th>
<th>Rate</th>
<th>Volume Infused</th>
<th>To Be Absorbed</th>
<th>End Time (HH:MM)</th>
<th>Initials</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>Urinary Output</th>
<th>Time (HH:MM)</th>
<th>Urine</th>
<th>Initial</th>
<th>Time (HH:MM)</th>
<th>Urine</th>
<th>Initial</th>
<th>Time (HH:MM)</th>
<th>Urine</th>
<th>Initial</th>
</tr>
</thead>
</table>

|------------------------------|-------------|-------|---------|-------------|-------|---------|-------------|-------|---------|

**Medications**

<table>
<thead>
<tr>
<th>Time (HH:MM)</th>
<th>Medications</th>
<th>Dose</th>
<th>Route</th>
<th>Initials</th>
<th>Time (HH:MM)</th>
<th>Effect</th>
<th>Initials</th>
</tr>
</thead>
</table>

- **Airway** □ N/A □ Distention
  - □ Guarding □ Rigidly
- **Genitourinary** □ N/A □ Hematuria
  - □ Frequency_________ □ Flank pain □ Right □ Left
- **Gynecologic** □ N/A
  - □ Last menstrual period ________ □ Gravida/Para ________
  - □ Estimated Date of confinement __________
  - □ Vaginal bleeding Amount ________ Duration ________
- **Musculoskeletal** □ N/A
  - □ Limb ________ Tender @ ________
  - □ Deformity @ ________ Weight Bearing ________
  - □ Distal Pulse ________ Sensation ________
  - □ Parasthesia ________
- **Pain** □ N/A
  - □ Location ________ Radiation ________
  - □ Quality ________ Time of onset ________
  - □ Provoked/Alleviated by ________ Pain Score ________/10
- **Symptoms** □ N/A
  - □ Nausea ________ Vomiting ________ Diarrhea ________
  - □ Last bowel movement ________

<table>
<thead>
<tr>
<th>Initials</th>
<th>Initials</th>
<th>Initials</th>
<th>Initials</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>Site #1</th>
<th>Site #1</th>
<th>Initials</th>
<th>Initials</th>
</tr>
</thead>
<tbody>
<tr>
<td>Site #1</td>
<td>Site #1</td>
<td>Initials</td>
<td>Initials</td>
</tr>
</tbody>
</table>
### Vital Signs

<table>
<thead>
<tr>
<th>Time (HH:MM)</th>
<th>Heart Rate</th>
<th>Cardiac Rhythm</th>
<th>Blood Pressure</th>
<th>Respiratory Rate</th>
<th>SpO2</th>
<th>Oxygen LPM %</th>
<th>Temperature</th>
<th>Eyes</th>
<th>Verbal</th>
<th>Motor</th>
<th>Pupils</th>
<th>Strength</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

### Eyes

- Pupils: Size (mm), React (+/-)
- Size (mm), React (+/-)
- Size (mm), React (+/-)

### Glasgow Coma Scale

<table>
<thead>
<tr>
<th>Size</th>
<th>mm</th>
<th>React</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

### Temporal Arterial Lines (R, L)

- Right Arm
- Left Arm
- Right Leg
- Left Leg

### Transfer Summary

- Date (YYYY-MM-DD)
- Time (HH:MM)
- Unit
- Report given by
- Report given to

### Discharge Summary

- Date (YYYY-MM-DD)
- Time (HH:MM)
- Accompanied by
- Method of transportation

### Follow-up Plan

- Written Instructions
- Health Teaching

### Discharge Nurse

- Name
- Initials
- Name
- Initials
- Name
- Initials
- Name
- Initials

**This template has been adapted from the Emergency Department Medical Directives Implementation Kit.**

www.oha.com/edmedicaldirectives
MODULE 5

CEDIS Presenting Complaints Table
ED Prototype Medical Directives Module 5 of 6

Use this table as a quick reference for summarizing directive orders by presenting complaints

THIS IS NOT A MEDICAL DIRECTIVE
This table presents a partial CEDIS Presenting Complaint List (Version 1.0, 2007) categorized by ED Prototype Medical Directives. It has been developed:

1. As a practice prompt to guide use of the ED prototype directives, and
2. To facilitate evaluation of the directives’ effect on patient flow.

It is not, and may not be used as a medical directive because it lacks mandatory information necessary to qualify as one. Therefore, it cannot be used to authorize performance of procedures identified in it. Refer to Module 3 – Prototype Directives for prototype medical directives.

### Draft CEDIS Presenting Complaints Table

April 10, 2008

<table>
<thead>
<tr>
<th>CEDIS Presenting Complaint (Version 1.0)</th>
<th>Corresponding Prototype Medical Directives (See Module 3)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Cardiovascular</strong></td>
<td><strong>Adults</strong></td>
</tr>
<tr>
<td>• Chest Pain</td>
<td>• RBW</td>
</tr>
<tr>
<td></td>
<td>• INR/Ptt</td>
</tr>
<tr>
<td></td>
<td>• Cardiac Markers</td>
</tr>
<tr>
<td></td>
<td>• 12 Lead ECG</td>
</tr>
<tr>
<td>• Palpitations/ Irregular heartbeat</td>
<td>• RBW</td>
</tr>
<tr>
<td></td>
<td>• INR/Ptt</td>
</tr>
<tr>
<td></td>
<td>• Cardiac Markers</td>
</tr>
<tr>
<td></td>
<td>• 12 Lead ECG</td>
</tr>
<tr>
<td>• General Weakness</td>
<td>• RBW</td>
</tr>
<tr>
<td></td>
<td>• Cardiac Markers</td>
</tr>
<tr>
<td></td>
<td>• Urinalysis (POCT)</td>
</tr>
<tr>
<td></td>
<td>• If POCT +ve, then Urine – R&amp;M</td>
</tr>
<tr>
<td>CEDIS Presenting Complaint (Version 1.0)</td>
<td>Corresponding Prototype Medical Directives (See Module 3)</td>
</tr>
<tr>
<td>----------------------------------------</td>
<td>-----------------------------------------------------</td>
</tr>
<tr>
<td></td>
<td>Adults</td>
</tr>
<tr>
<td></td>
<td>Laboratory Tests and Diagnostic Procedures</td>
</tr>
<tr>
<td>Syncope/Pre-syncope</td>
<td>12 Lead ECG</td>
</tr>
<tr>
<td>Environmental</td>
<td>RBW</td>
</tr>
<tr>
<td>Gastrointestinal</td>
<td>RBW, Liver Profile (LFTs) (for right upper or epigastric pain)</td>
</tr>
<tr>
<td>Abdominal Pain</td>
<td>RBW, Group and Screen, Serum Quantitative HCG as/hospital policy (female patients who may be pregnant)</td>
</tr>
<tr>
<td>Diarrhea</td>
<td>RBW</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
</tbody>
</table>

This table is not and may not be used as a medical directive. See Module 3 - Prototype Directives for prototype medical directives.
<table>
<thead>
<tr>
<th>CEDIS Presenting Complaint (Version 1.0)</th>
<th>Corresponding Prototype Medical Directives (See Module 3)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Adults</td>
</tr>
<tr>
<td></td>
<td>Laboratory Tests and Diagnostic Procedures</td>
</tr>
<tr>
<td>• Nausea and/or vomiting</td>
<td>• RBW</td>
</tr>
<tr>
<td>• Parietal pain</td>
<td>• RBW</td>
</tr>
<tr>
<td>• Blood in stool/melena</td>
<td>• RBW</td>
</tr>
<tr>
<td>• Vomiting</td>
<td>• RBW</td>
</tr>
<tr>
<td>• Blood in stool/melena</td>
<td>• RBW</td>
</tr>
<tr>
<td>• Genitourinary</td>
<td></td>
</tr>
<tr>
<td>• Flank pain</td>
<td></td>
</tr>
<tr>
<td>• Hematuria</td>
<td></td>
</tr>
<tr>
<td>• Urinary retention</td>
<td></td>
</tr>
<tr>
<td>• UTI complaints</td>
<td></td>
</tr>
<tr>
<td>CEDIS Presenting Complaint (Version 1.0)</td>
<td>Corresponding Prototype Medical Directives (See Module 3)</td>
</tr>
<tr>
<td>----------------------------------------</td>
<td>----------------------------------------------------------</td>
</tr>
<tr>
<td></td>
<td>Laboratory Tests and Diagnostic Procedures</td>
</tr>
<tr>
<td>Neurologic</td>
<td>RBW</td>
</tr>
<tr>
<td></td>
<td>Seizure</td>
</tr>
<tr>
<td></td>
<td>Extremity weakness/symptoms of CVA</td>
</tr>
<tr>
<td>Ob-Gyn</td>
<td>Vaginal bleed</td>
</tr>
<tr>
<td></td>
<td>Ophthalmology</td>
</tr>
</tbody>
</table>
### CEDIS Presenting Complaints Table (Version 1.0)

<table>
<thead>
<tr>
<th>CEDIS Presenting Complaint</th>
<th>Corresponding Prototype Medical Directives (See Module 3)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adults</td>
<td>Paeds</td>
</tr>
<tr>
<td>Laboratory Tests and Diagnostic Procedures</td>
<td>Diagnostic Imaging</td>
</tr>
<tr>
<td>• Foreign body, eye</td>
<td>• Td</td>
</tr>
<tr>
<td>• Eye pain</td>
<td>• Analgesic eye drops</td>
</tr>
<tr>
<td>Orthopedic</td>
<td>• Urinalysis (POCT)</td>
</tr>
<tr>
<td>• Traumatic back/spine injury</td>
<td>• If POCT +ve, then Urine – R&amp;M</td>
</tr>
<tr>
<td>• Upper or lower extremity injury</td>
<td>• Serum Quantitative HCG as per hospital policy</td>
</tr>
<tr>
<td>Respiratory</td>
<td>• Urine Qualitative HCG as hospital policy</td>
</tr>
<tr>
<td>• Shortness of breath</td>
<td>• Extremity x-rays</td>
</tr>
<tr>
<td>Skin</td>
<td>• • Acetaminophen or Ibuprofen</td>
</tr>
<tr>
<td>• Abrasion</td>
<td>• Acetaminophen or Ibuprofen</td>
</tr>
</tbody>
</table>
### CEDIS Presenting Complaint (Version 1.0)

<table>
<thead>
<tr>
<th>CEDIS Presenting Complaint</th>
<th>Corresponding Prototype Medical Directives (See Module 3)</th>
<th>Adults</th>
<th>Medications</th>
<th>Therapeutic Procedures</th>
<th>Paeds</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Laboratory Tests and Diagnostic Procedures</strong></td>
<td><strong>Diagnostic Imaging</strong></td>
<td><strong>Medications</strong></td>
<td><strong>Therapeutic Procedures</strong></td>
<td><strong>Paeds</strong></td>
<td></td>
</tr>
<tr>
<td>Laceration/Puncture</td>
<td></td>
<td></td>
<td></td>
<td>Acetaminophen or Ibuprofen</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>LET</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Td</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Burn</td>
<td></td>
<td></td>
<td>Td</td>
<td>Acetaminophen or Ibuprofen</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Acetaminophen or Ibuprofen</td>
<td></td>
</tr>
<tr>
<td><strong>Substance Misuse</strong></td>
<td></td>
<td></td>
<td>12 Lead ECG</td>
<td></td>
<td>Saline Lock/ IV/Access VAD O₂</td>
</tr>
<tr>
<td>Substance misuse/intoxication</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>RBW, Liver Profile (LFTs), Serum alcohol and/or drug screen as per hospital policy</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cap glucose</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Overdose ingestion</td>
<td></td>
<td></td>
<td>12 Lead ECG</td>
<td></td>
<td>Saline Lock/ IV/Access VAD O₂</td>
</tr>
<tr>
<td>RBW</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Liver Profile (LFTs) (For ASA OD, consider AST only as per hospital policy), Serum alcohol and/or drug screen as per hospital policy</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cap glucose</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*This table is not and may not be used as a medical directive. See Module 3 - Prototype Directives for prototype medical directives.*
### CEDIS Presenting Complaint (Version 1.0)

<table>
<thead>
<tr>
<th>Trauma</th>
<th>Adults</th>
<th>Paeds</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Laboratory Tests and Diagnostic Procedures</td>
<td>Diagnostic Imaging</td>
</tr>
<tr>
<td><strong>Major trauma – penetrating or blunt</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>• RBW</td>
<td>• 12 Lead ECG</td>
<td></td>
</tr>
<tr>
<td>• G&amp;S</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• INR/Ptt</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Serum alcohol and/or drug screen as/hospital policy</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Serum Quantitative HCG (female patient who may be pregnant)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Liver Profile (LFTs) for blunt abdominal trauma</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

| | | | |
| **General & Minor** | | | |
| | | | |
| • Fever | | | |
| (Note: Patients with sepsis may present with hypothermia, same orders may apply) | | | |
| • RBW | | | |
| • Blood Cultures | | | |
| • Serum Lactate (patients who meet SIRS criteria) | | | |
| • Urinalysis (POCT) if UTI symptoms or patient ≥ 65 yrs. | | | |

| | | | |
| • Hyperglycemia | | | |
| • Cap Glucose | | | |

---

This template has been adapted from the Emergency Department Medical Directives Implementation Kit

[www.oha.com/edmedicaldirectives](http://www.oha.com/edmedicaldirectives)
<table>
<thead>
<tr>
<th>CEDIS Presenting Complaint (Version 1.0)</th>
<th>Corresponding Prototype Medical Directives (See Module 3)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Adults</td>
</tr>
<tr>
<td></td>
<td>Paeds</td>
</tr>
<tr>
<td><strong>Laboratory Tests and Diagnostic Procedures</strong></td>
<td><strong>Diagnostic Imaging</strong></td>
</tr>
<tr>
<td><strong>Hypoglycemia</strong></td>
<td>Cap Glucose</td>
</tr>
<tr>
<td><strong>Pain Management</strong>¹</td>
<td></td>
</tr>
<tr>
<td>• Earache</td>
<td></td>
</tr>
<tr>
<td>• Dental/Gum problem</td>
<td></td>
</tr>
<tr>
<td>• Facial trauma (minor)</td>
<td></td>
</tr>
<tr>
<td>• Sore throat</td>
<td></td>
</tr>
<tr>
<td>• URTI complaints</td>
<td></td>
</tr>
<tr>
<td>• Headache</td>
<td></td>
</tr>
<tr>
<td>• Upper Extremity Pain</td>
<td></td>
</tr>
<tr>
<td>• Lower Extremity Pain</td>
<td></td>
</tr>
<tr>
<td>• Back Pain</td>
<td></td>
</tr>
<tr>
<td>• Pre-procedure anaesthetic</td>
<td></td>
</tr>
</tbody>
</table>

¹ Pain Management is not on the CEDIS list. In the interests of being complete, it appears in the table because the orders and corresponding indications appear in 2 prototype directives: Medication (Adults) and Paediatrics. See Module 3: Prototype Directives.
MODULE 6

Sample Approval Forms
ED Prototype Medical Directives Module 6 of 6

Use these samples to develop or augment forms for obtaining approvals. Includes forms for locums and for Chief to approve by proxy.
Sample Approval Forms:

1. Physician Approval Form – Staff Physicians
2. Agreement for Chief to Approve Directive on Behalf of ED Physicians (Approval by Proxy)
3. Stakeholder/Administrative Approval Form
4. Physician Approval Form – Locums and Temporary Physicians
5. Physician Approval Form – New Staff

Acknowledgements:

Physician and Stakeholder/Administrative Approval Form templates from: An Interprofessional Guide on Orders, Directives and Delegation; Federation of Health Regulatory Colleges of Ontario (January 2007)
Physician Approval Form – Staff Physicians

Title and Number of Directive: 

Each undersigned physician agrees:

- With the content of the directive and that it is an intervention that can be implemented safely and effectively given the circumstances in the ED as understood by the physician,
- To assume the care of patients who have had an intervention performed as authorized by the directive, and
- S/he knows how the staff will document or communicate when a directive has been implemented so s/he can assume care appropriately.

<table>
<thead>
<tr>
<th>Name of Physician</th>
<th>Signature</th>
<th>Date</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Template from: An Interprofessional Guide on Orders, Directives and Delegation; Federation of Health Regulatory Colleges of Ontario (January 2007)
Sample Agreement for Chief to Approve Directive(s) on Behalf of ED Physicians

For the Period: ____________________________________________________________

Either party may terminate this agreement at any time with notice.

Chief Agreement:
I, ____________________________ (Name of Chief), accept proxy from the
undersigned physician(s) and agree to:
• Approve a medical directive that is agreed to by each physician, and
• Ensure each physician is kept informed of the directive.

_____________________________                     ______________________
Signature of Chief                  Date

ED Physician Agreement:
I, an undersigned physician, give proxy to the Chief named above to approve a medical directive on my behalf
under the conditions identified above and agree to:
• Review the directive to fully understand the conditions under which it will be implemented, including
  knowing how the staff will document or make me aware that the directive has been implemented so I
  can assume care appropriately, and
• Assume the care of patients who have had an intervention performed as authorized by the directive

_____________________________                     ______________________
Name of Physician                  Signature                  Date

_____________________________                     ______________________
Name of Physician                  Signature                  Date

_____________________________                     ______________________
Name of Physician                  Signature                  Date

_____________________________                     ______________________
Name of Physician                  Signature                  Date

_____________________________                     ______________________
Name of Physician                  Signature                  Date

_____________________________                     ______________________
Name of Physician                  Signature                  Date
Stakeholder/Administrative Approval Form

**Title and Number of Directive:**

In accordance with role responsibilities, the undersigned agree with the use of the directive:

<table>
<thead>
<tr>
<th>Name and Position</th>
<th>Signature</th>
<th>Date</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Template from: An Interprofessional Guide on Orders, Directives and Delegation; Federation of Health Regulatory Colleges of Ontario (January 2007)
Physician Approval Form – Locums and Temporary Physicians

Title and Number of Directive/Delegation: __________________________________________

Title and Number of Directive/Delegation: __________________________________________

Title and Number of Directive/Delegation: __________________________________________

Title and Number of Directive/Delegation: __________________________________________

Title and Number of Directive/Delegation: __________________________________________

Title and Number of Directive/Delegation: __________________________________________

Title and Number of Directive/Delegation: __________________________________________

Each undersigned physician has reviewed each directive above and agrees:

- With the directive, its content, and that it is an intervention that can be implemented safely and effectively given that it has been approved by the ED Chief, physicians and administrative authorities, and given the circumstances in the ED as they understand them
- To assume the care of patients who have had an intervention performed as authorized by the directive
- S/he knows how the staff will document or communicate that a directive has been implemented so s/he can assume care appropriately

<table>
<thead>
<tr>
<th>Name of Physician</th>
<th>Signature</th>
<th>Date</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Physician Approval Form – New Staff

Title and Number of Directive/Delegation: ______________________________

Title and Number of Directive/Delegation: ______________________________

Title and Number of Directive/Delegation: ______________________________

Title and Number of Directive/Delegation: ______________________________

Title and Number of Directive/Delegation: ______________________________

Title and Number of Directive/Delegation: ______________________________

Title and Number of Directive/Delegation: ______________________________

Title and Number of Directive/Delegation: ______________________________

Title and Number of Directive/Delegation: ______________________________

Title and Number of Directive/Delegation: ______________________________

Each undersigned physician has reviewed each directive above and agrees:

• With the directive, its content, and that it is an intervention that can be implemented safely and effectively given the circumstances in the ED as they understand them
• To assume the care of patients who have had an intervention performed as authorized by the directive
• S/he knows how the staff will document or communicate that a directive has been implemented so s/he can assume care appropriately

<table>
<thead>
<tr>
<th>Name of Physician</th>
<th>Signature</th>
<th>Date</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Name of Physician</th>
<th>Signature</th>
<th>Date</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>