How to respond to radiological, biological and chemical threats:
A guide for the European Front-Line Health Professional

The ETHREAT Scientific Committee
How to respond to radiological, biological and chemical threats:

A guide for the European Front-Line Health Professional

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ACKNOWLEDGEMENTS
This handbook is dedicated to the memory of
Georgios Gouvras  (1948-2006)

George Gouvras was head of Unit for
Health Threats at the European
Commission’s Directorate for Public
Health and he was the person behind
the EU Health Security programme,
set up in December 2001 to improve
preparedness, response and
cooperation in the event of biological
and/or chemical agent attacks. He also
guided the preparedness for the
response of the European Community
against pandemic influenza.

George helped create the legislation for the ECDC and was member
of its first Management Board. He had an astute understanding of the
needs of the European Community, and a talent for explaining the
need for policy change and paving the Commission’s road to actually
implement it in order to serve better the citizens of Europe.
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The new threats to health, resulting from deliberate releases, bring up new needs for the diffusion of existing knowledge. The training of health professionals stands out as a priority that should be coherently addressed at the European level as part of a long-term strategy to respond to the changing environment of public health threats. During its course (2005-2008), the European Training for Health Professionals on Rapid Response to Health Threats (ETHREAT) Project, developed an educational package for front line health professionals (FLHP) that specifically targets the new public health threats.

Six European Member States (Bulgaria, Czech Republic, Germany, Greece, Poland and the UK) participated at the ETHREAT consortium representing highly specialized organisations and institutes on public health issues. The National and Kapodistrian University of Athens was responsible for the management and coordination of the project as well as for the cooperation of the whole partnership with the European Commission (Directorate for Public Health), which co-financed the project.

The aim of the present handbook is to assist training institutions, universities and public health authorities in the education of health professionals, so as to enhance the European human capital on the timely identification, the management and response to events that could be the result of deliberate attacks with the use of biological, chemical and radiological agents. In other words, the educational package contains the basic necessary knowledge
and training material to empower European health professionals, including armed forces health personnel, to clinically recognise and to respond rapidly and adequately to new public health threats, like attacks with biological, chemical and radiological agents.

The material presented in this handbook originates on one hand from an inventory of existing courses and training material from the EU and other countries addressing health professionals, including armed forces and front line public health professionals on the same issues of new threats to public health and on the other from the Endnote© databank library we created, which includes all collected material on the education and training of health professionals on new public health threat issues and all identified sources of information related to the project with the relevant electronic links.

Moreover, the project team explored the opinions of their target audience and of European experts on the existence and appropriateness of currently available programmes, as well as the desired content of an educational package by surveying front line health professionals (FLHP) and public health (PH) and CBRN experts in the European Union (EU) member states (MS).

The enclosed CD-ROM contains the presentations and educational material presented in a pilot course that took place in Athens in May 2007. The course was organised in such a way so as to include all the necessary information and guidelines that public health organisations, governments, civil authorities, security agencies and armed forces will need to disseminate to front-line health professionals in emergency situations caused by biological, chemical or radiological forms of terrorism as well as the algorithm guide of actions to be undertaken in case of terrorist attacks for most European Member States.

This handbook is a user friendly basic awareness manual accompanied by electronic material aiming at empowering front-line health professionals (FLHP), to rapidly recognise and adequately respond to new public health threats.

We urge you to disseminate the material in your own country in its current form or to translate it in your own language. The ETHREAT team can assist you with the dissemination or organisation of possible courses. We are more than willing to help with such activities and we are also interested in hearing your opinion or comments regarding the contents of this package in the email address below.

We hope that this handbook provides you with sound basic knowledge and outlines the first steps that you need to take in order to handle the initial phase of an unknown and possibly dangerous situation for your staff and yourself.

On behalf of the ETHREAT Project Team

ATHENA LINOS, MD, MPH, PhD
Associate Professor, University of Athens Medical School, ETHREAT Project Coordinator

Email: alinos@med.uoa.gr
### Abbreviations - Glossary

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<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tr>
<td>ABC</td>
<td>Airway, Breathing, Circulation</td>
</tr>
<tr>
<td>ARDS</td>
<td>Acute Respiratory Distress Syndrome</td>
</tr>
<tr>
<td>ARS</td>
<td>Acute Radiation Syndrome</td>
</tr>
<tr>
<td>ASAP</td>
<td>As soon as possible</td>
</tr>
<tr>
<td>BAL</td>
<td>British Anti-Lewisite</td>
</tr>
<tr>
<td>bid or bd</td>
<td>Twice a day</td>
</tr>
<tr>
<td>BSA</td>
<td>Body Surface Area</td>
</tr>
<tr>
<td>CAP</td>
<td>Community-acquired pneumonia</td>
</tr>
<tr>
<td>CBC</td>
<td>Complete Blood Count</td>
</tr>
<tr>
<td>CBRN</td>
<td>Chemical Biological Radiological and Nuclear</td>
</tr>
<tr>
<td>CCHF</td>
<td>Crimean Congo Hemorrhagic Fever</td>
</tr>
<tr>
<td>CD</td>
<td>Communicable Diseases</td>
</tr>
<tr>
<td>CDC</td>
<td>Centers for Disease Control and Prevention, Atlanta</td>
</tr>
<tr>
<td>CNS</td>
<td>Central nervous system</td>
</tr>
<tr>
<td>COPD</td>
<td>Chronic Obstructive Pulmonary Disease</td>
</tr>
<tr>
<td>CSF</td>
<td>Cerebrospinal fluid</td>
</tr>
<tr>
<td>CXR</td>
<td>Chest X-ray</td>
</tr>
<tr>
<td>d</td>
<td>Day(s)</td>
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<tr>
<td>DIC</td>
<td>Disseminated Intravascular Coagulation</td>
</tr>
<tr>
<td>DMPS</td>
<td>2,3-dimercaptopropanesulfonic acid, which is a metal chelator</td>
</tr>
<tr>
<td>DMSA</td>
<td>2,3-dimercaptopropanoic acid, an organic metal chelator</td>
</tr>
<tr>
<td>DTPA</td>
<td>Diethylene-triamine penta-acetate</td>
</tr>
<tr>
<td>EC</td>
<td>European Commission</td>
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<tr>
<td>ECDC</td>
<td>European Centre for Disease Control and Prevention</td>
</tr>
<tr>
<td>ED</td>
<td>Emergency Department</td>
</tr>
<tr>
<td>EEE</td>
<td>Eastern Equine Encephalitis</td>
</tr>
<tr>
<td>EM</td>
<td>Electron microscopy</td>
</tr>
<tr>
<td>ETHREAT</td>
<td>European Training for Health Professionals on Rapid Response to Health Threats project</td>
</tr>
<tr>
<td>EU</td>
<td>European Union</td>
</tr>
<tr>
<td>FLHP</td>
<td>Front Line Health Professionals</td>
</tr>
<tr>
<td>g</td>
<td>Gram</td>
</tr>
<tr>
<td>Abbreviation</td>
<td>Description</td>
</tr>
<tr>
<td>--------------</td>
<td>-------------</td>
</tr>
<tr>
<td>Gram +/-</td>
<td>Gram positive/negative</td>
</tr>
<tr>
<td>Gy</td>
<td>Gray (radiation absorbed per unit mass of tissue)</td>
</tr>
<tr>
<td>HAZMAT</td>
<td>Hazardous Material</td>
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<tr>
<td>HCW</td>
<td>Health care worker</td>
</tr>
<tr>
<td>HF</td>
<td>Hemorrhagic Fever</td>
</tr>
<tr>
<td>hr(s)</td>
<td>hour(s)</td>
</tr>
<tr>
<td>IV</td>
<td>intravenous</td>
</tr>
<tr>
<td>JE</td>
<td>Japanese Encephalitis</td>
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<tr>
<td>m</td>
<td>meters</td>
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<tr>
<td>μ OR μm</td>
<td>micrometer (1 x 10^-6 m)</td>
</tr>
<tr>
<td>mg</td>
<td>milligram</td>
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<tr>
<td>min</td>
<td>minutes</td>
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<tr>
<td>mm</td>
<td>millimeter</td>
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<tr>
<td>MOF</td>
<td>multiple organ failure</td>
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<tr>
<td>OPCW</td>
<td>Organisation for the Prohibition of Chemical Weapons</td>
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<tr>
<td>PO</td>
<td>orally</td>
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<tr>
<td>PPE</td>
<td>Personal Protective Equipment</td>
</tr>
<tr>
<td>PTSD</td>
<td>Post Traumatic Stress Disorder</td>
</tr>
<tr>
<td>RBC</td>
<td>Radiological, Biological and Chemical agents</td>
</tr>
<tr>
<td>RDD</td>
<td>Radiation Dispersion Device</td>
</tr>
<tr>
<td>SARS</td>
<td>Severe Acute Respiratory Syndrome</td>
</tr>
<tr>
<td>SEB</td>
<td>Staphylococcal Enterotoxin B</td>
</tr>
<tr>
<td>sec(s)</td>
<td>second(s)</td>
</tr>
<tr>
<td>Sv</td>
<td>Sievert (equivalent dose), mSv=milliSievert</td>
</tr>
<tr>
<td>TB</td>
<td>tuberculous</td>
</tr>
<tr>
<td>TBE</td>
<td>Tick-borne encephalitis</td>
</tr>
<tr>
<td>TIC</td>
<td>Toxic Industrial Chemical(s)</td>
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<tr>
<td>US</td>
<td>United States</td>
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<tr>
<td>VEE</td>
<td>Venezuelan Equine Encephalitis</td>
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<tr>
<td>VHF</td>
<td>Viral Hemorrhagic Fever</td>
</tr>
<tr>
<td>w</td>
<td>with</td>
</tr>
<tr>
<td>WEE</td>
<td>Western Equine Encephalitis</td>
</tr>
<tr>
<td>WNV</td>
<td>West Nile virus</td>
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<tr>
<td>WW I and II</td>
<td>World War I and II</td>
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</tbody>
</table>
During the last two decades PH professionals have faced a number of serious PH threats and are increasingly worried over the possibility of international spread and the effects of Communicable Diseases (CD). Predictions based on climatic changes, as well as the global socioeconomic situation indicate that humanity as a whole is going to face in the future increasing number of international outbreaks, as humans on one hand invade more and more tropical forests and come in contact with new agents and on the other travel long distances in large numbers.

Recent experiences of outbreaks or other incidents (e.g. SARS outbreak (2003), Avian Influenza outbreaks (2005-2008) and melamine contaminated milk (2008)) have shown that no nation is completely safe or immune to a spreading epidemic or the effects of globalised trade. Collaboration at multiple levels and across multiple sectors is needed in order to enable efficient response to the challenge of these new threats.

### NEW HEALTH THREATS AND PUBLIC HEALTH

<table>
<thead>
<tr>
<th>Disease</th>
<th>Year of Outbreak</th>
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<tbody>
<tr>
<td>Human African trypanosomiasis</td>
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<tr>
<td>Cholera</td>
<td></td>
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<tr>
<td>Marburg haemorrhagic fever</td>
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<tr>
<td>MDR/XDR tuberculosis</td>
<td></td>
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<tr>
<td>Plague</td>
<td></td>
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<tr>
<td>Human monkeypox</td>
<td></td>
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<tr>
<td>Chikungunya fever</td>
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<tr>
<td>Enterovirus 71</td>
<td></td>
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<tr>
<td>Hendra virus</td>
<td></td>
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<tr>
<td>Nipah virus</td>
<td></td>
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<tr>
<td>Vancomycin-resistant Staphylococcus aureus</td>
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<tr>
<td>H5N1 influenza</td>
<td></td>
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<tr>
<td>Typhoid fever</td>
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<tr>
<td>Rift Valley fever</td>
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<tr>
<td>Diphtheria</td>
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<tr>
<td>Drug-resistant malaria</td>
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<tr>
<td>Ebola haemorrhagic fever</td>
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<tr>
<td>Cryptosporidiosis</td>
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<tr>
<td>West Nile virus</td>
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<tr>
<td>Cyclosporiasis</td>
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<tr>
<td>The French pox (syphilis), 1494</td>
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<tr>
<td>The American plague (yellow fever), 1793</td>
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<tr>
<td>Hueyzahuatl (smallpox), 1520</td>
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<tr>
<td>Anthrax, 1770</td>
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<tr>
<td>Cholera, 1832</td>
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<tr>
<td>HIV/AIDS, circa 1930</td>
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<tr>
<td>Spanish influenza, 1918</td>
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<tr>
<td>Measles, 1875</td>
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<tr>
<td>The Black Death (plague), 1347–50</td>
<td></td>
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<tr>
<td>The Plague of Athens (unidentified disease), 430 BC</td>
<td></td>
</tr>
<tr>
<td>Anthrax bioterrorism</td>
<td></td>
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<tr>
<td>Hantavirus pulmonary syndrome</td>
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<tr>
<td>Dengue</td>
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<td>Yellow fever</td>
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<td>HIV</td>
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<td>Lassa fever</td>
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<td>Lyme disease</td>
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<td>Hepatitis C</td>
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<td>vCJD</td>
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<td>The French pox (syphilis), 1494</td>
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<td>The American plague (yellow fever), 1793</td>
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<td>Lassa fever</td>
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<td>Lyme disease</td>
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<td>Hepatitis C</td>
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<td>vCJD</td>
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</table>
At the same time the global political situation and risk assessment constantly indicates the existing risk of intentional release of a biological, chemical or radioactive substance to cause harm. Although risk assessment and risk perception vary among the European Member States (MS), sometimes significantly, it is generally considered important to keep the awareness of this possibility in the differential diagnosis. It is important to underline that an overall strong Public Health system will be able to respond to various kinds of threats or incidents of natural (e.g. earthquakes, floods etc) or man-made (e.g. terrorist incidents) nature.

The basic principles for the PH system response include:

- Detection of a new incident
- Rapid laboratory diagnosis
- Epidemiological Investigation
- PH control measures.

Front-Line Health Professionals (FLHP) such as emergency medical services, emergency departments, and primary health care personnel, as well as regional public health personnel should be aware of the basic issues regarding new health threats to PH and RBC threats. FLHPs form the first vital link for the detection of any incident. Astute clinicians are usually the first to understand and notice new clinical syndromes (e.g. West Nile virus encephalitis) and the reporting of these to the regional PH authorities is the first step towards the recognition of the problem.

The ETHREAT scientific committee stresses the fact that generic preparedness builds strong public health systems that are able to respond to various types of threats both natural and man-made. Public Health response is based on common principles for all types of threats. The material in this handbook aims to provide FLHPs with an outline of the basic principles and knowledge regarding the high threat biological and chemical agents, as well as the principles of management of radiation exposure.

References
BASIC CONCEPTS IN HANDLING A HEALTH CRISIS

It must be noted that the basic principles that apply to the response of any type of health crisis, natural or technological, accidental or intentional are the same. For the level of a Front Line Health Professional (FLHP) in particular, only few simple facts need to be kept in mind in order to enable the management of the initial phases of such an event.

Health crisis can be defined as any event that threatens the health of citizens because of its nature, risk for spread, increased morbidity, mortality or severity.

Simple rules for crisis management include:

Defined planning: having a plan for the level of responsibility of the facility or authority is very important for the coordination of the personnel involved in their new role in the emergency response phase.

Basic planning rules include:

• Description of the roles of
  i) the different players,
  ii) the mechanism of alert and
  iii) the way of moving to the emergency phase, as well as the way back to normal function.

• Description of the function of a coordination structure (operations centre) for the hospital or facility

• Description of the back-up situation including built in redundancies (e.g. in the event of a communications or power failure)

• Training of the personnel on the implementation of the plan, as well as testing the plan via exercising at different levels.

• Communication planning: communication tools as well as a communication algorithm need to be in place as part of the preparedness plan. Crisis communication at the local level frequently involves the health care professionals.
- Good communication practices include:
  - simple, straightforward and truthful messages giving basic instructions to the public,
  - common messages across the different authorities involved at the regional level,
  - one spokesperson is preferred to deliver press communications at regular intervals.

Intersectoral collaboration: Inter-sectoral collaboration may be needed to respond to a health crisis, especially involving the deliberate release of an agent (e.g.: civil protection, law enforcement).
I Biological Agents

Information on biological agents of High Threat

Although the perception of risk for the intentional release of a biological agent to cause harm is different across the 27 EU MS, the risk is real and should not be ignored by the different levels of the health and public health sector.

Various attempts to prioritize the biological agents of highest concern for PH have been undertaken in order to assign appropriate resources for preparedness.

In the EU a specific matrix was constructed in order to be able to assess the vulnerability of the MS and the EU as a whole against various biological agents. The utility and thinking behind this tool is that provided it has updated information, each PH system can create its personalized priority list according to its own capacity and resources. An overall assessment of the threat posed by various biological agents, according to the knowledge available in 2003, is presented in the following Table I.1.

Table I.1: List of pathogens and diseases of high threat according to the EU matrix assessment

<table>
<thead>
<tr>
<th>List of diseases</th>
<th>Agents of VERY HIGH threat</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anthrax</td>
<td>Bacillus anthracis</td>
</tr>
<tr>
<td>Botulism</td>
<td>Clostridium botulinum toxin</td>
</tr>
<tr>
<td>Glanders</td>
<td>Burkholderia mallei</td>
</tr>
<tr>
<td>Haemorrhagic fever</td>
<td>Congo-Crimean virus, Ebola virus, Guanarito, Junin virus, Lassa virus, Machupo virus, Marburg virus</td>
</tr>
<tr>
<td>Plague</td>
<td>Yersinia pestis</td>
</tr>
<tr>
<td>Smallpox</td>
<td>Variola major</td>
</tr>
<tr>
<td>List of diseases</td>
<td>Agents of HIGH threat</td>
</tr>
<tr>
<td>----------------------------------</td>
<td>-----------------------</td>
</tr>
<tr>
<td>Toxic syndromes</td>
<td>Ricin, Tetrodotoxin</td>
</tr>
<tr>
<td>Tularemia</td>
<td>Francisella tularensis</td>
</tr>
<tr>
<td>Brucellosis</td>
<td>Brucella abortus, B. melitensis, B. suis</td>
</tr>
<tr>
<td>Chikungunya Fever</td>
<td>Chikungunya virus</td>
</tr>
<tr>
<td>Cholera</td>
<td>Vibrio cholerae</td>
</tr>
<tr>
<td>Coccidioidomycosis</td>
<td>Coccidioides immitis</td>
</tr>
<tr>
<td>Dysentery</td>
<td>Shigella dysenteriae</td>
</tr>
<tr>
<td>Hantavirus pulmonary syndrome</td>
<td>Hantaan virus</td>
</tr>
<tr>
<td>Haemorrhagic fever</td>
<td>Nipah, Rift Valley fever virus</td>
</tr>
<tr>
<td>Histoplasmosis</td>
<td>Histoplasma capsulatum</td>
</tr>
<tr>
<td>Hemolytic Uremic Syndrome</td>
<td>E. coli O157:H7</td>
</tr>
<tr>
<td>Influenza</td>
<td>Influenza virus (new strain)</td>
</tr>
<tr>
<td>Legionellosis</td>
<td>Legionella pneumophila</td>
</tr>
<tr>
<td>Melioidosis</td>
<td>Burkholderia pseudomallei</td>
</tr>
<tr>
<td>Meningitis</td>
<td>Neisseria meningitidis</td>
</tr>
<tr>
<td>Monkey pox fever</td>
<td>Monkey pox</td>
</tr>
<tr>
<td>Paratyphoid fever</td>
<td>Salmonella paratyphi</td>
</tr>
<tr>
<td>Psittacosis</td>
<td>Chlamydia psittaci</td>
</tr>
<tr>
<td>Q fever</td>
<td>Coxiella burnetii</td>
</tr>
<tr>
<td>Rocky mountain spotted fever</td>
<td>Rickettsia rickettai</td>
</tr>
<tr>
<td>Scrub typhus</td>
<td>Orientia tsutsugamushi</td>
</tr>
<tr>
<td>Toxic syndrome</td>
<td>Carotsaxin, Microcystin, Saotoxin, Palytoxin</td>
</tr>
<tr>
<td>Tuberculosis</td>
<td>Mycobacterium tuberculosis</td>
</tr>
<tr>
<td>Typhoid fever</td>
<td>Salmonella typhi</td>
</tr>
<tr>
<td>Typhus fever</td>
<td>Rickettsia prowazekii</td>
</tr>
</tbody>
</table>
Another important and well publicized attempt to categorize the biological threats is the one by the CDC, Atlanta USA, which according to expert consensus proposes three categories of biological agents A, B and C, according to the assessed threat to the US PH system.

### Table I.2: List of priority pathogens according to CDC, Atlanta

<table>
<thead>
<tr>
<th>Category A</th>
<th>Category B</th>
<th>Category C</th>
</tr>
</thead>
<tbody>
<tr>
<td>variola major (smallpox);</td>
<td>* Caxella burneti (Q fever);</td>
<td>Hantavirus</td>
</tr>
<tr>
<td>Bacillus anthracis (anthrax);</td>
<td>* Brucella species (brucellosis);</td>
<td>Tick-borne HF viruses</td>
</tr>
<tr>
<td>Yersinia pestis (plague);</td>
<td>* Burkholderia mallei (glanders);</td>
<td>Tick-borne Encephalitis (TBE) viruses</td>
</tr>
<tr>
<td>Clostridium botulinum toxin (botulism);</td>
<td>* Oncleaviruses,</td>
<td>Yellow fever</td>
</tr>
<tr>
<td>Francisella tularensis (tularemia);</td>
<td>* EEE, WEE;</td>
<td>Multi-drug resistant TB</td>
</tr>
<tr>
<td>filoviruses,</td>
<td>* ricin toxin from Ricinus communis (castor beans);</td>
<td></td>
</tr>
<tr>
<td>* Ebola hemorrhagic fever;</td>
<td>* epsilon toxin of Clostridium perfringens;</td>
<td></td>
</tr>
<tr>
<td>Marburg hemorrhagic fever;</td>
<td>* SEB</td>
<td></td>
</tr>
</tbody>
</table>
| arenaviruses, | * Footborne or waterborne pathogens | *
| * Lassa (Lassa fever), | * Salmonella, Shigella | |
| * Junin (Argentine HF) and related viruses | * V. cholerae | |
| | * Cryptosporidium parvum | |
Clues to keep in mind for the differentiation of natural versus man-made incidents, especially concerning a biological agent, include the following:

- Multiple clinical presentations of a disease
- Similar genetic type of agent from geographically distinct sources
- Unusual, atypical, or genetically engineered strain
- Endemic disease with unexplained increased incidence
- Simultaneous clusters of same illness in different from the usual
  - geographic areas
  - age groups
  - season
- Atypical route of transmission
- Concurrent animal disease

The following pages present an overview of the main disease presentation and management points for the biological agents of high threat from the perspective of a frontline health professional.

References
I.1 ANTHRAX

Agent
Bacillus anthracis: Gram +, rod-shaped, spore forming bacterium

Why anthrax?
Anthrax spores are extremely hardy and resistant to high temperatures, UV light and antiseptics. Anthrax has been favored as a biological weapon as it is easy to obtain, to culture and to maintain. If inhaled, it causes a serious illness with significant morbidity and mortality although it is not transmitted from person to person.

Disease and Clinical forms
Cutaneous, inhalational and gastrointestinal anthrax

Incubation Period
Usually 2-7d, (range:1-60d)

Epidemiology
Zoonosis (affects cattle); spores are very hardy and survive in the soil for decades
Human cases are rare in the EU. Cutaneous form is the most frequent naturally occurring type and a few cases are reported annually in South Europe, while it is endemic in the Middle East and Africa
Inhalational anthrax is extremely rare in the EU
Considered an occupational risk via contact with animals or animal products

Exposure
Naturally occurring human disease is usually connected to contact of abraded skin with infected animals or animal products or consumption of contaminated food
Occupational exposure for farmers, veterinarians and any profession handling animal products like leather or wool or laboratory personnel
Inhalation of the aerosolized anthrax spores during handling infected animals, in the dust of an infected area, as a laboratory accident or an act of deliberate release

**Transmission**
- No person to person transmission

**Signs and Symptoms**

**Cutaneous anthrax:**
- Usually on hands, forearms, neck, or face
- Initially small itching skin lesion (papule), which progressively enlarges and blisters
- Ulceration over 2-6 d to become a black eschar
- Marked local swelling
- Local lymphadenopathy
- Systemic symptoms: malaise, headache, chills, rarely fever

**Inhalational anthrax:**
- Febrile, flu-like illness
- Fever with marked sweating
- Malaise, myalgia
- Nausea, vomiting
- Non-productive cough
- Headache, confusion
- 1-2 days: later severe sepsis, acute dyspnea, chest pain, respiratory failure.
- Meningism
- Almost 100% fatal, if untreated. 40-60% mortality with full critical care support
Gastrointestinal anthrax:
- Severe abdominal pain (presentation similar to surgical acute abdomen)
- Nausea, vomiting
  - Bloody diarrhea
- Sepsis, shock
- High mortality even with treatment

Diagnosis of Anthrax
- Gram stain of blood, CSF or exudate
- Culture of blood, CSF, feces, tissue or exudate
- PCR testing
- Serology for antibodies
- Immunohistochemistry (skin, lymph nodes)

Management of Anthrax
- Standard precautions during hospitalisation
- Post-exposure prophylaxis: for persons, who have come in contact with suspicious packages or powder, or are involved in a deliberate release incident may be needed. (ciprofloxacin 500mg bid until environmental testing is complete or for 60d)
- Vaccine against anthrax is available for certain personnel in the UK and the USA

Treatment:
- 1st choice: Ciprofloxacin (adults: 400mg x 2 IV, children: 10mg/kg x 2 IV) or
- 2nd choice: Doxycycline (adults: 100mg x 2 IV, children >8yr: 2.2mg/kg PO x 2) or
- in combination with 1-2 additional antibiotics: clindamycin, penicillin or amoxicillin, vancomycin, rifampin, imipenem or meropenem, chloramphenicol
Anthrax Checklist for Front-line Health Professionals

Upon serious clinical suspicion of any clinical type of ANTHRAX disease:

- Protect yourself and your colleagues: standard precautions for infection control
- Inform senior clinician
  - Do you need to inform the police about a suspicious package?
- ID specialist consultation for your patient
- Report ASAP to PH services according to the established procedure in your area/region/state

A single case of inhalation anthrax in any EU MS should be fully investigated as deliberate release until proven otherwise.

References
3. CDC, Fact Sheet: Anthrax Information for Health Care Providers: http://emergency.cdc.gov/agent/anthrax/anthrax-hcp-factsheet.asp
1.2 BOTULISM

Agent
Toxin produced in 7 antigenic forms (A-G) in anaerobic conditions by the bacterium Clostridium botulinum. Gram +, rod-shaped, spore forming bacterium.

Why botulinum toxin?
Botulinum toxin is one of the most lethal substances known to man and acts by blocking the release of acetylcholine at the neuromuscular junction. It is also known to cause symptoms even in an inhaled form in laboratory accidents. Although it is hard to produce in a pure form and not transmitted from person to person, it is considered as a very high threat agent, because it causes a serious illness with significant morbidity and mortality and its antidote (botulinum antitoxin) is extremely expensive and relatively scarce in the market.

Disease and Clinical forms
Food borne, neonatal, wound botulism

Incubation Period
Usually 12-36 hrs, (range: 6 hrs-8d)

Epidemiology
- Food borne botulism is the most common type of the disease, but rather rare in Europe. Mostly caused by improper preservation or storage of food (e.g. home-canning)
- Wound botulism is reported in IV drug users in Europe and the USA, associated with black tar heroin injection
- Infant botulism results from production of toxin in the infant’s intestine from ingested Clostridium spores, sometimes associated with honey or corn syrup
Exposure

- Naturally occurring human disease is usually connected to consumption of contaminated food containing the toxin or contamination of open wound or IV drug injection site with Clostridium spores
- Inhalation of pure toxin as a laboratory accident or a deliberate release act.

Transmission

- No person to person transmission

Signs and Symptoms

All clinical forms of botulism have similar symptoms

Early Symptoms:
- No fever
- Ptosis, dry mouth, dilated pupils and sluggishly reacting to light
- 4 Ds: dysphonia, dysarthria, diplopia, dysphagia
- Alert and oriented patient
- Nausea, vomiting and diarrhea sometimes are related to food borne botulism

Late Symptoms:
- Descending paralysis progressively involving the respiratory muscles and diaphragm, in a proximal to distal pattern
- Autonomic disturbance
- Mortality is associated with airway obstruction and respiratory failure (up to 25% mortality, if left untreated, about 5% mortality with full critical care support)

Diagnosis of Botulism

- The diagnosis of botulism is clinical
- Toxin neutralization bioassay in mice
- Culture of clostridium botulinum as corroborating evidence (feces, food sample, wound swab)
- Detection of toxin in food sample/serum (<3 days after ingestion)

Management of Botulism

- Standard precautions during hospitalisation
- Toxoid vaccine available for laboratory personnel
- No post-exposure prophylaxis
Treatment:

- ABCs, and specifically respiratory support
- Critical care support: in severe cases mechanical ventilation may be needed for extended periods of time (2-8 weeks), as recovery follows the regeneration of new neuromuscular connections
  - Aminoglycosides should be avoided
- If a deliberate release incident is suspected involving aerosolized toxin:
  - Wash clothes and skin with soap and water
  - Contaminated objects or surfaces can be cleaned with 0.1% hypochlorite solution

Antitoxin:

- The decision to give botulinum antitoxin is clinical and should not await testing results
- Collect relevant specimens for testing before antitoxin infusion (serum/fees/food sample/wound swab need to reach a reference laboratory for toxin detection)
- Antitoxin is effective in reducing the severity of symptoms, if administered early.
- The currently available antitoxin is equine and states usually maintain a strategic stockpile in small quantities
- Antitoxin dose is decided in consultation with ID consultant and pharmaceutical producer
Botulism Checklist for Front-line Health Professionals

Upon serious clinical suspicion of any clinical type of BOTULISM disease:

- Protect yourself and your colleagues:
  standard precautions for infection control
- Inform senior clinician
- Do you have a suspected source for the botulism infection?
  If you suspect food borne botulism, try to collect the suspected foodstuff for testing
- ID specialist and critical care consultation for your patient
  Procurement of botulinum antitoxin
- Report ASAP to PH services, according to the established procedure in your state

A single case of clinical botulism is considered a PH emergency regardless of the suspicion of deliberate release

References

1.3 PLAGUE

Agent

Yersinia pestis: small, Gram (-), cocco bacillus

Why plaque?

Plague is transmitted by infected fleas from a rodent reservoir. Plague outbreaks contributed to massive panic in cities and countries where it appeared in the past. The agent has been favored as a biological weapon as it is relatively easy to obtain and culture, causes a serious illness with significant morbidity and mortality, which in its pneumonic form is easily transmitted from person to person. As the clinical presentation of plague is connected to the route of exposure, a deliberate release with aerosolized Yersinia pestis would cause widespread epidemic of pneumonic plague.

Disease and Clinical forms

Bubonic, pneumonic, septicemic

Incubation Period

Usually 2-4 d. (range: 1-8d)

Epidemiology

◆ Zoonosis, also considered an occupational exposure, which does not occur naturally in Europe. Cases are reported from the southwest states in the USA, Africa, India and South East Asia.

◆ Bubonic plague is the most frequent naturally occurring form of the disease.

◆ Symptoms are connected to the route of exposure
Exposure

- Naturally occurring human disease is usually connected to infected flea bites or consumption of contaminated food.
- Occupational exposure (veterinarians, hunters, laboratory personnel)
- Inhalation of the aerosolized bacteria as a laboratory accident or a deliberate release act

Transmission

- The pneumonic form is transmitted from person to person via droplets (contact <2m)

Signs and Symptoms

Bubonic Plague:
- High fever
- Swollen and quite painful regional lymph nodes (bubo), usually in groin, axilla or neck and usually unilateral
- Prostration, hypotension and confusion (may progress to pneumonic, septicemic or plague meningitis)

Pneumonic Plague:
- Fever, chills, severe malaise
- Nausea, vomiting and abdominal pain
- Cough with progressive dyspnea, chest pain and hemoptysis
- CXR findings: consolidation, infiltrates and effusion
- Progresses to shock, ARDS and respiratory failure
- Mortality 100%, if left untreated


PH10. PA chest radiograph of a patient with pneumonic plague showing both infiltrates and effusion.

PHOTO: Public Health Image Library (PHIL), CDC
Septicemic Plague:
- Fever, chills, sweats
- Gram (-) shock
- Purpura/peripheral gangrene ("black death")
- DIC
- May present as the primary form of the disease, but has usually advanced from untreated bubonic or pneumonic forms

Diagnosis of Plague
- Gram, Wright or Wayson stain of sputum/tracheal aspirate, pus or blood
- Culture of blood, pus or sputum/tracheal aspirate (preferably BEFORE antibiotics are started)
- PCR testing
- Serology for detection of antibodies

Management of Plague
- Standard and Droplet precautions during first 72 hours after initiation of treatment
- No vaccine available
- Post-exposure prophylaxis: for close contacts of a patient with pneumonic plague (ciprofloxacin 500mg bid or doxycycline 100mg bid x 7-10d)

Treatment:
- Critical care support
- 1st choice: Streptomycin (adults: 1g x 2 IM, children: 7.5mg/kg x2 IM) or Gentamicin (80mg x 3 IV)
- 2nd choice: Ciprofloxacin (adult: 400mg x 2 IV, children: 15mg/kg x 2 IV)
- Chloramphenicol is still considered 1st choice drug for plague meningitis
References


3. CDC, Plague Information Page: http://www.bt.cdc.gov/agent/plague/#fact


Plague Checklist for Front-line Health Professionals

Upon serious clinical suspicion of any clinical type of PLAGUE disease:

- Protect yourself and your colleagues:
  - standard and droplet precautions for infection control for the first 72 hours of treatment of any clinical form
  - droplet precautions for pneumonic plague
- Inform senior clinician
- ID specialist consultation for your patient
- Report ASAP to PH services, according to the established procedure in your state

A single case of plague in any EU MS is considered a PH emergency, regardless of the suspicion of deliberate release.
1.4 TULARAEMIA

Agent
Francisella tularensis: very small, Gram (-), coccobacillus with several biovars of varying pathogenicity.

Why tularaemia?
Tularaemia is transmitted by infected arthropods (fleas, mosquitoes, ticks) from a small mammal reservoir (e.g., hares, voles). Tularaemia has been favored as a biological weapon as it is relatively easy to obtain and causes a serious illness with significant morbidity. As the clinical presentation of tularaemia is connected to the route of exposure, a deliberate release with aerosolized Francisella tularensis would cause large numbers of pneumonic tularaemia.

Disease and Clinical forms
Ulceroglandular, glandular, ocuglandular, oropharangeal, pneumatic, septicemic

Incubation Period
Usually 2-5 d, (range: 1-14 d)

Epidemiology
- Zoonosis, also considered an occupational exposure, which occurs naturally in Scandinavia and Central Europe, the Americas, Asia and Australia
- Ulceroglandular tularaemia is the most frequent naturally occurring form of the disease
- Symptoms are connected to the route of exposure
Exposure
- Naturally occurring human disease is usually connected to bites from infected arthropods, handling infected animals or consuming contaminated food or water
- Occupational exposure (veterinarians, hunters, laboratory personnel)
- Inhalation of the aerosolized bacteria during handling infected animals, in the dust of an infected area, as a laboratory accident or a deliberate release act

Transmission
- No person to person transmission

Signs and Symptoms

Ulceroglandular and glandular Tularaemia:
- Fever, chills, headache, myalgia
- Ulcer at the site of inoculation, which has progressed from a papule
- Swollen and tender regional lymph nodes, usually near the insect bite/inoculation site

Oculoglandular Tularaemia:
- Results from inoculating the agent in the eye mucosa
- Fever, chills, headache, myalgia
- Unilateral painful conjunctivitis with exudates
- May present with corneal ulcer
- Periauricular regional lymphadenopathy

Oropharyngeal Tularaemia:
- Fever, chills, headache, myalgia
- Sore throat
- Exudative tonsillitis/pharyngitis/stomatitis
- Cervical lymphadenopathy

Pneumonic Tularaemia:
- Follows inhalation of organism or is secondary from other site
- Fever, chills, severe malaise, sore throat
- Cough with dyspnea, chest pain
- CXR findings: variable but may show infiltrates, hilar adenopathy and effusion
- Progresses to respiratory failure and death
- Mortality 30%, if left untreated
Septicemic Tularaemia:
- Usually secondary from other site
- Fever, chills, headache, myalgia
- Nausea, vomiting, diarrhea, abdominal pain
- Confusion, coma
- Septic shock, ARDS
- DIC and haemorrhage

Diagnosis of Tularaemia
- Gram stain of sputum, pharyngeal or ulcer exudate swab
- Culture of blood, sputum/tracheal aspirate, wound (preferably BEFORE antibiotics are started - multiple sets are needed as organism is hard to grow)
- PCR testing
- Serology for detection of antibodies.

Management of Tularaemia
- Standard precautions during hospitalisation
- Vaccine available for laboratory personnel
- Post-exposure prophylaxis: for persons involved in a deliberate release incident (ciprofloxacin 500mg bid or doxycycline 100mg bid x 14d)

Treatment:
- 1st choice: Streptomycin (1g x 2 IM) or Gentamicin (80mg x 3 IV)
- 2nd choice: Ciprofloxacin (adults: 400mg x 2 IV, children: 5mg/kg x 2 IV)
- Relapses are common
Tularaemia Checklist for Front-line Health Professionals

Upon serious clinical suspicion of any clinical type of TULARAEMIA disease:

- Protect yourself and your colleagues: standard precautions for infection control
- Inform senior clinician
- ID specialist consultation for your patient
- Report ASAP to PH services, according to the established procedure in your state

In some EU member states a single case of tularaemia may be a PH emergency

References
3. CDC. Tularaemia Information Page: http://www.bt.cdc.gov/agent/tularemia
I.5 VIRAL HEMORRHAGIC FEVERS

Agents
Viruses from families: arena-, filo, bunya- and flaviviruses

Why VHF?
Viral hemorrhagic fevers describe a heterogeneous family of diseases with the common clinical presentation of fever and bleeding diathesis. Some of these diseases carry significant morbidity and mortality (nearing 90% in the case of Ebola fever), along with the possibility of person-to-person transmission.

Disease and Clinical forms
Multiple VHF such as: Marburg fever, Lassa fever, Ebola fever, Machupo (Bolivian hemorrhagic fever), Guanarito (Venezuelan hemorrhagic fever), Junin (Argentinian hemorrhagic fever), Rift Valley fever, Crimean-Congo hemorrhagic fever (CCHF), yellow fever, dengue and others.

Incubation Period
Varies depending on the individual virus, range: 1-21d

Epidemiology
◆ All are zoonoses with varying geographical distribution depending on their animal reservoir.
◆ The vast majority of VHFsd does not occur naturally in any EU member state, with the exception of CCHF, which occurs in South East Europe.
Exposure

- Naturally occurring human disease is connected to bites from infected arthropods (mosquito or tick), inhaling infected dust, or consuming contaminated primate meat, depending on the life cycle of each virus.
- Occupational exposure for healthcare or laboratory personnel
- Inhalation of the aerosolized virus particles in the dust of an infected area, as a laboratory accident or a deliberate release act

Transmission

- Person to person transmission through contact with secretions from symptomatic patients (saliva, diarrhea, vomit, urine or blood) or needle sticks
- Asymptomatic patients are rarely infectious

Signs and Symptoms

All VHFs exhibit a prodrome with fever, malaise, headache, myalgia, nausea and vomiting.

Ebola/Marburg Hemorrhagic fevers:
- Abrupt onset of febrile prodrome
- Prostration
- Diarrhea (bloody), vomiting
- Maculopapular rash (3-8 d after onset)
- Bleeding
- Confusion, coma, multiple organ failure
- Ulcer at the site of inoculation, which has progressed from a papule
- Swollen and tender regional lymph nodes, usually near the insect bite/inoculation site
- Mortality 30-90% in the recent epidemics in central Africa

Lassa Fever:
- Slow onset febrile prodrome
- Prostration
- Sore throat, conjunctivitis, face edema and chest pain
- Vomiting, diarrhea
- Bleeding
- Effusions (pleural, ascites), encephalopathy
- Mortality <30%, residual sensorineural deafness common
CCHF:
• Abrupt onset febrile prodrome
• Vomiting, diarrhea, abdominal pain
• Sore throat, conjunctivitis
• Lethargy, face edema
• Petechiae and bleeding (usually after day 4)
• Hepatomegaly, encephalopathy
• Mortality in epidemics 30-50%

Diagnosis of VHF
- Viral culture of blood, tissue, sputum/tracheal aspirate (Important: the viral culture of these agents is possible only in a BSL-4 laboratory)
- PCR testing
- Serology for detection of antibodies

Management of VHF
- Isolation of patient
- Possible need for High Security Isolation Unit admission and hospitalisation in a negative pressure room, by specialized personnel
- Standard, contact and airborne precautions
- No vaccine available
- Post-exposure prophylaxis: ribavirin PO for exposure to Lassa fever or other arenavirus and observation for 21 d

Treatment:
- Supportive care
- Ribavirin for Lassa and CCHF (adults: 30mg/kg IV initial dose, followed by 15mg/kg TID x 4d, 7.5mg/kg IV x 6d)
VHF Checklist for Front-line Health Professionals

Upon serious clinical suspicion of any VHF:

☑ Protect yourself and your colleagues:
   - Isolate suspected patient
   - Standard, contact and droplet precautions for infection control

☑ Inform senior clinician

☑ Urgent ID specialist consultation
   - Decision on need for airborne precautions
   - Decision on need for admission in high security isolation unit (under negative pressure ventilation)

☑ Report ASAP to PH services, according to the established procedure in your state
   - A single case of VHF is a PH emergency and warrants full PH investigation regardless of the suspicion of deliberate release

References
3. CDC, Viral Hemorrhagic Fever Information Page: http://emergency.cdc.gov/agent/vhf/#related
1.6 VIRAL ENCEPHALITIDES

Agents
Viruses from various families: togaviruses, flaviviruses, arenaviruses, bunyaviruses

Why encephalitis?
Viral encephalitides describe a heterogeneous family of diseases with the common clinical presentation of fever and encephalitis syndrome (confusion, paralysis, convulsions etc). Some of these diseases carry significant morbidity and/or mortality. In particular Venezuelan Equine Encephalitis (VEE) has been studied in programs of biological weapon production in the past as an incapacitating agent. The VEE virus can be readily aerosolized and infects almost 100% of exposed.

Disease and Clinical forms
Multiple encephalitides such as: Venezuelan Equine Encephalitis (VEE), Eastern Equine (EEE), and Western Equine Encephalitis (WEE), St. Louis Encephalitis (SLE), Japanese Encephalitis, and others.

Incubation Period
Varies depending on the individual virus, range: 1-15d

Epidemiology
- Mostly zoonoses with varying geographical distribution depending on their animal reservoir
- Some encephalitides occur in Europe, such as Tick-borne complex Encephalitis (TBE) in central Europe, West Nile Encephalitis in South Europe

Exposure
- Naturally occurring human disease is connected to bites from infected arthropods (EEE, WEE, VEE and WNV by Aedes or Culex mosquitoes, TBE by ticks), to exposure to tissues/secretions of infected animals or to inhalation of infected dust
Occupational exposure for laboratory personnel

Inhalation of the aerosolized virus particles in the dust of an infected area, as a laboratory accident or a deliberate release act

Transmission
- No person to person transmission

Signs and Symptoms
- Infection may be asymptomatic or cause non-specific influenza-like illness

Venezuelan Equine Encephalitis (VEE):
- Incubation 2-6d
- Fever, headache, malaise
- Sore throat
- Nausea, vomiting, diarrhea
- Headache and photophobia
- May recover or deteriorate
- Severe headache and backache
- Prostration
- Confusion, altered mental status and coma
- Convulsions, ataxia, paralysis
- Mortality up to 20%, children tend to exhibit severe form in outbreaks and survivors have high frequency of neurological sequelae

West Nile virus Encephalitis (WNV):
- Incubation 3-14d
- Fever, chills, malaise
- Rash
- Nausea, vomiting, diarrhea
- Headache and photophobia
- Cough
- Confusion, altered mental status and coma
- Mortality 12-14% in encephalitis patients and higher in elderly with predisposing morbidity, residual neurological sequelae in >50% of survivors

Tick-borne Encephalitis (TBE):
- Incubation 7-14 d
- Influenza-like, febrile prodrome with headache and myalgia
- Asymptomatic phase lasting 1-3 d to 3wks
- Paralysis
- Confusion, coma, encephalitis
- Mortality may be 4-25%
Diagnosis of Viral Encephalitis
- Viral culture from blood, tissue, CSF
- PCR or RT-PCR testing
- Serology for detection of antibodies

Management of VHF
- Standard precautions for hospitalisation
- There are vaccines available for some viruses (such as TBE and JE)

Treatment:
- Supportive care

Encephalitis Checklist for Front-line Health Professionals

Upon serious clinical suspicion of any viral encephalitis:
- Protect yourself and your colleagues:
  Standard precautions for infection control
- Inform senior clinician
- ID specialist consultation
- Report ASAP to PH services, according to the established procedure in your state
  In most EU MS viral encephalitides are not endemic, although occasional cases may be diagnosed

References
1.7 Q FEVER

Agent
Rickettsia Coxiella burnetii. Gram (-), pleomorphic, intracellular coccobacillus

Why Q Fever?
While Q fever is not a lethal disease but rather an incapacitating one, it is considered a high threat agent as very few C. burnetii organisms are able to cause human infection. As an agent it is quite hardy, resistant to heat, dryness and many disinfectants. It has been known to cause natural outbreaks in wide areas via exposure to contaminated dust from infected animals.

Disease and Clinical forms
Infection may be asymptomatic, acute or chronic

Incubation Period
Usually 18-21 d, range: 4-40 d

Epidemiology
- Zoonosis with worldwide geographical distribution and reservoirs in sheep, cattle, domesticated pets, small mammals, pigeons and ticks
- Animals are asymptomatic but shed the agent in feces, urine, reproductive fluids and placentas

Exposure
- Naturally occurring human disease is connected to exposure to tissues/secretions of infected animals or to inhalation of infected dust and rarely to arthropod bites (ticks)
- Occupational exposure for farmers, veterinarians, laboratory personnel
- Inhalation of the aerosolized agent in the dust of an infected area, as a laboratory accident or a deliberate release act.
Transmission
- No person to person transmission

Signs and Symptoms

Acute disease
- High fever, chills and sweats
- Headache, confusion and lethargy
- Myalgia and pharyngitis
- Nausea, vomiting, diarrhea and abdominal pain
- Cough, chest pain, pneumonia (25%)
- Hepatitis

Chronic disease
- Fever, weight loss, malaise
- Aseptic meningitis/meningoencephalitis
- Endocarditis (75%)

Diagnosis of Q Fever
- Serology for detection of antibodies
- Culture from blood, tissue (difficult)
- PCR testing
- Immunohistochemistry

Management of Q Fever
- Standard precautions for hospitalisation
- No licensed vaccine available
- Post-exposure prophylaxis: for persons involved in a deliberate release incident (doxycycline 100mg PO BID x 7d or co-trimoxazole 960mg PO BID x 7d)

Treatment:
- Supportive care
- 1st choice: Doxycycline (adults: 100mg IV/PO BID x 14-21d, children >8yr: 2.2mg/kg BID x 14-21d)
- 2nd choice: Co-trimoxazole or fluoroquinolones
- In chronic disease multi-drug treatment protocol is needed with doxycycline in combination with fluoroquinolone or doxycycline and hydroxychloroquine
Q Fever Checklist for Front-line Health Professionals

Upon serious clinical suspicion of Q FEVER:

- Protect yourself and your colleagues:
  - Standard precautions for infection control
- Inform senior clinician
- ID specialist consultation
- Report ASAP to PH services, according to the established procedure in your state

References
I.8 GLANDERS

Agent
Burkholderia mallei, small Gram (-) bacillus

Why glanders?
Glanders is quite rare and was considered a bioweapon in the World Wars I and II, when horses and mules were vital parts of the movement of personnel and equipment. It is considered a high threat agent as it can infect humans via inhalation but at the same time there is currently very little clinical experience in recognizing and treating the disease.

Disease and Clinical forms
Localized, pulmonary, septicemic, chronic

Incubation Period
Usually 10-14 d

Epidemiology
- Zoonosis mainly affecting horses, donkeys, mules but also goats, dogs and cats
- Endemic in Africa, Asia, Central and South America, while in the EU it has not been reported since the '40s
- Immuno-compromised patients are more susceptible

Exposure
- Naturally occurring human disease is connected to exposure to infected animals or their carcasses and products
- Occupational exposure for farmers, veterinarians, abattoir workers, laboratory personnel
- Inhalation of the aerosolized agent in the process of handling infected animals, as a laboratory accident or an act of deliberate release
Transmission

- Person to person transmission very rare

Signs and Symptoms

Localized disease

- Fever, chills and sweats
- Headache, myalgia
- Local and generalized pustular rash that ulcerates
- Lymphadenopathy
- Nasal discharge

Pulmonary glanders

- Fever, chills and sweats
- Headache, myalgia
- Cough, chest pain, dyspnea
- CXR: multifocal consolidation, effusion, cavitiation, lung abscess

Septicemic glanders

- Fever, chills and sweats
- Headache, myalgia
- Septic shock
- Multiple abscesses (liver, kidney, spleen)
- Multi organ failure

Diagnosis of Glanders

- Culture from blood, sputum, pus, urine

Management of Glanders

- Standard precautions for hospitalisation
- No vaccine available
- Post-exposure prophylaxis: for persons involved in a deliberate release incident (doxycycline 100mg PO BID x 7d or co-trimoxazole 960mg PO BID x 7d)

Treatment:

- Supportive care
- 1st choice: Ceftazidime (adults: 2g IV TID x 14d, children: 20mg/kg TID x 14d)
- 2nd choice: Meropenem/Imipenem (1g IV TID x 14d)
- 3rd choice: Gentamicin (8mg/kg IV QD x 14d) and Co-trimoxazole (8/40mg/kg/d PO x 14d)
  to be continued to complete 20 weeks with
  - doxycycline PO and co-trimoxazole PO or
  - amoxicillin/clavulanic acid PO
- Relapses are common, long term follow up is needed
Glanders Checklist for Front-line Health Professionals

Upon serious clinical suspicion of GLANDERS:

- Protect yourself and your colleagues: Standard precautions for infection control
- Inform senior clinician
- Exclude immuno-compromised staff from patient care
- ID specialist consultation
- Report ASAP to PH services, according to the established procedure in your state

A single case of glanders may be a PH emergency regardless of suspicion of deliberate release

References

3. CDC, Glanders page: http://www.cdc.gov/nczved/dfbmd/disease_listing/glanders_ti.html
1.9 MELIOIDOSIS

Agent
Burkholderia pseudomallei
small Gram (-) bacillus

Why melioidosis?
Melioidosis is quite rare in Europe, <10 imported cases are reported annually. It is considered a high threat agent as it can infect humans via inhalation, it causes a high morbidity and mortality disease but at the same time there is currently very little clinical experience in recognizing and treating it.

Disease and Clinical forms
Localized, pulmonary, septicemic, chronic

Incubation Period
Usually 1-21 d (range: 1 day- years after exposure)

Epidemiology
• Melioidosis occurs in South and SE Asia, Northern Australia, but also in Africa and the Americas.
• Immuno-compromised patients are more susceptible

Exposure
• Naturally occurring human disease is connected to contaminated water exposure through skin lesions or consumption/aspiration of contaminated water
• Occupational exposure for farmers, rice workers and laboratory personnel
• Inhalation of the aerosolized agent in the process of work in rice fields, as a laboratory accident or a deliberate release act
Transmission

- Person to person transmission is very rare, occurring via contact with blood and patient fluids

Signs and Symptoms

Skin, soft tissue disease
- Fever, chills and sweats
- Headache, myalgia
- Subcutaneous nodules
- Multiple soft tissue abscesses
- Local lymphadenopathy
- Skin pustules

Pulmonary melioidosis
- Fever, chills and sweats
- Headache, myalgia
- Cough, chest pain, dyspnea
- CXR: multifocal consolidation, effusion, cavitation, lung abscesses

Septicemic melioidosis
- Fever, chills and sweats
- Headache, myalgia
- Septic shock
- Multiple abscesses(liver, kidney, spleen, brain)
- Multi organ failure
- Mortality 100%, if untreated and 40% with treatment

Diagnosis of Melioidosis
- Culture from blood, sputum, pus, urine
- Serology for detection of antibodies
Management of Melioidosis

- Standard precautions for hospitalisation
- No vaccine available
- Post-exposure prophylaxis: for persons involved in a deliberate release incident (doxycycline 100mg PO BID x 7d or co-trimoxazole 960mg PO BID x 7d)

Treatment:

- Supportive care
- 1st choice: Ceftazidime (adults: 2g IV TID x 14d, children:120mg/kg TID x 14d)
- 2nd choice: Meropenem/Imipenem (1g IV TID x 14d) to be continued to complete 20 weeks with
  - doxycycline PO and co-trimoxazole PO or
  - amoxicillin/clavulanic acid PO
- Relapses are common, long term follow up is needed

Melioidosis Checklist for Front-line Health Professionals

Upon serious clinical suspicion of MELIOIDOSIS:

- Protect yourself and your colleagues: Standard precautions for infection control
- Inform senior clinician: Exclude immuno-compromised staff from patient care
- ID specialist consultation
- Report ASAP to PH services, according to the established procedure in your state
  - A case of melioidosis merits full investigation regardless of suspicion of deliberate release
References


3. CDC, Melioidosis page: http://www.cdc.gov/ncezid/dfhdbd/disease_listing/melioidosis_g.htm

I.10 SMALLPOX

Agent
Variola virus: the largest DNA virus

Why smallpox?
Smallpox (variola) is the only known communicable disease that has been eradicated from our earth through international PH cooperation and collaboration. The eradication meant also interruption of the vaccination programs and loss of clinical expertise for the recognition and management of the disease. Smallpox has caused significant outbreaks in the past, claiming the lives of millions of people and it is known that the virus was studied in bioweapons programs in the past. The modern society is deemed highly susceptible to this virus, which has a very low infectious dose (10-100 virions) and causes a disease with significant morbidity and mortality. As the period of contagiousness is quite long, the disease is able to be readily transmitted from person to person.

Disease and Clinical forms
• Variola major, which may present as ordinary, modified, flat and hemorrhagic smallpox
• Variola minor

Incubation Period
12-14d, range: 7-17d

Epidemiology
• Smallpox does not exist as a natural disease since 1977. In most EU member states the last cases were reported in the ‘40s or ‘50s
• The smallpox virus is only stockpiled in two laboratories in the world, in the USA and the Russian Federation
• There is no known animal reservoir for smallpox
Exposure

- Currently there is no naturally occurring human disease
- Inhalation of the aerosolized virus particles as a laboratory accident or an act of deliberate release

Transmission

- Person to person transmission through prolonged face-to-face contact, contact with patient fluids (saliva, vesicle fluid, scabs) or contaminated objects such as clothing
- Few reported cases with airborne transmission from patients with significant cough

Signs and Symptoms

Variola Major:

- **Ordinary Smallpox (>90%)**
  - Abrupt onset febrile prodrome with high fever, headache and backache (2-4 d)
  - Prostration
  - Enanthem (rash) in the mouth marks the beginning of the contagious period
- **Maculopapular rash (+2-4 d after fever)** that spreads to the whole body (24hrs) but appears to have centrifugal distribution and may involve palms and soles
- The rash gradually but uniformly develops to vesicles (3-5d after onset), then deep embedded pustules with central dimple (6-12 d after onset)
- Gradually the rash crusted over and scabs form, which fall off about 3 weeks after the appearance of the rash leaving scars
- Only after the scabs have fallen off, the patient is not contagious anymore
- Average mortality 25-30%, highest in infants and the elderly
Uncommon forms of Smallpox (about 5% each):

- Abrupt onset febrile prodrome
- Modified: fewer lesions, faster evolution of the rash maybe without the pustular phase
- Hemorrhagic: Rash becomes hemorrhagic and DIC develops
- Flat: the rash remains flat and soft, was reported in infants
- Historically mortality 95-100% for the flat and hemorrhagic types

Variola Minor

- Clinically undistinguishable from variola major
- Fewer systemic symptoms
- Fewer lesions without residual scarring
- Mortality about 1%

Diagnosis of Smallpox

- Clinical picture is quite specific, but new generations of medical personnel have no clinical experience with this disease any more. The most important fact is to differentiate from varicella (chickenpox)
- Electron Microscopy for orthopoxvirus recognition
- PCR testing
- Viral culture of vesicle fluid, sputum/tracheal aspirate (Important: the viral culture of smallpox is possible only in a BSL-4 laboratory)
- Serology for detection of antibodies
### Table I.10: Differential diagnosis of Smallpox (Variola) and Chickenpox (Varicella)

<table>
<thead>
<tr>
<th></th>
<th>Chickenpox (Varicella)</th>
<th>Smallpox (Variola)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Incubation Period</strong></td>
<td>14-21d</td>
<td>12-14d</td>
</tr>
<tr>
<td><strong>Febrile prodrome</strong></td>
<td>None to mild</td>
<td>2-4d duration: Severe headache, backache</td>
</tr>
<tr>
<td><strong>Rash</strong></td>
<td>Centripetal: dense on trunk, less on face and extremities</td>
<td>Centrifugal: dense on face and extremities, less on trunk</td>
</tr>
<tr>
<td></td>
<td>Itchy rash, evolves quickly in crops from superficial papules to vesicles to pustules</td>
<td>uniformly develops to vesicles (1-2d), deep embedded pustules (+5-14d)</td>
</tr>
<tr>
<td></td>
<td>Usually spares palms and soles</td>
<td>Present in palms and soles</td>
</tr>
<tr>
<td></td>
<td>Rash dries quickly and scabs fall (&lt;1d)</td>
<td>Rash dries (+10-14d) and scabs fall off (+14-28d) slowly</td>
</tr>
<tr>
<td><strong>Agent</strong></td>
<td>DNA herpes virus</td>
<td>DNA orthopox virus</td>
</tr>
<tr>
<td><strong>Treatment</strong></td>
<td>Acyclovir is effective, if given&lt;72hrs</td>
<td>Supportive care</td>
</tr>
</tbody>
</table>

**Note**: Comparative evaluation of the rash in smallpox and chickenpox.
Management of Smallpox

- Isolation of patient.
- Possible need for High Security Isolation Unit admission and hospitalisation in a negative pressure room, by specialized personnel.
- Standard, contact and airborne precautions.
- Vaccinia vaccine is available and effective (95%), but its use is connected with significant side effects.
- Trained and vaccinated medical or emergency response personnel exist in EU member states according to their individual national planning.
- Vaccinia vaccine can also be effective as a post-exposure measure, if given soon (<7 days) after exposure. Therefore close contacts of patients should be immediately traced, isolated and vaccinated.

Treatment

- Supportive care
Smallpox Checklist for Front-line Health Professionals

Upon serious clinical suspicion of SMALLPOX:

- Protect yourself and your colleagues:
  Isolate suspected patient
  Standard, contact and droplet precautions for infection control

- Inform senior clinician
  Trained or vaccinated staff only to care for this patient
  Exclude immuno-compromised staff from the care of this patient
  Follow protocols according to existing national smallpox management plan

- Urgent ID specialist consultation
  Exclusion of varicella (chickenpox) and herpes virus infection
  Decision on need for airborne precautions
  Decision on need for admission in high security isolation unit (under negative pressure ventilation)
  Follow protocols according to existing national smallpox management plan

- Report IMMEDIATELY to PH services, according to the established procedure in your state

- A single case of SMALLPOX is a global PH emergency and warrants full PH investigation

References
   http://www.hpa.org.uk/
   http://www.eurosurveillance.org/ViewArticle.aspx?ArticleId=502
Although this chapter will focus in providing a concise overview of the four basic guidelines for infection control, which are needed to respond to biological agents of high threat, it needs to be noted that infection control also involves all the processes, the mentality and the organisational framework that prevents the transmission of diseases.

- from patient to patient in the health care setting
- from health care workers to patients
- from patients to health care workers

Drawing from this definition infection control is not just about the four types of precautions used by health care workers for patients with certain diseases. It includes:

- working practices (hand hygiene, sharp disposal etc)
- training and availability of PPE, as well as monitoring of PPE use
- environmental measures (disinfection, laundering etc)
- patient placement protocols
- isolation options (appropriately ventilated rooms etc)
- monitoring of health care workers for occupational injuries, vaccination status, prophylaxis etc

In the following pages the four basic infection control types are outlined:

- Standard Precautions
- Droplet Precautions
- Contact Precautions
- Airborne Precautions
Standard Precautions

Wash hands
- Before and After any patient contact
- Before and After any medical procedure on a patient and contact with patient secretions
- Before and After you eat or touch your face
- After removing your gloves
- After handling/touching equipment
- After using the lavatory

Wear gloves
- for invasive procedures
- for any anticipated contact with potentially infectious body fluids or material

Change gloves after procedure, wash hands before attending next patient

Eye and face protection, if there is risk of spray or splash to your eyes, nose and mouth

Wear a gown or disposable plastic apron, appropriate to the task, to protect skin and prevent soiling or contamination of clothing during procedures

Dispose of sharps and soiled PPE appropriately

Respiratory Hygiene/Cough Etiquette
- For patients, accompanying persons and Health Care Workers

- Standard Precautions apply to ALL PATIENTS, for ALL STAFF, at ALL SETTINGS regardless of suspected or confirmed infection status, ALL THE TIME
- Select your PPE according to the extent of anticipated blood, body fluid, or pathogen exposure

Attention!
Contact Precautions
in addition to Standard Precautions

- A single patient room is preferred
- Limit transport and movement of patients outside of the room to medically-necessary purposes
- Instruct patient to wear a mask and follow respiratory hygiene/cough etiquette depending on respiratory symptoms and diagnosis
- Single-use disposable patient equipment preferred

Wash Hands

- Before and After any patient contact
- Before and After any medical procedure on a patient and contact with patient secretions
- Before and After you eat or touch your face
- After removing your gloves
- After handling/touching equipment
- After using the lavatory

Use running water and antimicrobial soap, count slowly up to 50 or use alcohol-based hand rub

Wear Gloves

- Wear gloves upon entry into patient room
- Change gloves after procedure
- Remove gloves before exiting the patient room

Single-use (latex, vinyl or nitrile)

Eye and Face Protection, if there is risk of spray or splash to your eyes, nose and mouth

Use single-use surgical mask or mask with face shield or mask and goggles

If not available consult infection control team

Ensure proper decontamination of regular equipment and environment
Contact Precautions in addition to Standard Precautions cont’d

Always wear a gown, or disposable plastic apron (depending on patient’s symptoms) upon entry into the patient room

Dispose of sharps and soiled PPE appropriately

Respiratory Hygiene/Cough Etiquette

For patients, accompanying persons and Health Care Workers

Attention! • Remember to wash your hands after removing PPE

Contact Precautions

Are intended for agents that spread via
• direct contact with a patient or
• indirect contact with contaminated environment, especially when there is
  • wound drainage,
  • diarrhea or incontinence
  • other secretions.

Some examples where Contact Precautions are needed:

- SARS
- Multi-drug resistant bacteria (e.g. MRSA, VRE)
- Salmonella, Shigella infections
- RSV infections
- Skin infestations (e.g. scabies, lice)
- C. difficile infections
- Rotavirus infections
- Hepatitis A infection
Droplet Precautions
in addition to Standard Precautions

- A single patient room is preferred
- In multi-patient rooms, cohorting is advisable and a distance 1-2m between patients is desirable.
- Instruct patient to always wear a mask and follow respiratory hygiene/cough etiquette
- Limit transport and movement of patients outside of the room to medically-necessary purposes.
- Single-use disposable patient equipment preferred

Wash hands
- Before and After any patient contact
- Before and After any medical procedure on a patient and contact with patient secretions
- Before and After you eat or touch your face
- After removing your gloves
- After handling/touching equipment
- After using the lavatory

Use running water and antimicrobial soap, count slowly up to 50 or use alcohol-based hand rub

Wear gloves
- Wear gloves upon entry into patient room
- Change gloves after procedure
- Remove gloves before exiting the patient room

Single use (latex, vinyl or nitril)

Eye and face protection
- Always wear a mask upon entry into the patient room or patient contact <2m
- Eye protection if there is risk of spray or splash to your eyes, nose and mouth

Patient to wear single-use surgical mask

Ensure proper decontamination of regular equipment and environment

If not available consult infection control team
Droplet Precautions are intended to prevent transmission of pathogens spread through close respiratory or mucous membrane contact with respiratory secretions. Respiratory droplets are large-particle droplets >5μ in size that are generated by a patient who is coughing, sneezing or talking. Due to their large size, respiratory droplets are not able to travel long distances airborne and fall with 1-2 m from the patient.

Examples where droplet precautions are needed include:
- SARS
- influenza
- pneumonic plague
- monkeypox
- Mycoplasma pneumoniae
- adenovirus
- pertussis (whooping cough)
- group A streptococcal infections
- rubella
- meningococcal infections (Neisseria meningitidis)
- Parvovirus B19 infections
- mumps
- rubella
- meningococcal infections (Neisseria meningitidis)

Always wear a gown, or disposable plastic apron upon entry into the patient room.

Dispose of sharps and soiled PPE appropriately.

Respiratory Hygiene/Cough Etiquette
- patient and accompanying person to wear a mask.

- Donning PPE sequence: gown, mask, goggles, gloves
- Remove PPE before exiting the patient room, according to the order shown above
- Remember to wash your hands after removing PPE

Droplet Precautions in addition to Standard Precautions cont'd

• Donning PPE sequence: gown, mask, goggles, gloves
• Remove PPE before exiting the patient room, according to the order shown above
• Remember to wash your hands after removing PPE

Attention!
Airborne Precautions
in addition to Standard Precautions

- Single patient room with negative pressure ventilation
- Restrict entry to essential personnel and visitors
- Limit transport and movement of patients outside the room to medically necessary purposes
- Single-use disposable patient equipment preferred

If not available consult infection control team
All entering room to wear appropriate PPE
Patient to wear single-use surgical mask and follow respiratory hygiene/cough etiquette
Ensure proper decontamination of regular equipment and environment

Wash hands
- Before and After any patient contact
- Before and After any medical procedure on a patient and contact with patient secretions
- Before and After you eat or touch your face
- After removing your gloves
- After handling/touching equipment
- After using the lavatory

Use running water and soap, count slowly up to 50
or
Use alcohol-based hand rub

Wear gloves
- Single use (latex, vinyl or nitril)
- Change gloves after procedure
- Remove gloves before exiting the patient room

Eye and face protection
- Always wear a mask and
- Always wear eye protection (goggles) upon entry into the patient room

Use properly fitted FFP3 respirator
Wear appropriate dermal protection; water resistant surgical gown or whole-body suit from water resistant material.

Dispose of sharps and soiled PPE appropriately.

Remove PPE before exiting the patient room.

**Attention!**

- Donning PPE sequence: gown, FFP3 respirator, goggles, gloves
- Removing PPE sequence as follows:
  - Step 1: Remove gloves
  - Step 2: Remove goggles/visor
  - Step 3: Remove gown/suit by peeling
  - Step 4: Remove respirator

Remember to wash your hands after removing PPE.

Airborne Precautions are needed for agents that are transmitted via core respiratory particles (<5μm) that can remain suspended in air and travel long distances with air currents or through ventilation systems in a building. Examples where droplet precautions are needed include:

- Rubeola virus [measles]
- Varicella virus [chickenpox]
- Mycobacterium tuberculosis
- SARS-CoV, in some instances
- Smallpox, in some instances
- Viral haemorrhagic fevers (VHFs) in the last stages of clinical disease

**References**

III CHEMICAL AGENTS

Chemical agents of High Threat

There are reports of chemical substance use in war in the histories of many civilizations such as the Chinese, the Greeks and Byzantium.

Unfortunately World War I marks the landmark use of gas warfare in such extent that approximately 1 million of the 26 million casualties of this conflict is attributed to use of chemical warfare. Chlorine, phosgene, cyanogenes and vesicants were used in battlefields across Europe by both sides and caused significant number of deaths and injuries with residual effects in the armed forces.

In the years between World War I and World War II, new chemical weapons and protective gear were developed but were not used in battlefields. Nevertheless all of the major nations involved in WW I and several other countries developed chemical weapons programs. Cyanide was consistently used in the Nazi concentration camps.

Throughout the remainder of the 20th century, chemical weaponry continued to develop and used in major wars around the world. During the Persian Gulf War, the alleged use of nerve and mustard agents by Iraqi troops has been reported. Following that war, there have been reports of chemical agent use ordered by Saddam Hussein against Kurd and Shiite Muslims as well as proliferation of chemical weapons development in Libya.

The Chemical Weapons Convention treaty was finalized in 1993 and is currently signed by more than 140 countries around the world. It prohibits the development, production, stockpiling, and use of chemical weapons and provides for the verification and destruction of known stockpiles.

Despite diplomatic efforts, chemical weapons will remain a threat in warfare and have become a potential weapon of the terrorist in recent years. In 1994, a religious cult, Aum Shinrikyo, released nerve gas in a residential area in Matsumoto, Japan, and in 1995 the same group undertook the well-publicized sarin attack in the Tokyo subway.

Chemicals are manufactured daily in huge quantity by the industry and many substances are highly toxic and dangerous for humans, animals let alone the environment. The ease with which chemical substances are accessible and may be manufactured into weapons increases the concern that they may be used for terrorist purposes and cause mass casualties.

The following pages present a brief overview of the background, the symptoms and the management of the classes of chemical agents considered of highest threat.
Agents
Phosgene, chlorine, ammonia, hydrogen sulphide, hydrogen chloride

Why pulmonary agents?
Phosgene and chlorine were among the agents widely used during WWI causing thousands of deaths and incapacitated among the troops. Currently many of these agents are widely used in the modern chemical industries (plastics, pesticides) and therefore stored or transported through communities. This class of agents is the most commonly involved in industrial chemical releases.

Odor
- Phosgene: white to pale yellow gas, smells like mown grass or musty hay
- Chlorine: greenish gas, smells like bleach
- Hydrogen sulphide: smells like rotten egg

Latent Period
Usually minutes to 48hrs

Mechanism of Action
- These are gases that are heavier than air and therefore accumulate in low-lying areas.
- Reaction with water in mucous membranes and production of corrosive substances, such as hydrochloric acid (chlorine, phosgene) or nitric acid (ammonia)
- Destruction of the alveolar-capillary membrane of the respiratory tract and leak of fluid in the interstitial tissue, resulting in ARDS

Exposure
- Inhalation of gas
- Skin and eye contact with gas
- In a liquid form the agents may contaminate water or food and people can be exposed via consumption

Effects and clinical syndrome
Damage depends on the water solubility and direct tissue reactivity, the dose and the duration of exposure to each agent.

Early:
- Eye irritation and tearing with blurred vision, chemical conjunctivitis or corneal injury
- Nose and throat irritation
- Skin irritation and burn-like lesions, when in contact with liquid
- Cough, choking, chest pain, chest pressure
- Nausea and vomiting
- Laryngeal edema in massive exposures
Effects and clinical syndrome of pulmonary agent exposure

Latent:
- Dyspnea on exertion progressing to dyspnea at rest
- Bronchoospasm, frothy or blood tinged sputum
- Hypoxia and pulmonary edema
- ARDS

Diagnosis of pulmonary agent exposure
- The diagnosis of exposure to a pulmonary agent is CLINICAL

Management
Prompt decontamination of exposed patients is recommended as follows:
- Remove patient from the agent/area of exposure and
- Remove agent from the patient
  - Remove clothes and dispose appropriately in double, sealed bags
  - Wash patient’s skin with warm soap water or 0.1% hypochlorite solution
  - Remove contact lenses and irrigate eyes thoroughly with NaCl 0.9%, check for corneal injury
  - Do not induce emesis

Treatment
- NO specific antidote
- ABCs and supportive care, with frequent reassessments
- Aggressive respiratory support: administration of high flow O₂, inhaled bronchodilators and corticosteroids
- Bed rest and observation at least for 24hrs
Upon serious clinical suspicion of exposure to PULMONARY AGENTS:

- Protect yourself and your colleagues:
  - Make sure that patient is decontaminated  
  - Wear appropriate PPE
- Inform senior clinician
  - Poison centre
  - Pulmonary medicine
  - Ophthalmology
- Report ASAP to PH, law enforcement or other services (e.g. civil protection) according to the established procedure in your state

References
   http://www.hpa.org.uk/
3. CDC, Chlorine and Phosgene Information Pages:
   http://emergency.cdc.gov/agent/chlorine/ and
   http://emergency.cdc.gov/agent/phosgene/basics/facts.asp
Agents
Cyanide, cyanogen chloride, cyanide salts (sodium or potassium cyanide)

Why cyanide?
Cyanide was manufactured as a chemical weapon in large quantities between WWI and II, and was used in concentration camps and in the Iran-Iraq war in the '80s. Cyanide has been used often for assassinations and suicides. Cyanide is widely used in the industry (plastics, fertilizers, photography) and is also a combustion product in house fires, considered to play a significant role in smoke inhalation morbidity.

Odor
• Colorless gas or white solids, smells of bitter almonds

Note: only some people are genetically able to smell it

Latent Period
Immediate, seconds to minutes

Mechanism of Action
• Usually gases that are lighter than air and very volatile liquid or solid salts
• Chemical asphyxiants are agents that replace oxygen in the hemoglobin molecule and inhibit oxygen transport to the cells causing tissue hypoxia
• Some cyanide salts may also be corrosive to skin and eyes

Exposure
- Inhalation of gas
- Skin and eye contact with liquid or solid
- Ingestion
**Effects and clinical syndrome**

Damage depends on route of exposure, concentration and duration of exposure

**Severe Exposure:**
- Gagging
- Convulsions
- Coma and dilated pupils
- Respiratory arrest
- Sudden collapse and death

**Mild Exposure:**
- Eye and nose irritation from cyanogens
- Dizziness, headache
- Nausea, with vomiting in moderate exposure
- Confusion and agitation
- Dyspnea, chest tightness
- Persistent hypotension and acidemia, despite good arterial oxygen levels

**Diagnosis of cyanide exposure**

- Blood cyanide level

**Note:** the decision to administer antidote is CLINICAL and should not await test results

**Management**

- Prompt decontamination of exposed patients is recommended as follows:
  - Remove patient from the agent/area of exposure and
  - Remove agent from the patient
    - Remove clothes and dispose appropriately in double, sealed bags
    - Wash patient’s skin with warm soap water or 0.1% hypochlorite solution
  - Remove contact lenses and irrigate eyes thoroughly with NaCl 0.9%, check for corneal injury
  - Do not induce emesis, consider gastric lavage or activated charcoal PO for ingestions
Treatment

- Administer Cyanide Antidote IMMEDIATELY IF PATIENT SYMPTOMATIC
  - Altered mental status (GCS<8) and/or
  - Respiratory depression

- Cyanide Antidote
  - Hydrosocobalamin 5% (CyanoKit©): 5g IV diluted in 0.9%NaCl over 15min. Dose may be repeated once, if severe exposure (max 10g)
  - Dicobalt Edetate: 300mg IV over 1 minute followed by 50ml glucose 50%
  - Amyl Nitrite with Sodium Nitrate and Sodium Thiosulphate (Cyanide Antidote Package): 1 ampule Amyl Nitrite broken and inhaled in front of the face of the patient every 1min, until able to administer 300mg (or 10ml of 3%) sodium nitrite IV over 5-20 minutes, followed by 12.5g (or 25ml of 50%) sodium thiosulphate IV over 10 minutes
  - Need simultaneous blood pressure monitoring
- ABCs and specifically administration of 100% O₂
- Supportive care, ECG and pulse oximetry monitoring

Cyanide Checklist for Front-line Health Professionals

Upon serious clinical suspicion of exposure to CYANIDE:

- Protect yourself and your colleagues:
  - Make sure that patient is decontaminated
  - Wear appropriate PPE
- Administer ANTIDOTE
- Inform senior clinician
  - Poison centre
  - Critical Care Medicine
  - Consultations may be necessary
- Report ASAP to PH, law enforcement or other services (e.g. civil protection) according to the established procedure in your state
References

3. CDC, Cyanide Information Page: http://emergency.cdc.gov/agent/cyanide
III.3 VESICANTS (Blister Agents)

Agents
Mustards (nitrogen and sulphur), organic arsenicals (Lewisite, Phosgene Oxime)

Why vesicants?
Vesicants were manufactured as chemical weapons in large quantities between WWI and II, and mustards were used in the Iran-Iraq war in the 80s. These agents are not used in industry, but quantities of them still exist in the arsenals of various countries and are under the process of destruction or have been dumped in the sea and frequently show up on the shores or in fishermen’s nets in the Baltic Sea.

Odor
- Lewisite: may smell of geraniums
- Mustards: may smell like garlic, fresh onion or mustard

Latent Period
- Immediate for Lewisite
- 4-12 hrs for mustards

Mechanism of Action
- Oily volatile liquids, pale yellow to amber that in gas form are heavier than air and accumulate in low-lying areas
- Vesicants cause tissue damage by alkylation, similarly to radiation, affecting all rapidly replicating cells

Exposure
- Inhalation of gas
- Absorption through intact skin
- Absorption through eye contact with liquid or gas
Effects and clinical syndrome

Damage depends on concentration and duration of exposure, humidity and environmental temperature.

Eyes:
- Eye irritation: tearing, redness, blepharospasm, photophobia
- Periorbital edema
- Corneal ulceration and clouding
- Temporary or permanent loss of vision
- Globe perforation may complicate severe eye exposure to liquid form

Respiratory System:
- Runny nose with burning pain, ulceration of nose and oropharynx
- Loss of voice
- Cough, mucosal sloughing, dyspnea, hemoptysis
- Fever
- Chemical pneumonitis and ARDS
- Chemical pneumonitis and ARDS are the most common cause of death

Gastrointestinal tract:
- Nausea, vomiting, diarrhea

Systemic Effects:
- Arsenic toxicity after exposure to Lewisite (liver failure, nephritis, neuropathy, hemolysis, encephalopathy)
- Bone marrow depression and pancytopenia, leading to secondary bacterial infections in the skin or respiratory tract
- CNS depression
- Cardiac arrhythmias

Note: skin blisters do not contain mustard

PH31. Tearing, redness and periorbital edema in a patient exposed to mustard

PH32. Skin lesions in patients exposed to mustard

PH31. Tearing, redness and periorbital edema in a patient exposed to mustard

PH32. Skin lesions in patients exposed to mustard
Diagnosis of vesicant exposure

- The diagnosis is CLINICAL
  - Urine mustard metabolites (thiodiglycol) may be measured in specialised laboratories
  - Urine arsenic after suspected exposure to Lewisite

  **Note:** laboratory tests can be used at a later stage to CONFIRM exposure and should not delay treatment or treatment decisions

Management

- Decontamination is CRITICAL:
  - Remove patient from the agent/area of exposure and
  - Remove agent from the patient
    - Remove clothes and dispose appropriately in double, sealed bags
    - Wash patient’s skin with warm soap water or 0.1% hypochlorite solution
    - Remove contact lenses and irrigate eyes thoroughly with NaCl 0.9%, check for corneal injury
    - Do not cause emesis in cases of ingestion.

Treatment

- There is NO antidote for mustards
- There is a specific antidote for Lewisite only, if there is clinical suspicion of exposure and
  - Pulmonary edema
  - Chemical burn with history of late decontamination (>15 min from exposure)
  - Skin damage >5% BSA:
    - Administer Dimercaprol or British Anti-Lewisite (BAL): 3mg-5mg/kg by deep IM injection every 4 hours for 4 doses
    - Alternatives: 2, 3-DMSA and 2, 3-DMPS
- ABC and supportive care, intensive respiratory support
- Do not patch eyes: atropine eye drops for blepharospasm and ophthalmic ointment to prevent eyelids from sticking together
- Burn care for the skin damage: analgesia, debridement, dressings
Vesicant Checklist for Front-line Health Professionals

Upon serious clinical suspicion of exposure to VESICANTS:

✔ Protect yourself and your colleagues:
    Make sure that patient is decontaminated or
    Wear appropriate PPE

✔ Inform senior clinician
    Do you need to activate emergency response plan in your hospital?
    Poison centre
    Critical Care Medicine (Burn Unit)
    Plastic Surgery
    Ophthalmology

✔ Report ASAP to PH, law enforcement or other services (e.g. civil protection)
  according to the established procedure in your state

References
3. CDC, Blister Agents/Vesicants Information Page: http://emergency.cdc.gov/agent/vesicants
Agents

G-agents (tabun, sarin, soman, cyclosarin), V-agents (VX, Russian VX)

Why nerve agents?
Nerve agents are extremely toxic chemical weapons and were manufactured as chemical weapons in large quantities between WWI and II. These agents were used in the Iran-Iraq war in the '80s, as well as in terrorist attacks in Matsumoto (1994) and Tokyo (1996) in Japan. These agents are not used in industry, but quantities of them still exist in the arsenals of various countries and are under the process of destruction. Organophosphate pesticides in general have been banned from use in the EU, but accidental or deliberate (e.g. suicide) exposures are not uncommon in almost all EU MS.

Odor
- G-agents: clear, colorless liquids, odorless or may smell fruity
- V-agents: brown oily liquid at room temperature, odorless

Latent Period
- Immediate

Mechanism of Action
- Volatile liquids, colorless to brown at room temperature, vapors are heavier than air and accumulate in low lying areas
- Nerve agents act similarly to organophosphate pesticides by inhibiting acetylcholinesterase enzymes, causing extreme cholinergic stimulation of CNS, and peripheral muscarinic and nicotinic receptors by the accumulating acetylcholine.
Exposure
- Inhalation of gas or aerosol
- Absorption through intact skin
- Absorption through eye contact with liquid or gas

Effects and clinical syndrome
Damage depends on route, dose and duration of exposure

Severe Exposure:
- Pinpoint pupils
- Confusion, agitation
- Convulsions
- Increased respiratory secretions
- Cardiac arrhythmias
- Respiratory arrest and coma
- Death

Mild to Moderate Exposure:
- Pinpoint pupils, red eyes, lachrymation (tearing), blurred vision
- Dizziness, confusion, headache
- Sneezing, coughing, bronchorrhea, wheezing, dyspnea
- Drooling, abdominal cramping, vomiting, diarrhea, urination
- Muscle twitching/tremors, muscle weakness and eventually paralysis
- Tachycardia and hypertension

Note: progression of symptoms should alert you for continued exposure, inadequate decontamination or inadequate treatment

Diagnosis of nerve agent exposure
- The diagnosis is CLINICAL
  - Red cell (RBC) cholinesterase activity
  - Plasma cholinesterase

Note: laboratory tests can be used at a later stage to CONFIRM exposure and should not delay treatment or treatment decisions

Management
- Decontamination is CRITICAL:
  - Remove patient from the agent/area of exposure and
  - Remove agent from the patient
    - Remove clothes and dispose appropriately in double, sealed bags
    - Wash patient's skin with warm soap water or 0.1% hypochlorite solution
    - Remove contact lenses and irrigate eyes thoroughly with NaCl 0.9%.
    - Do not induce emesis in cases of ingestion.

Note: inadequate decontamination may cause continuing exposure and progression of symptoms for the patient, as well as secondary cases in the emergency response and medical personnel
There is specific antidote for nerve agents, which is lifesaving if administered:
- **Atropine**: (2 mg for adults or 0.05 to 0.1 mg/kg for children, IM/IV every 5-10 min, titrating with respiratory secretions and dyspnea)
- **Pralidoxime**: 600-1800 mg IM or 1 g infusion over 30 min
- **Benzodiazepines**: Diazepam (5 to 10 mg in adults and 0.2 to 0.5 mg/kg in children), for seizures as needed

**Note 1**: treating physicians should be able to recognize and treat possible atropinization

**Note 2**: organophosphate poisoning may demand large amounts of atropine

**Note 3**: pralidoxime should be administered ASAP, esp. if nerve agent is suspected, to avoid the “aging” of the agent

- ABC and supportive care, intensive respiratory support

### Nerve Agent Checklist for Front-line Health Professionals

- Upon serious clinical suspicion of exposure to NERVE AGENTS or ORGANOPHOSPHATES:
  - Protect yourself and your colleagues: Make sure that patient is decontaminated or Wear appropriate PPE
  - Inform senior clinician
  - Poison centre
  - Critical Care Medicine consultations may be necessary
  - Ophthalmology
  - Report ASAP to PH, law enforcement or other services (e.g., civil protection) according to the established procedure in your state
References
   http://www.hpa.org.uk/web/HPAweb&Page&HPAwebAutoListName/Page/1158934607980
3. CDC, Nerve Agents Information Page: http://emergency.cdc.gov/agent/nerve
III.5 RIOT CONTROL AGENTS (incapacitating agents, tear gas)

Agents
BZ, hallucinogens (LSD), lachrymators (CS, CN, Chloropicrin, pepper spray, mace), vomiting agents (adamante)

Why riot control agents?
Riot control agents are frequently used by law enforcement around the world for crowd control in demonstrations or for capturing missions. Although not lethal, except in specific circumstances such as confined spaces and for patients with pre-existing morbidity, they can be adequately incapacitating and cause significant mass morbidity.

Odor
Usually white odorless agents, tear gases may smell like pepper or apples

Latent Period
• Immediate (secs to min)

Mechanism of Action
• BZ: anticholinergic agent, antagonist of muscarinic receptors
• Hallucinogens: 5-HT serotonin receptor agonist, action on sympathetic system
• Opioids: liquid, synthetic opioids
• Lachrymators: irritants to mucosa and skin
• Vomiting agents: action on CNS centre for vomiting

Exposure
• Inhalation of spray/dust/aerosolized agent
• Ingestion (rare)
Effects and clinical syndromes
Damage depends on concentration and duration of exposure

- **BZ**: latent period 30min-24hrs, mydriasis and blurred vision, agitation, disorientation and hallucinations, initially tachy- then bradycardia, flushing, hyperthermia, dry skin and mucous membranes, ataxia

- **Hallucinogens**: latent period min-12hrs, visual, acoustic and tactile hallucinations, confusion, hyperthermia, vertigo, ataxia, vomiting, tachycardia, hypertension

- **Opioids**: miosis, seizures and chest rigidity, respiratory depression/arrest, loss of consciousness, coma

- **Lachrymators**: latent period few min, eye and mucous membrane burning, excessive tearing, blepharospasm, corneal ulceration, skin irritation, blistering in prolonged exposure, painful runny nose, loss of voice, salivation, chest tightness, ARDS in severe exposure

- **Vomiting agents**: latent period few min, eye and mucous membrane irritation, burning and tearing, vomiting and malaise for hours

Diagnosis of incapacitating agent exposure
- The diagnosis is CLINICAL
- Opioids and BZ can be detected in the urine

Management
- Decontamination:
  - Remove patient from the agent/area of exposure and
  - Remove agent from the patient
  - Remove clothes and dispose appropriately
  - Wash patient's skin with warm soap water

  **Note**: hypochlorite solution exacerbates symptoms from lachrymators

- Remove contact lenses and irrigate eyes thoroughly with NaCl 0.9%, check for corneal injury
Treatment

- BZ: physostigmine 30-45mg/kg IM/IV, repeat as needed depending on mental status, every 60 min.
  
  **Note**: test dose 1mg IM/IV if diagnosis doubtful. IV administration may cause significant bradycardia.

- Hallucinogens: benzodiazepines (e.g. diazepam, lorazepam) IV as needed.

- Opioids: naloxone 0.2-0.4 mg IM/IV and repeat as needed.

- Lachrymators and Vomiting agents: there is NO antidote for these agents. Topical hydrocortisone for skin irritation and antihistamines or burn care if blistering. Respiratory support. Symptoms last for a few hours at the most.

- ABC and supportive care.

Incapacitating Agent Checklist for Front-line Health Professionals

Upon serious clinical suspicion of exposure to INCAPACITATING AGENTS:

- **Protect yourself and your colleagues:**
  - Make sure that patient is decontaminated
  - Wear appropriate PPE

- **Inform senior clinician**
  - Poison centre
  - Ophthalmology consultations may be necessary

- **Report ASAP to PH, law enforcement or other services (e.g. civil protection)** according to the established procedure in your state.
References
Agents

Toxins are proteins or peptides produced by various organisms as part of their defense mechanisms or other physiologic process:

- Botulinum toxin from bacterium Clostridium botulinum (see I.2 p24-27)
- Staphylococcal Enterotoxin B (SEB) from bacterium Staphylococcus aureus
- Ricin and abrin from castor oil (Ricinus communis) and rosary pea (Abrus precatorius) plants respectively
- Terodotoxin, saxitoxin and conotoxin from marine organisms (puffer fish and marine snails)
- Mycotoxins produced from various fungus species such as Fusarium sp, Aspergillus or Penicillium

Why toxins?

Toxins are in general proteinic substances, which are lethal to humans in very small quantities. They are considered as very high threat agents, although they are found in nature in minute quantities and they are in general hard to produce. Toxins produce nonspecific clinical syndromes which are hard to diagnose, with significant morbidity and mortality and have no known (or hard to obtain) antidotes. Their victims require high-quality critical care support which is extremely costly. Finally toxins pose a serious diagnostic challenge, as they are difficult to diagnose either clinically or in a laboratory. Little expertise is available at European or international level.

Odor

Odorless and tasteless substances

Latent Period

- Immediate to delayed in hrs (see below)
Mechanism of Action

- **SEB**: latent period 1-6hrs, exotoxin, super-antigen, activation of cytokine reaction, shock
- **Ricin and Abrin**: latent period depends on the route of exposure, for ingestion ranges typically between 6-12hrs, may be less for inhalation (4-6hrs). Both are water soluble, tasteless glycoproteins, cell poisons inhibiting ribosome or mitochondrial function and protein synthesis
- **Tetrodotoxin and Saxitoxin**: latent period minutes to a few hours, neurotoxins, inhibit Na+ influx intracellularly and disrupt nerve conduction
- **Mycotoxins**: latent period minutes from skin exposure or 3-12 hrs from ingestion, inhibit protein synthesis, which is followed by a secondary disruption of DNA and RNA synthesis and skin irritant

Epidemiology

- **SEB**: commonly associated with food poisoning from S. aureus but due to its stability, was studied as an incapacitating bioweapon
- **Ricin and Abrin**: while the castor oil plant has a worldwide distribution, the rosary plant is native to Indonesia and grows in tropical and subtropical areas of the world, and sporadic cases of poisoning are reported from consumption of its fruit
- **Tetrodotoxin and Saxitoxin**: the toxin is produced by bacteria contaminating the intestines of the puffer fish, which live in the Atlantic and Pacific tropical and temperate waters. Sporadic cases of tetrodotoxin food-borne poisoning are reported annually in the Far East, where these fish are considered particular delicacies
- **Mycotoxins**: Trichothecenes may be contaminants of grain crops such as wheat or corn, and are detected in environmental samples in buildings affected by mould, but are also considered incapacitating chemical weapons

Exposure

- Consumption of contaminated food (e.g. fish) or plant fruit/seeds (e.g. castor beans or rosary peas)
- Skin contamination for mycotoxins or contact with the toxin producing organism
- Injection of the toxin in an attempt for assassination
- Inhalation of aerosolized toxins as a laboratory accident or an act of deliberate release

Transmission

- No person to person transmission
Signs and Symptoms

SEB Toxin Syndrome:
- Non-specific flu like illness,
- Headache, myalgia
- High fever with chills
- Dyspnea, chest pain, cough, may progress to respiratory failure and ARDS (especially if inhaled)
- Multiple organ failure and septic shock in high level exposure

Ricin & Abrin Toxic Syndrome:
- Fever
- Cough, dyspnea, chest tightness
- Myalgia, arthralgia
- Non cardiogenic pulmonary edema, respiratory failure and ARDS
- Abdominal pain, cramps
- Nausea, vomiting and bloody diarrhea related to ingestion
- Hematuria, liver failure
- Hypovolemia, DIC, multiple organ failure
- Death in 36-72 hrs

Tetrodotoxin & Saxitoxin:
- Oral paresthesias progressing to arms and legs
- Respiratory failure due to
- Cranial nerve dysfunction
- Weakness progressing to paralysis, causing respiratory failure and death
Trichothecene Mycotoxins:
• Skin and mucous membrane burning pain
• Skin vesicles progressing to necrosis
• Nausea, vomiting, diarrhea,
• Cough, dyspnea progressing to respiratory insufficiency and ARDS
• Bleeding diathesis
• Convulsions and coma

Diagnosis of Toxins

☐ The diagnosis of most toxin related syndromes is clinical.
☐ Ricin: antibody fluorescent immunoassay and PCR in environmental samples, measurement of ricinine in urine by Mass Spectrometry.
☐ Detection of SEB, Mycotoxins or Tetrodotoxin in food sample/serum/urine in specialized laboratories.

Management of Toxin-related Syndromes

☐ Standard precautions during hospitalisation
☐ Decontamination if a deliberate release incident is suspected involving aerosolized toxin:
  • Remove clothes and dispose appropriately in double, sealed bags
  • Wash patient's skin with warm soap water or 0.1% hypochlorite solution
  • Remove contact lenses and irrigate eyes thoroughly with NaCl 0.9%
  • Contaminated objects or surfaces can be cleaned with 0.1% hypochlorite solution

☐ No vaccines available
☐ No post-exposure prophylaxis available

Treatment

☐ ABCs
☐ Critical care support
☐ Collect relevant specimens for testing, which need to reach a reference laboratory for toxin detection

No vaccines available
No post-exposure prophylaxis available
Toxin Checklist for Front-line Health Professionals

Upon serious clinical suspicion of any clinical TOXIN syndrome:

✔ Protect yourself and your colleagues:
  standard precautions for infection control

✔ Inform senior clinician:
  If you suspect food borne toxin syndrome, try to collect the suspected foodstuff for testing
  ID specialist
  critical care
  Poison centre/toxicology consultations may be needed for your patient

✔ Report ASAP to PH services, according to the established procedure in your region/state:
  A single case of toxin poisoning in any EU MS is considered a PH emergency and should be investigated fully regardless of the suspicion of deliberate release

References

2. CDC, Ricin Information Page: http://emergency.cdc.gov/agent/ricin/
Puffer fish producing tetrodotoxin

Ricinus communis, a fungus that produces trichothece mycotoxins.
III.7 TOXIC INDUSTRIAL CHEMICALS (TIC)

Agents
A wide variety of substances used in large quantities in the chemical industry, such as acids, ammonia, bases, chlorine, other inorganic substances. These commonly also include carbon disulfide, allyl alcohol, hydrazine, nitrobenzene, di-nitro-toluene, hydrogen sulphide, chloride and many others.

Exposure
Depending on the agent and the type of exposure (occupational accident or accidental environmental release) any route is possible (inhalation, dermal, eye, ingestion).

Signs and Symptoms
Mainly respiratory irritation, burns (skin, ocular or respiratory tract), anxiety

Diagnosis of Exposure to TICs
Diagnosis is mainly clinical

Management of related Syndromes
- Decontamination at the site of accident or release is preferable especially, if heavy exposure to liquid is suspected involving any chemical agent:
  - Remove clothes and dispose appropriately in double, sealed bags
  - Wash patient’s skin with warm soap water or 0.1% hypochlorite solution
  - Remove contact lenses and irrigate eyes thoroughly with 0.9% NaCl solution
Treatment

- ABCs
- Critical care support
- Symptomatic treatment
- In some cases antidotes are available e.g. calcium gluconate and corticosteroids for hydrofluoric acid

TIC Checklist for Front-line Health Professionals

Upon clinical suspicion or information of exposure to TIC:

✔ Protect yourself and your colleagues:
   Select appropriate Protective Equipment
   May need proper skin protection (water resistant PPE)

✔ Inform senior clinician
   Critical care consultations may be needed for your patient
   Poison centre/toxicology (tox database, material safety data sheets (MSDS) of the agent involved, medical management advice)

✔ Report ASAP to Poison Centre or other responsible agency (e.g. civil protection), according to the established procedure in your region/state

References

   http://www.hpa.org.uk/
Basic Information and Terms

Radiation is a type of ionizing energy, emitted by certain materials which cannot be detected by human senses. It is important to differentiate between a nuclear and a radiological event:

- A nuclear event results from the fusion or fission of atoms, which produces a significant and highly destructive wave of heat, light and radiation.
- A radiological event may involve an explosion and release of generally smaller amounts of radiation compared to a nuclear event.

The radiation injury suffered by persons involved in a radiological incident depends on:

- The dose of radiation received
- The type of radiation (alpha, beta or gamma)
- Whether the exposure involves internal or external contamination

Cells that multiply regularly are in general more sensitive to radiation, which in turn means that some body organs, like the bone marrow, are more radiosensitive compared to others.

Why radiological threats?

All types of radioactive sources or material used for industrial or medical purposes pose radiological threat and may cause a radiological emergency. Such events may involve the misuse of abandoned sources, transport emergencies, accidental leaks or spills of radioactive material or intentional use of radioactive material in conjunction with explosives (Radiation Dispersion Device (RDD) or “dirty bomb”).

Latent Period

Minutes to days after the exposure, depending on the dose of radiation absorbed.
Mechanism of Action

• Direct: radiation acts directly on tissues and causes biological changes

• Indirect: radiation acts on tissue water which becomes ionized and by creating free radicals it binds to proteins, enzymes and other molecules and causes biological changes

The effects of radiation on live cells are

• Stochastic: where the dose is related to the increasing possibility of occurrence of an effect (cancerogenesis, genetic effects).

• Non-stochastic: which are directly dose dependant and outlined in Table IV.1.

Exposure

• Radiation exposure occurs when this particular type of energy penetrates the human body to cause its effects.

• Factors determining the exposure to radiation are mainly:
  - **Time**: shorter time means shorter exposure
  - **Distance**: the longer the distance from the source means less exposure
  - **Shielding**: barrier between the body and the source means less exposure.

• Internal contamination signifies inhalation, ingestion or contamination of open wounds with radioactive dust or other material.

• External contamination implies the existence of radioactive dust or other material on the skin, hair or clothing of the exposed person.

Acute radiation syndrome (ARS)

• Acute radiation exposure in high dose, usually in short time, on large body surface area, to penetrating radiation results in acute radiation syndrome.

• Symptoms occur in 4 phases: prodromal, latent, illness, recovery/death and depend on the amount of the absorbed radiation dose.

The following tables IV.1 and IV.2 present an overview of the symptoms per radiation dose and a clinical score for the prediction of outcome of radiation exposed patients.
Table IV.1: Effects and Symptoms according to the dose of exposure

<table>
<thead>
<tr>
<th>Dose in Sv¹</th>
<th>Effects and Symptoms</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.005 (5mSv)</td>
<td>Asymptomatic</td>
</tr>
<tr>
<td>0.1 (100 mSv)</td>
<td>Asymptomatic, minimum dose detected by chromosome analysis</td>
</tr>
<tr>
<td>0.5 (500 mSv)</td>
<td>Asymptomatic, recurrent nausea and vomiting in &lt;10% exposed, transient lymphocyte and platelet depression</td>
</tr>
<tr>
<td>1.0</td>
<td>15% presents recurrent nausea and vomiting within 2d after exposure</td>
</tr>
<tr>
<td>2.0</td>
<td>Nausea and vomiting in the majority of exposed, anemia, skin damage (itching, erythema), hair loss</td>
</tr>
<tr>
<td>4.0</td>
<td>Nausea, vomiting and diarrhea within 48hrs, skin damage (burns), hair loss, serious bone marrow depression (hematopoietic syndrome), 50% mortality without treatment</td>
</tr>
<tr>
<td>6.0</td>
<td>Severe hematopoietic syndrome, 100% mortality within 30d without treatment, 50% mortality with treatment</td>
</tr>
<tr>
<td>7.0</td>
<td>Gastrointestinal syndrome: early nausea, vomiting, fatigue, severe bone marrow depression, death in 2-3 weeks</td>
</tr>
<tr>
<td>&gt;20.0</td>
<td>CNS syndrome: immediate explosive vomiting and diarrhea, headache, altered consciousness, coma, convulsions, shock, death in 24-72 hrs</td>
</tr>
</tbody>
</table>

¹ Equivalent dose measured in Sievert (Sv), to account for the different effects of the different types of radiation (equivalent dose = absorbed dose x radiation weighting factor), where 1 Sv = 1 Gy for gamma rays.
Table IV.2: Clinical Score for patient management in the first 48hrs after exposure

The score assumes whole body or large parts of the body external exposure. Therapeutical Management according to the European Consensus Conference (EBMT, European group for Bone Marrow Transplantation, Ulm University-Germany and IRS, Institut de Radioprotection et de Sûreté Nucléaire-France).

### Diagnosis of radiation exposure

- Radiation can be detected only by specific equipment, e.g. Geiger counter.
- Radiation detected on nasal or oral swabs indicate internal contamination by inhalation or ingestion.
- Complete Blood Count (CBC) with white cell differential (immediately as baseline and then every 4hrs for 12hrs followed by every 6hrs for 48hrs) to monitor absolute lymphocyte count (Andrews Lymphocyte Normogram, to predict severity of radiation injury).
Management

- **Decontamination:**
  - Remove patient from the agent/area of exposure
  - Remove agent from the patient
    - Survey patient with appropriate radiation detection equipment, document result
      - Carefully remove clothes and dispose appropriately in double, sealed bags
    - Repeat patient survey with radiation detection equipment, document result
      - Wash patient’s skin carefully with warm soap water or 0.1% hypochlorite solution
      - Remove contact lenses and irrigate eyes thoroughly with NaCl 0.9% solution, check for corneal injury
    - Repeat patient survey with radiation detection equipment, document result
  
  **Note 1:** may need to repeat washing until radiation level is twice the background or remains unchanged.

  **Note 2:** decontamination of contaminated victims should be done as soon as possible BUT it should not delay/interfere with life threatening interventions. (Fig. IV.1).

  **Note 3:** persons who have been exposed to radiation but are NOT externally contaminated do not require decontamination.
Treatment

- ABCs
- In the event of internal contamination
  - Specific agents may be available to dilute or compete with their radioactive counterpart or help eliminate specific radionuclides (e.g., potassium iodide for radioactive iodine (I131) to protect the thyroid gland, Prussian Blue for radioactive thallium or cesium), DTPA for plutonium and americium, deferoxamine for plutonium and iron, stable strontium, calcium, zinc and iron to compete with the respective radioactive elements
  - Diuresis helps with the removal of tritium, as well as radioactive sodium and potassium
  - Gastric lavage and cathartics may also be needed for the excretion of the radioactive agent
- Frequent reassessment and monitoring blood tests, as per experts
- Supportive care:
  - Analgesia,
  - Symptomatic treatment for nausea, vomiting and diarrhea,
  - Aggressive prevention/treatment of infections,
  - Use of hematopoietic growth factors.
- In whole body exposure, surgeries need to be performed within 48hrs or after recovery of the bone marrow
Fig. IV.1: Algorithm for the management of persons involved in a radiological emergency

Life-threatening wounds and burns should be treated first. Irradiation is not contamination – An irradiated person is not a source of radiation.

- **External Contamination?**
  - **NO**
  - **YES**

- **ABCs and stabilization**

- **Internal Contamination?**
  - **NO**
  - **YES**

  - **External Contamination?**
    - **NO**
    - **YES**

    - **Specific Diagnostic Tests Expert advice**

    - **Clinical picture indicates severe exposure?**
      - **NO**
      - **YES**

      - **Decontamination**

      - **Supportive Care**

      - **Expert Advice**
Radiation Emergency Checklist for Front-line Health Professionals

Upon serious clinical suspicion of exposure to RADIATION:

✔ Protect yourself and your colleagues:
  Make sure patient is decontaminated BUT do not delay ABCs
  Need for radiation detection equipment
  Standard precautions are usually enough
  FFP3 masks are preferred

✔ Inform senior clinician
  Hospital Radiation Safety
  Haematology
  Consultations may be necessary

✔ Report ASAP to PH services and Radiation Protection Authority
  according to the established procedure in your state

References
This chapter aims to give a brief overview of the process and the issues involved in the decontamination of persons exposed to biological or chemical agents or radioactive dust for the Front Line Health Professional (FLHP). It is underlined that decontamination of exposed persons follows the same basic principles for all types of agents in accidental or deliberate exposures. Exposure to biological agents is usually not perceived by humans and it remains to be detected by the clinical syndrome developed after the incubation period of the particular agent. In this case the patients presenting in the health care facilities do not need to undergo decontamination. In particular cases though, where there is a believable threat (e.g. powder sent by mail) there may be need to also decontaminate persons exposed to biological agents.

In the case of an overt incident involving a chemical or radiological threat a command and control structure will need to be rapidly established. The first act of the incident commander is to establish control of the site for the protection of attending emergency services, casualties and the wider public. This involves the establishment of an exclusion zone ("the hot zone"), where there is a risk of direct contamination from chemical exposure. It is therefore accessed only by specifically trained rescue personnel wearing gas tight (also referred to as Type A) PPE and self-contained breathing apparatus (SCBA).

Casualties are brought by the rescue team to the contamination reduction zone ("warm zone") where decontamination takes place. The risk in this zone is from indirect exposure to chemicals from contaminated casualties and equipment. Therefore, workers in this zone (ambulance, paramedical and medical staff) require gas tight PPE. Following decontamination, casualties enter the support zone ("cold" zone) before transportation to hospital, if required. The command post as well as any medical-aid and decontamination stations are always established upwind, uphill and up water of the incident, to prevent dissemination of the agent throughout the environment.
further contamination or exposure of the operating personnel. It needs to be noted that ambulatory patients from any incident (accidental or terrorist) may present to the closest medical facility and therefore they always present a challenge for the control of the incident and the proper response of the health sector in the area.

All patients exposed to RBC agents through accidental or intentional release should be decontaminated before entering the Emergency Department or any treatment area in order to protect

- The health care staff
- The other patients in the hospital
- The hospital environment, avoiding its contamination

The decontamination area should be prepared in advance or created ad hoc, but it needs to provide on one hand some privacy and on the other full access to running water. Various solutions are available for portable or permanent structures for the health care facilities and

The single most important step for the decontamination of patients exposed to biological, chemical or radioactive material is the prompt and complete removal of the patient’s clothing. In order to achieve this promptly and smoothly FLHPs should consider the need for triage of the exposed persons before decontamination primarily in ambulatory and non-ambulatory patients and secondly in the groups according to the triage system used in the particular health care facility. A simple overview presenting the principles of the START triage system is presented in the following chapter.
Non-ambulatory patients
Health care professionals need to assist or perform the decontamination of these patients in a different line from the ambulatory patients.

Ambulatory patients
Special considerations for ambulatory patients include:

- Provide clear instructions and a number of health professionals to oversee the process, especially if you are dealing with multiple patients.
- Provide privacy in order to unclothe.
  - Separation of sexes may be needed, especially if people with different religious beliefs are involved in the incident.
  - Avoid separation of family members to minimize their anxiety.
- The process of applying soap and rinsing should each last at least 5-10 minutes.
- Provide clothing for the decontaminated, “clean” victims.

V.2 Basic Principles for human decontamination

- Remove clothes carefully.
  - **Attention:** you may need to cut the clothes off in non-ambulatory victims.
- Double-bag clothes and seal them.
- Wash skin carefully and thoroughly with copious amounts of warm water and soap.
  - **Attention:** skin folds (e.g., axilla, under breasts etc).
  - **Attention:** irrigate eyes and open wounds with normal saline (0.9% NaCl) solution to avoid irritation.

V.3 Special Considerations for the Decontamination Process

- Removal of clothes is usually adequate decontamination of patients exposed to gases, with the exception of patients exposed to nerve agents.
- Remove first any solid particles (dust or larger particles) from naked skin by sweeping, before rinsing with water.
- In case of decontamination of persons exposed to a biological agent, the removal of clothing should be careful so as not to create aerosol or alternatively it should be done under running water.
• Decontamination of patients exposed to mercury
  - Mercury is highly toxic and volatile, even a household or laboratory thermometer contains dangerous quantities of mercury.
  - Removal of contaminated clothing needs to be very careful to avoid inhalation by the patient and the assisting team.

• Decontamination of radioactive material (dust or particles)
  - Need for Geiger counter to assess complete decontamination.
  - Some emergency response services or emergency departments may have available chemical detection equipment to assess complete decontamination for certain chemical agents.
Mass casualty situations create a discrepancy between the medical resources available for treatment and the number of casualties. Depending on the capacity of each healthcare facility the threshold number of casualties varies widely.

In mass casualty situations it is frequently assumed by the authorities that:
- there is prompt formation of a unified command for the on-site management
- there is coordinated transport of patients from the site to the receiving health care facilities
- first aid is given by medical and paramedical personnel.

But in reality the situation develops as follows:
- the on-site unified command forms late or not at all
- first aid is given by survivors of the incident and bystanders
- patients are transported with any available means to the closest health care facility.

These facts that have been noted in multiple disasters and terrorist incidents may bring a health care facility to an emergency situation and the front line health professionals in the position of having to care for a large number of casualties in a short period of time.

Triage is the procedure applied to resolve the discrepancy between the number of casualties and the capacity for care by applying a simple rule in order to “do good to the largest possible number of persons”. The aim is to prioritize according to severity of injuries and the availability of medical care, in order to assist in the management of more than one casualty (e.g. priority for evacuation, for transportation to health care facilities, for surgical or other specific treatment).

Triage is performed both on-site of a mass casualty incident but also at the receiving health care facilities. Triage priorities should be re-evaluated frequently especially in mass casualty situations, as patients tend to change priority category.

Special considerations during triage decisions should be: children and elderly, as well as patients exposed to NBC-agents, as decontamination factors in the decision.
In mass casualty situations simple solutions such as relevant colour bracelets or simple coloured tape can be used to expedite the procedure, as well as help indicate the decontamination status of the patient (i.e. one colour for triage priority and one for decon status).

START (Simple Triage And Rapid Treatment) triage system is one of the many available triage systems used in mass casualty incidents aiming to ensure care for as many casualties possible according to the available resources.

START triage is a simple algorithm based on the ABCs of Basic Life Support for the decision making. The priority categories for casualties are:

- **Immediate**: Describing casualties that need immediate attention for life-saving procedures (e.g., intubation, pneumothorax management etc.)
- **Urgent**: Describing casualties that need urgent attention to prevent loss of limb or life (e.g., head injuries, limb fractures, vascular or ocular injury)
- **Minimal**: Describing casualties with simple injuries or psychological effects only (e.g., cuts and abrasions)
- **Expectant**: Describing casualties that are dead or dying, cannot be cared for in the current situation and with the available resources.
Fig VI.1: START Triage algorithm

The assignment of a patient in Green or Yellow priority category will then depend on other concomitant injuries, e.g. limb fractures.
Terrorism has been defined as “the illegal use or threatened use of force or violence; an intent to coerce societies or governments by inducing fear in their populations”. The key word in this definition is the word “fear”, which reveals that apart from causing mass casualties, terrorism aims at causing widespread confusion, fear and psychological stress.

These indirect effects have lasting consequences on people’s mental health and may affect the social and economic life of a society for months or years following a “traumatic” event, act of terrorism or not. The effects of any traumatic incident affect not only the immediate casualties, but also their families and colleagues, the first responders and health professionals who care for them.

On the other hand, all traumatic incidents (natural or man-made) that cause fear can be accompanied by physical symptoms of anxiety that may mimic exposure to an agent and require differentiating by the treating physicians.

In the following paragraphs, we aim to outline the key points with regards to:

• the characteristics of the vulnerable groups, which experience psychological trauma depending on whether they are directly or indirectly affected by the event
• the symptoms of Post Traumatic Stress Disorder (PTSD) that may follow a terrorist act or other traumatic incident and
• ways to prevent and cope with psychological effects and avoid massive panic.

VII  PSYCHOLOGICAL EFFECTS OF TERRORISM

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• ways to prevent and cope with psychological effects and avoid massive panic.
Vulnerable groups:

- Individuals affected by and/or involved in the incident
- Families of individuals affected by and/or involved in the incident
- Children
- First responders: usually referring to police, fire-fighters and emergency medical personnel
- Health care professionals (front line health professionals, usually in the Emergency Departments (ED))
- Persons with pre-existing illnesses or psychological disorders

Remember that psychological effects on an individual depend on their:

- Personality
- Age and personal experiences in life
- Cultural and religious or philosophical background
- Educational background
- Physical and mental health status (history of trauma, unresolved anxieties, and pre-existing chronic illnesses)
- Social support
- Whether affected by and/or involved
- Size and personal distance from the event
- Physical injuries
- Personal losses (Relatives, friends, job loss, material loss, loss of trust and/or faith to civil structures, traumatic experiences)
Symptoms after a traumatic incident:

**Emotions**
- Feeling nothing
- Sadness, grief, fear, anger, irritability, volatile emotions
- Anxiety disorders
- Avoidance of thoughts, places, actions and people

**Cognition**
- Difficulty in concentrating
- Poor or loss of memory
- Disorientation
- Loss of or reduced interest
- Re-experiences: pictures, sounds, scents, motion experiences

**Physical Symptoms**
- Tiredness, exhaustion
- Dizziness, nausea
- Various aches and pains, mainly headaches
- Shaking or tremor
- Breathing difficulties
- Palpitations or other heart problems
- Problems with substance use
- Refuse self-care: not eating, not washing/bathing, not changing clothes
- Suicidal or homicidal thoughts, feelings or plans

**Other**

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Wide Public Health Impact
Preventing / coping with psychological effects and PTSD:

A. Education and Preparedness

- Train mental health professionals in your hospital, especially for assisting after acute traumatic incidents
- Involve your mental health professionals in the preparedness phase of an incident
- Instruct, inform, train and sensitize all possible crisis helpers (even informal care providers in your hospital) about the specific needs of acute traumatized people, as well as how to handle vulnerable groups such as children
- Assist and participate in relevant training activities for the public in your community to respond to a disaster

B. During the Response to an acute incident

- Involve your mental health professionals right from the first hours of an incident
- Induce proper crisis communication at your level in order to avoid mass panic reactions

References

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VIII ADDITIONAL SOURCES OF INFORMATION

A large number of references are included in the CD-ROM that accompanies this handbook, but a short selection of trusted sources for biological, chemical and radiological agents, as well as other health threats is referenced here for more convenience.

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