

CLEAPSS STUDENT SAFETS

2nd edition 2019

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EAPSS

Student safety sheets

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Microorganisms

See also CLEAPSS Student Safety Sheet 76, Bioreactors and Fermenters

Source	Hazard	Comment
Samples from the environment	BIOHAZARD	Air, water and soil samples could be used, but do not samples from high-risk areas, eg, toilets or the floors of changing rooms. All environmental samples could be contaminated with pathogens (organisms which cause disease).
Samples from humans	BIOHAZARD	'Finger dabs' could be used, samples could, however, be contaminated with pathogens (see above).
Foods	BIOHAZARD	Any uncooked animal product (eggs, meat, cheese etc) may be contaminated with bacteria, especially <i>Salmonella</i> and <i>Escherichia coli</i> (<i>E. coli</i>) from the gut, which can cause food poisoning. Take care to prevent cross contamination between cooked and uncooked foods. Thorough cooking will destroy bacteria.
Purchased cultures (ie, samples of microorganisms bought from suppliers)	BIOHAZARD	Cultures bought from reputable suppliers (but not those from hospitals, etc) should be safe but may have become contaminated. <i>E. coli</i> is often studied in schools, but this is not the same strain of bacterium that causes food poisoning.

Typical control measures to reduce risk

- When culturing, if possible, grow bacteria and fungi on solids (agar) rather than liquids (broth) to avoid spills and aerosol formation. Choose culture media, eg, nutrient agar, that do not favour the growth of pathogens. Incubate at temperatures which do not encourage growth of pathogens (avoid temperatures 30°C to 42°C)
- Do **not** seal cultures completely *before* incubation (otherwise hazardous anaerobic bacteria are encouraged) but make sure they cannot be opened accidentally.
- Incubated cultures taken from the environment or humans must never be opened.
- Use sterile equipment and aseptic technique(eg, by flaming loops and mouths of bottles, etc).
- Avoid draughts (from open windows and doors) which could contaminate cultures and cause spores from fungi (eg, mould) to spread.
- Work near Bunsen-burner flames so that the updraught helps to prevent contamination of cultures.
- After work is complete, treat surfaces using a suitable disinfectant, for a sufficient length of time.
- Dispose of all cultures (including mould on food e.g. mouldy bread) by sterilisation in an autoclave (pressure cooker).
- Always wash hands after handling cultures and before handling food.
- Wear a clean lab coat or overall to protect cultures and food from microbes on the skin, clothing, etc.
- In cooking, ensure that food is heated to at least 70 °C for at least 2 minutes.
- Do not reheat cooked food; prepare, store and display cooked and uncooked foods separately.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong? eg, could a food or a culture be, or become, contaminated? Could microorganisms or their spores escape?
- How serious would it be if something did go wrong? eg, could material pathogenic to humans be released? Could food poisoning result?
- How can the risk(s) be controlled for this activity? eg, can it be done safely? Does the procedure need to be altered?

Emergency action

Spilt on the floor, For spills of cultures, place paper towels over the spill, pour disinfectant (eg, Virkon) on top and leave for at least 15 minutes. Bleach is usually suitable in the home. bench, etc

Enzymes

includes amylase, catalase, cellulase, diastase, lipase, proteases (eg, pepsin, trypsin), urease

Substance	Hazard	Comment
Enzymes Powders 'Biological' detergents contain enzymes.	CORROSIVE HEALTH IRRITANT HAZARD	DANGER: Most enzymes are sensitisers (see below) and may cause allergy or asthma symptoms, or breathing difficulties if inhaled. DANGER: Some enzymes can cause serious eye damage. WARNING: Many enzymes irritate the eyes, skin and
Enzymes Concentrated solutions	CORROSIVE HEALTH IRRITANT HAZARD	respiratory system. DANGER: Most enzymes are sensitisers (see below) and may cause allergy or asthma symptoms, or breathing difficulties if inhaled. DANGER: Some enzymes can cause serious eye damage. WARNING: Many enzymes irritate the eyes, skin and respiratory system.
Enzymes Dilute solutions (less than 1% w/w) or 5% Lipase	LOW HAZARD	Most at these concentrations are unlikely to offer any significant risk. Biological systems are rich sources of enzymes, eg, liver (catalase), saliva (amylase).

Note: Some people are *allergic* to particular substances; their bodies' immune system reacts to these substances to an unusual extent. Asthma is one type of allergy which results in breathing difficulties. A sensitiser is a substance that may produce only a small or even no allergic reaction when humans are first exposed to it (sometimes over an extended period of time) but can produce a much more severe reaction on subsequent occasions, even when the body is exposed to much smaller amounts.

Typical control measures to reduce risk

- Use the lowest concentration/smallest amount possible.
- Wear eye protection for enzyme at 1% or higher concentration
- Avoid powdered enzymes escaping into the air; use a fume cupboard (not switched on) when handling enzyme powders.
- Do not spray enzyme solutions.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong? eg, could enzyme dust be breathed in?
- How serious would it be if something did go wrong?
- How can the risk(s) be controlled for this activity? eg, can it be done safely? Does the procedure need to be altered?

•	In the eye	Flood the eye with gently-running tap water for 10 minutes. Consult a medic if powder involved.
•	Swallowed	Wash out the mouth with drinking water. Do not induce vomiting. Consult a medic.
•	Dust breathed in	Remove the casualty to fresh air. Consult a medic if breathing is difficult
•	Spilt on the skin or clothing	Remove contaminated clothing. Wash off the skin with soap and plenty of water. Rinse contaminated clothing.
•	Spilt on the floor, bench, etc	Scoop up powders (take care not to raise dust). Wipe up solution spills or any traces of powders with a damp cloth.



Human body fluids and tissues

includes cheek cells, blood, saliva, sweat & urine

Source	Hazard	Comment
Cheek cells	BIOHAZARD	There is a very tiny risk of transmission of HIV or hepatitis virus but only if contact is made with samples other than your own. Sampling may be banned in some schools although this is now very unlikely.
Blood	BIOHAZARD	There is some risk of transmission of HIV or hepatitis virus if contact is made with blood other than your own. Taking blood samples is possible if stringent precautions are taken but may not be permitted in a few schools. Never share hypodermic needles or become 'blood brothers'.
Saliva	LOW HAZARD	There is negligible risk of transmission of HIV or hepatitis virus even if you come into contact with saliva other than your own. Kissing is rarely banned for reasons of hygiene!
Sweat	LOW HAZARD	There is negligible risk of transmission of diseases even if you come into contact with sweat other than your own but this is no excuse for poor hygiene!
Urine	BIOHAZARD	There is a very tiny risk of transmission of various diseases if you come into contact with urine other than your own, although urine is normally sterile. In investigations involving urine, take care when obtaining and transporting samples. Wash hands after using the toilet.

Typical control measures to reduce risk

- Only handle samples from your own body.
- After use, hygienically dispose of samples, disinfect contaminated containers by immersion for 30 minutes in a solution of 1% Virkon or sodium chlorate(I) (hypochlorite) for 10 minutes; treat benches for a sufficient length of time with a suitable disinfectant (Virkon is preferred) and wash hands. Any swabs, slides or other equipment contaminated with blood should be collected in a suitable container, then autoclaved. If necessary, use a 'sharps' container (eg, a sturdy box, clearly labelled and sealed and wrapped before disposal).
- Treat clinical thermometers, mouthpieces, etc in Milton for 30 minutes before and after use (unless disposable).
- In first aid, minimise contact with blood by wearing disposable surgical gloves or by asking the casualty to carry out her/his own treatment, eg, by applying pressure to a wound using a pad of cloth.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong?

eg, could somebody else come into contact with samples from your body, or vice versa? If first-aid treatment was **not** applied, could the casualty's condition put his or her life at risk?

- How serious would it be if something did go wrong? eg, could HIV or hepatitis virus or other pathogens be transmitted?
- How can the risk(s) be controlled for this activity? eg, can it be done safely? Does the procedure need to be altered?

•	Spilt on the floor, bench, etc	For spills of most body fluid samples, place paper towels over the spill, pour disinfectant (eg, 1% <i>Virkon</i>) on top and leave for at least 15 minutes. For blood, wear disposable gloves, wet paper towels or cloth with freshly-prepared sodium chlorate(I) (hypochlorite, bleach) containing at least 10 000 ppm chlorine or 1% <i>Virkon</i> . Add more disinfectant to soak area and leave for 15-30 minutes. Still wearing
•	Disposal of bloody tissues, etc	gloves rinse towels or cloth under running water Wash hands thoroughly with soap and water. Avoid skin contact with tissues etc contaminated with blood. In school, place in a container for sanitary towels, etc. Then incinerate or use a clinical waste-collection service. At home, flush down the toilet or wrap carefully and place in the refuse.

Food testing (1)

See also CLEAPSS Student Safety Sheet 5, Food Testing (2)

Substance	Hazard	Comment
Food		Uncooked samples of food may be contaminated with microbes. (See CLEAPSS Student Safety Sheet 1, Microorganisms.) Some people are allergic to some foods, especially peanuts.
Fehling's solution Used to test for reducing sugars	CORROSIVE	See CLEAPSS Student Safety Sheets 31 and 40. Solution A contains slightly-acidic copper sulfate solution. Solution B contains 3M sodium hydroxide solution. Solution A DANGER: Causes serious eye damage and skin irritation. Solution B DANGER: Causes severe eye damage and skin burns. As the mixture has to be heated in a test tube, there is a risk of alkali spitting out.
Benedict's solution Used to test for reducing sugars	IRRITANT	See CLEAPSS Student Safety Sheet 40. Contains slightly-alkaline 0.07M copper sulfate solution. WARNING: Irritant to skin and eyes. Some risk of spitting when heating test tubes.
Ethanol Used to test for fats (lipids)	HIGHLY HARMFUL HEALTH FLAMMABLE HAZARD	Most schools in fact use IDA (industrial denatured alcohol) rather than pure ethanol. See CLEAPSS <i>Student Safety Sheet 60</i> . DANGER: highly flammable, harmful by ingestion, may cause damage to organs. If Bunsen burners are being used nearby for other food tests, there is a serious fire risk.
lodine solution Used to test for starch	LOW HAZARD	See CLEAPSS Student Safety Sheet 56. The solution is so dilute (about 0.01M) that it presents only a LOW HAZARD.
Biuret test Used to test for proteins	CORROSIVE	See CLEAPSS Student Safety Sheets 31 and 40. DANGER/WARNING: The very dilute copper sulfate solution is LOW HAZARD. Sodium hydroxide solution may be CORROSIVE, IRRITANT or LOW HAZARD depending on concentration.

Typical control measures to reduce risk

- Do not taste foods in laboratories; avoid using products containing peanuts etc if there is a known allergy.
- Wear eye protection and use the smallest possible amounts of chemicals.
- Use Benedict's solution rather than Fehling's solution and heat with a water bath.
- Do not use ethanol if there are naked flames nearby.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong? eg, could chemicals spit out of a heated test tube, or might somebody using a Bunsen be unaware of ethanol being used nearby?.
- How can the risk(s) be controlled for this activity? eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

Emergency action

•	in the eye	Flood the eye with gently-running tap water for 10 minutes. Consult a medic.
•	Swallowed	Do no more than wash out the mouth with drinking water. Do not induce vomiting. Consult a medic.
		Description of the state of the

Spilt on skin or Remove contaminated clothing. Then drench the skin with plenty of water. If a large area is affected or clothing blistering occurs, consult a medic.

Push casualty to the floor, roll the body or smother flames on clothing or skin with fire blanket or other Clothing material. Cool burnt skin with gently running tap water for 10 minutes. Unless trivial consult medic. catches fire Other ethanol Allow fires in sinks, etc to burn out. Fires at the top of test tubes, beakers etc can be smothered with a damp cloth or heat-resistant mat if this can be done safely. fires

For small amounts, use a damp cloth. Rinse well. For larger amounts, cover with mineral absorbent (eg, Spilt on floor, cat litter) and scoop into a bucket. Neutralise alkali with citric acid. Rinse with water. bench, etc



Food testing (2)

See also CLEAPSS Student Safety Sheet 4, Food Testing (1)

Substance	Hazard	Comment
Millon's reagent Used to test for proteins and Cole's modification (Millon's reagent A)	TOXIC HEALTH HAZARD CORROSIVE ENVIRON.	DANGER: It contains a high concentration of a mercury compound and concentrated nitric acid (see CLEAPSS <i>Student Safety Sheets 21</i> and <i>44</i>). <i>Cole's modification</i> is less hazardous because it contains less concentrated sulfuric acid and also uses sodium nitrate(III) (nitrite). DANGER: Fatal if swallowed or on skin contact; causes severe skin burns and eye damage; suspected of causes genetic defects and damaging fertility. Because of toxicity, all residues must be collected for licensed disposal. There is a risk of spitting when it is heated in a test tube.
Sakaguchi test Used to test for proteins	CORROSIVE HIGHLY FLAMMABLE ENVIRONMENT	The test involves mixing three solutions: sodium hydroxide (~1.3 M), napthalen-1-ol in ethanol (~0.07 M) and sodium chlorate(I) (hypochlorite) (~1.5 M). See relevant CLEAPSS <i>Student Safety Sheets</i> . Despite these hazards, it is safer to use than either form of Millon's reagent because it does not need to be heated and only a few drops are required. The biuret test is safer still (see CLEAPSS <i>Student Safety Sheet 4</i>). DANGER: corrosive to skin and eyes, highly flammable
DCPIP Used to test for Vitamin C	Low hazard	See CLEAPSS Student Safety Sheet 70, Dyes and indicators. (Also known as PIDCP.)
Saliva Used to break down starch	Low hazard	See CLEAPSS Student Safety Sheet 3, Human body fluids and tissues. Negligible risk.
Clinistix Used to test for glucose	Low hazard	The tip of the stick contains a minute amount of a known carcinogen and should not be touched. The sticks should be stored and disposed of safely. It is normally used for testing urine.
Albustix Used to test for proteins	Low hazard	This will not detect all proteins. It is normally used for testing urine.

Typical control measures to reduce risk

- Wear eye protection and use the smallest possible amounts of chemicals.
- Use the least-hazardous substance that achieves the required effect.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong? eg, could chemicals spit out of a heated test tube?
- How serious would it be if something did go wrong? eg, could ethanol (in the Sakaguchi test) catch fire, or acid splash into the eye?
- How can the risk(s) be controlled for this activity? eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

•	In the eye	Flood the eye with gently-running tap water for 10 minutes. Consult a medic.
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- Do no more than wash out the mouth with drinking water. Do not induce vomiting. Consult a medic. **Swallowed**
- Spilt on skin or Remove contaminated clothing. Drench the skin with plenty of water. If a large area is affected or clothing blistering occurs, consult a medic.
- Push casualty to the floor, roll the body or smother flames on clothing or skin with a fire blanket or Clothing catches fire other material. Cool burnt skin with gently running tap water for 10 minutes. Unless trivial Consult a
- Other ethanol Allow fires in sinks, etc to burn out. Fires at the top of test tubes, beakers etc can be smothered with a fires damp cloth or heat-resistant mat if this can be done safely.
- Spilt on floor, For small amounts, use a damp cloth. Rinse well. For larger amounts, cover with mineral absorbent (eg, cat litter) and scoop into a bucket. Neutralise acid with sodium carbonate. Rinse with water. bench, etc



Humans as the subject of investigation (1)

including simple measurements on the body and senses and psychological effects See also CLEAPSS Student Safety Sheets 3, 7, 8 and 9

Hazard Source Comment

All investigations on pupils



Pupils should never be pressurised to take part in investigations on themselves. Individual pupils may become alarmed or distressed, or be subject to unpleasant comments from others in the class, if the results for their bodies are significantly different from others. In fact, a wide spread of figures may be regarded as normal and, in any case, measurement methods used in schools may not be very accurate or reliable.

Biohazards may result if body fluids are used (see Sheet 3).

Investigations of body mass and size, sight, colour blindness, teeth, hearing, reaction time, touch sensitivity, ear lobe attachment, eye colour, etc



Hazards include excessively loud sounds, bright light sources, sharp objects (touch testing), falling objects (reaction timing). Hygiene issues may arise with shared dental mirrors, etc.

Individual pupils may become alarmed or distressed, or be subject to unpleasant comments from others in the class, if genetic tests raise questions about biological or adoptive parentage, although in fact blue-eyed children can occasionally arise from biological parents with brown eyes because the inheritance of eye colour is a complex issue.

Typical control measures to reduce risk

- Ensure only willing volunteers are used and re-assure them if results are exceptional or surprising.
- Be aware of any medical conditions that could affect pupils' ability to participate and/or the results.
- Use hair grips rather than pins for touch sensitivity tests.
- Avoid sounds more than 85 dB(A).
- Ensure dental mirrors etc are properly disinfected between each use (see Student Safety Sheet 9).
- Use non-LED torches as light sources.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong?

eg, could somebody be injured by the equipment?

- How serious would it be if something did go wrong?
- How can the risk(s) be controlled for this activity?

eg, can it be done safely? Does the procedure need to be altered?

Emergency action

Minor cuts

Wash the wound. Get the casualty to apply a small, sterilised dressing.

Exposure to very loud sounds

Brief exposure to sounds of the loudness likely to be achieved in a school laboratory is unlikely to cause lasting damage. Reassure the casualty. Repeated exposure to loud music on portable media players or at festivals is much more damaging,

Exposure to very bright lights

Although looking directly at the Sun and other very bright lights can cause permanent damage, the natural aversion response of the eyes of pupils will usually prevent this unless deliberately overridden. Reassure the casualty.

Impact injury

Rest the injured part; apply ice to reduce the bruising and pain; get attention from a first-aider.

Humans as the subject of investigation (2)

including exercise, breathing and blood pressure

See also CLEAPSS Student Safety Sheets 3, 6, 8 and 9

Source	Hazard	Comment
Investigating effects of exercise	DANGER	Over-exertion may be a hazard, especially for those with certain medical conditions. Competitive situations can lead to careless behaviour and accidents. Unsuitable footwear, uneven surfaces, running up and down stairs and unstable equipment may be hazards.
Investigating breathing	DANGER	Shared mouthpieces are sources of infection. When using manometers, fluid may be taken into the mouth. Use of spirometers which have a large chamber filled with air or oxygen must be closely supervised by the teacher. Use of lung-volume bags (or even water-filled bell jars), data-logging sensors and peak-flow meters are much safer. It is dangerous to carry out investigations involving rebreathed air for more than 1 minute. Lung volume bags need to be sterilised and dried between use.
Investigating blood pressure	DANGER	Using a sphygmomanometer with a mercury manometer and a stethoscope requires great skill. Electronic models, especially with automatic cuff inflation, are much more suitable but still require close teacher supervision.

Typical control measures to reduce risk

- Ensure only willing volunteers are used and reassure them if results are exceptional or surprising.
- Make sure you know how to use any equipment safely.
- When taking exercise, use step-ups on stable equipment rather than running up stairs, do not exercise more than in PE and be aware of students with asthma, diabetes, circulatory problems or those advised not to take part in normal PE lessons.
- Change and disinfect mouthpieces after each pupil.
- Ensure manometers have a trap to prevent fluid being taken into the mouth.
- Only use equipment for measuring blood pressure under the direct supervision of trained staff.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong?

eq, could somebody over-exert themselves or trip dangerously?

If first-aid treatment were not applied, could the casualty's condition put his or her life at risk?

How serious would it be if something did go wrong?

eg, could it become life-threatening?

How can the risk(s) be controlled for this activity?

eg, can it be done safely? Does the procedure need to be altered?

Emergency action

difficulties

Rest the injured part; apply ice to reduce the bruising and pain; get attention from a first-Impact injury

Students should use their own inhalers (if any) and inform teacher Asthma or breathing



Humans as the subject of investigation (3)

including tasting, eating & drinking and chemicals on the skin

See also CLEAPSS Student Safety Sheets 3, 6 and 7

Source

Hazard

Comment

Investigations where chemicals are placed on skin



HAZARD

IRRITANT

Hazardous chemicals may be placed on the skin during evaporation tests or when investigating perspiration. Where possible, it is safest to use substances which are deliberately intended for skin use, e.g. alcohol based gels.

FNVIRON.

HAZARD



FLAMMABLE

DANGER: Cobalt chloride (or thiocyanate) may cause allergic skin reaction or asthma-like symptoms if inhaled. Suspected of causing genetic defects or cancer if inhaled. May damage fertility. Toxic to aquatic life. These risks are negligible when impregnated on paper but skin contact must be minimised, although there is no evidence of problems from typical school use.

Investigations involving tasting, eating and drinking



Eating and drinking should not take place in laboratories and in some circumstances it may be illegal under the COSHH Regulations.

Use of disclosing tablets in accordance with the manufacturer's instructions is considered low hazard. Tasting small amounts of dilute solutions (eg, sugar, salt) must be done under hygienic conditions, using a drinking straw, cotton bud or impregnated filter paper.

DANGER: PTC (phenyl thiocarbamide, phenylthiourea, PTU), used in genetics testing, is fatal if swallowed and may cause an allergic skin reaction. It must be used only in carefully controlled, very small, amounts e.g. on brought in test strips

Consuming alcohol in any form or caffeine in the form of *ProPlus* tablets is not acceptable. Use caffeine in cola drinks or coffee instead but limit total caffeine intake by avoiding it before and after testing. Limited amounts of Red Bull might be used if students drink it anyway, although it contains other components as well.

Typical control measures to reduce risk

- Ensure only willing volunteers are used and reassure them if results are exceptional or surprising.
- Only use water or alcohol based hand gels for testing evaporation on skin.
- If using cobalt chloride (or thiocyanate) paper on the skin, handle with forceps and wash skin afterwards.
- Do not conduct taste tests in laboratories; use dining areas or food technology rooms.
- Observe scrupulous hygiene no sharing of cups, spoons, straws, etc.
- Use only specially-purchased food- or pharmaceutical-quality chemicals for tasting.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong? eg, could somebody taste the wrong substance by accident or as a result of mischief?
- How serious would it be if something did go wrong? eg, could it become life-threatening?
- How can the risk(s) be controlled for this activity? eg, can it be done safely? Does the procedure need to be altered?

Emergency action

Hazardous chemicals in mouth Wash out mouth

Hazardous chemicals swallowed Wash out mouth. Do NOT make victim vomit. Consult a medic.

Disinfectants

Source	Hazard	Comment
Disinfectants containing active chlorine, eg Domestos, Milton	varies	Effective against wide range of microorganisms. Less suitable for dirty conditions because disinfectant is degraded by organic matter. They corrode metals and generate toxic chlorine with acids. A good choice where blood is involved but use is discouraged by some employers. Requires a contact time of at least 15 minutes. See CLEAPSS Student Safety Sheet 41, Sodium chlorate(I) (sodium hypochlorite).
Disinfectants containing methanal (formaldehyde)	varies	Very effective at killing microorganisms but more concentrated solutions are hazardous. See CLEAPSS Student Safety Sheet 67, Methanal.
Ethanol, 70%, industrial denatured alcohol (IDA), 70%	HIGHLY IRRITANT HEALTH FLAMM. HAZARD	Active against most bacteria, viruses and fungi, within 5 minutes. Mainly useful for swabbing benches immediately before preparing cultures or for rapid disinfection of mouthpieces, dental mirrors, etc – rinse with clean water after sterilisation. DANGER: highly flammable, harmful by ingestion, may cause damage to organs See CLEAPSS Student Safety Sheet 60, Ethanol.
VirKon , powder	CORROSIVE	Main active component is dipotassium peroxomonosulfate, which oxidises the infective material. Powder can be poured directly onto liquid spills, but is usually dissolved in water. DANGER: causes severe skin irritation and serious eye damage.
VirKon, 1% aqueous solution	LOW HAZARD	Very effective with 10 minutes contact against bacteria, fungi and viruses. Solution attacks metal after 10 minutes. The best choice for most situations.
Phenolic disinfectants	varies	Clear phenolics have been phased out under the <i>Biocidal Products Regulations</i> . Other phenolics (white or black, eg <i>Jeyes Fluid, Izal, Lysol</i>) are less effective and some are TOXIC/CORROSIVE. Not generally recommended for laboratory use.

Typical control measures to reduce risk

- Where possible, avoid the use of disinfectants classed as TOXIC or CORROSIVE and use freshly-made solutions.
- Wear eye protection, including when making or disposing of solutions.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong? eg, Is the disinfectant effective against the organisms likely to be present and will it be left long enough to be effective?
- How serious would it be if something did go wrong? eg, could a failure of disinfection lead to a serious illness?
- How can the risk(s) be controlled for this activity?

eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

Emergency action

See CLEAPSS Student Safety Sheet 41 if sodium chlorate(I) used. Sodium chlorate(I) incidents

Immediately rinse the eye with gently-running tap water for at least 10 minutes. Hazardous chemicals in the

Consult a medic. eve

Hazardous chemicals Do no more than wash out the mouth with water. Do not induce vomiting. Consult a swallowed medic.

Brush off solids. Rinse skin or clothing thoroughly with water. Spilt on the skin or clothing

Spilt on the floor, bench, etc Wear eye protection. Scoop up any solid and wipe up small amounts with a damp

cloth and rinse it well. Cover solutions with mineral absorbent (eg, cat litter) and

scoop into a bucket. Rinse well.



Electricity

Voltage / current	Hazard	Comment
230 V ac and above at high currents (over 5 mA).	ELECTRIC SHOCK /	In non-school contexts: Electric power distribution, eg over-head power lines and local sub-stations could cause accidents if children behave foolishly.
Above 30V AC or DC and at currents over 5 mA. This includes the 230 V ac mains supply.	ELECTRIC SHOCK / BURN	In school science: problems may arise from terminals of high voltage (high tension, HT) supplies or low-voltage units with an HT outlet (as some supply 150 mA); also in activities involving electrophoresis, model transformers or conductivity of molten glass. In non-school contexts: problems arise due to poor insulation (damaged wiring and plugs), incorrect wiring, over-loaded circuits, poor earthing or vandalism.
Less than 30 V AC or DC and at currents over 5 mA. This includes almost all work with batteries in school or elsewhere.	ELECTRICAL HAZARD, LOW TOXIC CORROSIVE	In school science: most school circuit work, including electrolysis, is in this category (although problems could arise if currents over 10 A were used). Some cells, batteries and accumulators contain TOXIC or CORROSIVE materials.
Any voltages at very low currents (well below 5 mA) Eg, Static electricity	LOW HAZARD	In school science: examples include the Van de Graaff generator (but not induction coils which may give over 5 mA). Electronic equipment nearby may be damaged by static discharges or electromagnetic fields.

Typical control measures to reduce risk

- Use the lowest voltage possible (and, for electrolysis, the lowest current and concentration that gives good results).
- Avoid exposed conductors which are live above 30 V AC or DC.
- Avoid the possibility of water coming into contact with conductors which are live above 30 V AC or DC.
- Check that primary and secondary insulation (ie, both layers of plastic coating) are in good condition.
- Avoid over-loaded circuits, too many plugs in one socket, etc.
- Check that plugs are correctly wired with appropriate fuses.
- Ensure good earth connections where necessary.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong? eg, accidentally touching a live component through poor design or poor maintenance.
- How serious would it be if something did go wrong? eg, could a current flow through the heart? How large a voltage and/or current?
- How can the risk(s) be controlled for this activity? eg, can it be done safely? Does the procedure need to be altered?

Emergency action

Electric shock

Take care for your own safety.

Break contact by switching off or removing the plug. If this is not possible, use a wooden broom handle or wear rubber gloves to pull the casualty clear. Consult a medic.

If the casualty is unconscious, check that airways are clear and that the casualty is breathing and has a pulse. If so, place the casualty in the 'recovery position'. If a pulse is found but the casualty is not breathing, artificial ventilation is necessary. If no pulse is found and the casualty is not breathing, cardio-pulmonary resuscitation is necessary.

Radioactive materials

Substance	Hazard	Comment
Alpha sources (α radiation) Low-activity α sources are used in domestic smoke alarms.	IONISING RADIATION	Produce heavy ionisation (see <i>Additional Information</i> , below) but range is less than 1 mm in living tissue. Little problem if source is kept outside body.
Beta sources (β radiation) All samples of potassium and its compounds (including granite rocks, clays, etc) contain very small amounts of a natural β emitter.	IONISING RADIATION	Produce medium ionisation (see Additional Information, below) but the range is a few centimetres in living tissue. There is little problem if source is kept away from the body.
Gamma sources (γ radiation) Very active γ sources are used in hospitals for killing cancer cells. Domestic smoke alarms emit a little γ radiation.	IONISING RADIATION	Produce little ionisation (see Additional Information, below) but the range is long in living tissue, some passing right through the body. In schools, use a weak source and keep well away from the body.

Additional information:

- When (ionising) radiation is absorbed by living tissue, mostly it results in a few extra hydrogen ions and hydroxide ions in the cytoplasm of cells. These rapidly recombine to form water.
- Some radiation is absorbed by more complex molecules and the ions from these can result in the death of the cell.
- Low levels of ionising radiation have little noticeable effect because biological organisms are continually replacing cells which die for other reasons anyway.
- High doses of radiation can result in skin burns (like sunburn) or radiation sickness (where so many cells have been killed that an organ ceases to function properly).
- Ionising radiation can also affect DNA in cells and change the genetic code. In reproductive organs, this could cause abnormal offspring but has never been confirmed in humans.
- Modified DNA may allow cells to reproduce out of control and form a cancer.

Typical control measures to reduce risk

- Use the lowest-activity source possible (only low-level sources are permitted in schools).
- Keep as far away from the source as possible (For school sources, use a handling tool which keeps the source at least 10 cm from the hand. Observers of demonstrations should generally keep at least 2 m away.).
- Have a clear set of local rules, including "Sources must not be handled by under-16s".
- When radioactive solids, liquids or gases (open sources) are in use, prevent contamination of people (by use of lab coats and disposable plastic gloves), of benches (by use of trays and sheets of absorbing paper) and apparatus (by handling equipment with disposable tissues).

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong?
 - eg, source is dropped, spilt or stolen or somebody moves too close to the source.
- How serious would it be if something did go wrong?
- How can the risk(s) be controlled for this activity? eg, can it be done safely? Does the procedure need to be altered?

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•	Sealed source dropped	Do not look directly at source, but use mirror to examine source for damage. Check area
		for radioactivity where source was dropped.
•	Spilt on the skin or	Wash the affected area thoroughly and check for radioactivity. If swallowed, go to a
	swallowed	hospital specialising in radiation incidents.
•	Open source spilt on	Wipe up small amounts with damp tissues. Wipe the area until count rate is less than 50%
	the floor, bench, etc	above background. Place tissues in a plastic bag and dispose of it in solid waste.

Electromagnetic radiation

Type of radiation	n Hazard	Comment		
Radio waves LOW HAZARD		There is no evidence that electric and magnetic fields at ordinary		
Wavelength 10 ³ m		intensities affect the human body.		
Microwaves	\wedge	These are strong sources which produce local heating. Microwave		
Wavelength 10 ⁻² m	((6,2))	ovens should be operated and maintained according to instructions,		
		so that microwaves cannot leak out. Using mobile phones is safe if		
	NON-IONISING RADIATION	the power is less than 100 W m ⁻² . Microwave radiation has been used		
		medically to give relief from pain.		
Infra-red		It can produce skin burns and strong sources (eg, furnaces for melting		
Wavelength 10 ⁻⁵ m	((a))	iron) may damage the cornea of the eye. It may ignite combustible		
Traverengen zo	NON-IONISING RADIATION	materials. Infra-red radiation has been used medically to give relief		
		from pain. TV remote control units are safe.		
Visible	\wedge	Intense sources (eg, from the Sun and narrow beams from class 3 & 4		
Wavelength	((₁))	lasers) may damage the retina. Class 1(but not 1M) lasers (eg, in laser		
0.5 x 10 ⁻⁶ m	NON-IONISING LASER	printers) are totally enclosed. Class 2 (but not 2M) lasers are low		
0.5 X 10 III	RADIATION RADIATION	power (less than 1 mW) and safe unless the 'blink response' is over-		
	RADIATION RADIATION	ridden or ignored. Some cheap laser pointers are wrongly classified		
		and may be class 3B (typical power 3 mW). Some are even 100 mW.		
		These are dangerous, especially if misused. Direct observation of		
		eclipses of the Sun often results in blindness.		
Ultra violet		It can cause sunburn. There is a risk of skin cancer. Do not sunbathe		
Wavelength 10 ⁻⁸ m	((a))	without adequate protection. Only class 1 UV lasers are safe.		
UV-A, 315-400 x 10 ⁻⁹ ;	NON-IONISING RADIATION	UV-A is used in sun beds. UV-B and UV-C can damage the eyes.		
UV-B, 218-315 x 10 ⁻⁹ ;		Although UV-A is safer, many sources produce all wavelengths. UV is		
		produced in electric-arc or oxy-acetylene welding. Glass which is		
UV-C, 100-280 x 10 ⁻⁹ .		4 mm thick stops UV-B and UV-C. Wear snow goggles when skiing.		
X-rays		X-rays are produced in low-pressure systems by sparks and discharges		
Wavelength 10 ⁻¹⁰ m		at more than 6 kV. Large doses cause burns and may induce cancer.		
	IONISING RADIATION	X-rays are used in medicine for diagnosis and treatment (but should		
	A TOMORIO RADIATION	be avoided during pregnancy).		
Gamma (γ) rays		See CLEAPSS Student Safety Sheet 11, Radioactive materials.		
Wavelength 10 ⁻¹² m				
	IONISING RADIATION			

Typical control measures to reduce risk

- X-ray sets are permitted in schools only where there are suitably-qualified staff.
- Ultra-violet lamps must be screened or personal protective equipment used (also for welding).
- Lasers for use in schools must be class 1 or class 2, when the rule is "Do not stare down the beam".
- Use sun lotions with a suitable protection factor and do not sunbathe without adequate protection.
- Never look directly at the Sun (even during an eclipse) or through a lens, filter or pin hole.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong?
- How serious would it be if something did go wrong?
- How can the risk(s) be controlled for this activity? eg, can it be done safely? Does the procedure need to be altered?

- Radiation burns to the skin Treat as for a heat burn, ie, cool burnt skin under gently-running tap water for 10 minutes. Consult a medic if more than an area the size of a small coin is affected.
- Suspected eye damage Cover and consult a medic.

Hydrochloric acid

also applies to Hydrogen chloride gas

Substance	Hazard	Comment
Hydrogen chloride Gas	CORROSIVE TOXIC	DANGER. It causes severe skin burns and eye damage. It is toxic if breathed in. For a 15-minute exposure, the concentration in the atmosphere should not exceed 8 mg m ⁻³ . Effects of exposure by inhalation may or may not be immediately apparent and can develop and/or increase over time. Inhalation by those with known breathing difficulties (eg asthma) may exacerbate such pre-existing conditions.
Concentrated hydrochloric acid (If 6.8 M or more)	CORROSIVE IRRITANT	DANGER. It causes burns. The vapour irritates the lungs.
Moderately- concentrated hydrochloric acid	(1)	WARNING. It may irritate the eyes, and respiratory system.
(If less than 6.8 M but 2.7 M or more)	IRRITANT	
Dilute hydrochloric	LOW HAZARD	This includes stomach acid.
acid		Dilute acid may still cause harm in the eyes or in a cut.
(If less than 2.7 M)		

Typical control measures to reduce risk

- Use the lowest concentration possible.
- Use the smallest volume possible.
- Wear eye protection for all but the most-dilute solutions; goggles for concentrated acids.
- Wear protective gloves if anything more than tiny amounts of concentrated acid is handled.
- Avoid breathing the gas or fumes from concentrated solutions, eg by use of a fume cupboard.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong?

eg, hazardous products (such as chlorine) produced as a result of a reaction wit the acid or choking fumes if the concentrated acid is over-heated.

How serious would it be if something did go wrong?

NB There are occasional reports of students being taken to hospital as a result of breathing in chlorine.

How can the risk(s) be controlled for this activity?

eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

Emergency action

In the eye Flood the eye with gently-running tap water for 10 minutes. Consult a medic.

Vapour breathed in Remove to fresh air. Consult a medic if breathing difficult.

Swallowed Do no more than wash out the mouth with drinking water. Do not induce vomiting. Consult a

medic.

Spilt on the skin or clothing

Remove contaminated clothing. Then drench the skin with plenty of water. If a large area is affected or blistering occurs, consult a medic.

Spilt on the floor, bench, etc

For release of gas, consider the need to evacuate the laboratory and open all windows. For large spills, and especially for (moderately) concentrated acid, cover with mineral absorbent (eg, cat litter), then scoop into a bucket. Neutralise with sodium carbonate. Rinse with plenty of

water. Wipe up small amounts with a damp cloth and rinse well.

Nitric (v) acid

Substance	Hazard	Comment
Concentrated nitric acid (If 10 M or more)	CORROSIVE OXIDISING	DANGER: It causes severe skin burns and eye damage; skin is stained yellow and then peels. For a 15-minute exposure, the vapour concentration in the atmosphere should not exceed 2.6 mg m ⁻³ . Use a fume cupboard. May cause or intensify a fire in contact with combustible materials. Usually supplied in plastic bottles which will be attacked by the acid after a few years – transfer to borosilicate bottles with chemically-resistant cap.
Moderately-concentrated nitric acid (If less than 10 M but 0.8 M or more)	CORROSIVE	DANGER: It causes severe skin burns and eye damage.
Moderately dilute nitric acid (If less than 0.8 M but 0.1 M or more)	IRRITANT	It is irritating to the eyes and skin.
Dilute nitric acid (If less than 0.1 M)	LOW HAZARD	It may still cause harm in the eyes, in a cut or on the skin.

Typical control measures to reduce risk

- Use the lowest concentration possible.
- Use the smallest volume possible.
- Wear eye protection including when making or disposing of solutions.
- Wear protective gloves if concentrated acid is handled, especially if more than test tube amounts
- Ensure good ventilation if oxides of nitrogen could be formed

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong?

eg, could hazardous products (such as oxides of nitrogen) be formed in reactions with the acid or corrosive fumes produced if concentrated acid is over-heated?

- How serious would it be if something did go wrong?
 - eg, peeling skin, from burns caused by concentrated acid, may be very painful.
- How can the risk(s) be controlled for this activity?

plenty of water.

eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

Emergency action

•	In the eye	Flood the eye with gently-running tap water for 10 minutes. Consult a medic.
•	Fumes breathed in	Remove the casualty to fresh air. Keep him/her warm. Consult a medic even if no symptoms are apparent.
•	Swallowed	Do no more than wash out the mouth with drinking water. Do not induce vomiting. Consult a medic .
•	Spilt on the skin or clothing	Quickly use a dry cloth or paper towel to wipe as much liquid off the skin as possible. Then drench with plenty of water. If a large area is affected or blistering occurs, consult a medic . Remove contaminated clothing and rinse it well.
•	Spilt on the floor,	Wipe up small amounts with a damp cloth and rinse it well.
	bench, etc	For larger amounts, and especially for (moderately) concentrated acid, cover with mineral

absorbent (eg, cat litter) and scoop into a bucket. Neutralise with sodium carbonate. Rinse with

Sulfuric(VI) acid

Substance	Hazard	Comment
Concentrated sulfuric(VI) acid		DANGER. It causes severe skin burns and eye damage.
		It reacts violently, becoming very hot, when mixed with water.
	CORROSIVE	For a 15-minute exposure, the vapour concentration in the atmosphere should not exceed 0.15 mg m ⁻³ .
Moderately-concentrated sulfuric(VI) acid (If 1.5 M or more)		DANGER. It causes severe skin burns and eye damage.
Includes acid used in car batteries.	CORROSIVE	
Moderately-dilute sulfuric(VI) acid	<u>(i)</u>	WARNING. It may irritate the eye and skin.
(If less than 1.5 M but 0.5 M or more)	IRRITANT	
Dilute sulfuric(VI) acid		It may still cause harm in the eyes or in a cut.
(If less than 0.5 M)	LOW HAZARD	For many activities in school science, 0.4 M is adequate.

Typical control measures to reduce risk

- Use the lowest concentration possible.
- Use the smallest volume possible.
- Wear eye protection for all but the most-dilute solutions; goggles for concentrated acids.
- Wear protective gloves if anything more than tiny amounts of concentrated acid is handled.
- Add the concentrated acid slowly to cold water (or preferably ice) when diluting, never the reverse; stir frequently to ensure good mixing.

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong? eg, hazardous products are formed in reactions with the acid or corrosive fumes are produced if concentrated acid is over-heated.
- How serious would it be if something did go wrong? eg, skin and eyes can be seriously burned if not treated quickly.
- How can the risk(s) be controlled for this activity? eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

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Ξ	mergency action	1
•	In the eye	Flood the eye with gently-running tap water for 10 minutes. Consult a medic.
•	Swallowed	Do no more than wash out the mouth with drinking water. Do not induce vomiting. Consult a medic.
•	Spilt on the skin or clothing	Remove contaminated clothing. Especially with concentrated acid, quickly use a <i>dry</i> cloth or paper towel to wipe as much liquid as possible off the skin. Then drench the skin with plenty of water. If a large area is affected or blistering occurs, consult a medic.
 Spilt on 	Spilt on the floor,	Wipe up small amounts with a damp cloth and rinse it well.
	bench, etc	For larger amounts, and especially for (moderately) concentrated acid, cover with mineral absorbent (eg, cat litter) and scoop into a bucket. Neutralise with sodium carbonate. Rinse with plenty of water.

Ethanoic (acetic) acid

also applies to methanoic (formic) acid

Substance	Hazard	Comment			
Ethanoic acid (acetic acid) Pure liquid, 'glacial' Also methanoic (formic) acid usually supplied as 90% solution (about 20 M)	CORROSIVE FLAMMABLE	DANGER: both acids cause severe skin burns and eye damage. Ethanoic acid only: flammable vapour and liquid. For a 15-minute exposure, the concentration of methanoic acid vapour in the atmosphere should not exceed 28.8 mg m ⁻³ . Some descalers for kettles, irons etc use methanoic acid.			
Concentrated acids Solution in water (If 4 M or more ethanoic acid; 2.2 M or more methanoic acid)	CORROSIVE	DANGER: both acids cause severe skin burns and eye damage.			
Moderately-concentrated acids (If less than 4 M but 1.7 M or more ethanoic acid; less than 2.2 M but 0.4 M or more methanoic acid)	IRRITANT	WARNING: both acids are irritating to the eyes and skin. Ant venom is about 1.5 M methanoic acid; stinging nettles are less concentrated.			
Dilute acids (If less than 1.7 M ethanoic acid; less than 0.4 M methanoic acid)	LOW HAZARD	Both acids may still cause harm in the eyes or in a cut. This includes vinegar (about 0.8 M ethanoic acid).			

Typical control measures to reduce risk

- Use the lowest concentration possible.
- Use the smallest volume possible.
- Wear eye protection for all but the most-dilute solutions; goggles for concentrated acids.
- Wear protective gloves if anything more than tiny amounts of concentrated acid is handled.
- Avoid breathing gas or fumes from concentrated solutions, eg, by use of a fume cupboard.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong?

eg, choking fumes if concentrated acid is over-heated. .

How serious would it be if something did go wrong?

NB We put vinegar on chips!

How can the risk(s) be controlled for this activity?

eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

Emergency action

•	n the eye	Flood the eye wi	ith gently-runni	ing tap water f	or 10) minutes. (Consult a medic.
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Vapour breathed in Remove to fresh air. Consult a medic if breathing is difficult.

Swallowed Do no more than wash out the mouth with drinking water. Do not induce vomiting. Consult a

medic.

Spilt on the skin or clothing

Remove contaminated clothing. Then drench the skin with plenty of water. If a large area is affected or blistering occurs, consult a medic.

Spilt on the floor, bench, etc

Wipe up small amounts with a damp cloth and rinse it well.

For larger amounts, and especially for (moderately) concentrated acid, cover with mineral absorbent (eg, cat litter) and scoop into a bucket. Neutralise with sodium carbonate. Rinse with

plenty of water.

Phosphoric(V) acid

also known as orthophosphoric acid

Substance	Hazard	Comment
Concentrated phosphoric(V) acid (If 2.6 M or more) Usually sold as 85% w/w (about 9 M)	CORROSIVE	DANGER: cause severe skin burns and eye damage. For a 15-minute exposure, the concentration of phosphoric acid vapour in the atmosphere should not exceed 2.0 mg m ⁻³ . It reacts violently, becoming very hot, when mixed with water. It decomposes if heated strongly, forming toxic oxides of phosphorus. This includes phosphoric acid used in most descaling and rust treatments.
Moderately-concentrated phosphoric(V) acid (If less than 2.6 M nut 1.0 M or	<u>(1)</u>	WARNING: irritating to eyes and skin.
more)	IRRITANT	
Dilute phosphoric(V) acid (If less than 1.0 M)	LOW HAZARD	May still cause harm in the eyes or in a cut. This includes acid used in cola drinks, etc. Over long periods of time, the acid may attack teeth.

Typical control measures to reduce risk

- Use the lowest concentration possible.
- Use the smallest volume possible.
- Wear eye protection for all but the most-dilute solutions; goggles for concentrated acids.
- Wear protective gloves if anything more than tiny amounts of concentrated acid is handled.
- Add the concentrated acid slowly to cold water when diluting, never the reverse; stir frequently to ensure good mixing.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong? eg, hazardous products are formed in reactions with the acid or corrosive fumes are produced if concentrated acid is over-heated.
- How serious would it be if something did go wrong? eg, the skin and eyes can be seriously burned if not treated quickly.

plenty of water.

How can the risk(s) be controlled for this activity? eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

Ξ	Emergency action				
•	In the eye	Flood the eye with gently-running tap water for 10 minutes. Consult a medic.			
•	Swallowed	Do no more than wash out the mouth with drinking water. Do not induce vomiting. Consult a medic.			
•	Spilt on the skin or clothing	Remove contaminated clothing. Especially with concentrated acid, quickly use a dry cloth or paper towel to wipe as much liquid as possible off the skin. Then drench the skin with plenty of water. If a large area is affected or blistering occurs, consult a medic.			
•	Spilt on the floor,	Wipe up small amounts with a damp cloth and rinse it well.			
	bench, etc	For larger amounts, and especially for (moderately) concentrated acid, cover with mineral absorbent (eg, cat litter) and scoop into a bucket. Neutralise with sodium carbonate. Rinse with			



Citric, Oxalic & Tartaric acids

2-hydroxypropane-1,2,3-tricarboxylic acid, ethanedioic acid and 2,3-dihydroxybutanedioic acid

Substance	Hazard	Comment
2-hydroxypropane-1,2,3- tricarboxylic acid (citric acid), solid and most solutions (If 0.5 M or more)	IRRITANT	WARNING: causes serious eye and skin irritation and may cause respiratory irritation. It is an approved food additive, E330. Concentrated lemon juice may contain 2-hydroxypropane-1,2,3-tricarboxylic acid (citric acid) up to 1.7 M.
Dilute 2-hydroxypropane- 1,2,3-tricarboxylic acid (If less than 0.5 M)	LOW HAZARD	Even dilute solutions will cause discomfort in the eye. It is found in citrus fruits, eg, lemons, oranges, grapefruit. Lemons contain 2-hydroxypropane-1,2,3-tricarboxylic acid up to about 0.25 M.
Ethanedioic acid (oxalic acid), solid and most solutions (If 0.1 M or more)	HARMFUL	Warning: harmful in contact with the skin and if swallowed. May cause eye damage. It removes calcium ions from the blood, forming insoluble calcium ethanedioate (calcium oxalate); this can block kidneys. It is found in rhubarb, especially in the leaves and in unripe leaf stalks (ie, the part which is eaten). Cases of poisoning have been reported, although very rarely fatal. The toxic effects of rhubarb may be due to other substances.
Dilute ethanedioic acid (If less than 0.1 M)	LOW HAZARD	Even dilute solutions will cause discomfort in the eye.
2,3-dihydroxy-butanedioic acid (tartaric acid), solid and most solutions (If 0.7 M or more)	IRRITANT	WARNING: causes serious eye and skin irritation and may cause respiratory irritation. It is an approved food additive, E334. It is used in baking powder and is found in many food products.
Dilute 2,3-dihydroxy- butanedioic acid (If less than 0.7 M)	LOW HAZARD	Even dilute solutions will cause discomfort in the eye.

Typical control measures to reduce risk

- Use the lowest concentration possible.
- Wear eye protection for all but the most-dilute solutions; goggles for concentrated acids.
- Avoid the possibility of swallowing oxalic acid or its salts, eg, by using a safety pipette filler.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong? eg, specks of solid acid transferred into the eye, by rubbing with a contaminated finger.
- How serious would it be if something did go wrong?
- How can the risk(s) be controlled for this activity?

eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

•	In the eye	Flood the eye with gently-running tap water for 10 minutes. Consult a medic.
•	Swallowed	Do no more than wash out the mouth with drinking water. Do not induce vomiting. Consult a medic.
•	Spilt on the skin or clothing	Remove contaminated clothing. Then drench the skin with plenty of water.
•	Spilt on the floor, bench, etc	Wipe up small amounts with a damp cloth and rinse it well. For larger amounts, cover with mineral absorbent (eg, cat litter) and scoop into a bucket. Neutralise with sodium carbonate. Rinse with plenty of water.

Salicylic acid, aspirin, salol, oil of wintergreen

2-hydroxybenzoic acid, 2-ethanoyloxybenzoic acid, phenyl 2hydroxybenzoate, methyl 2-hydroxybenzoate

Substance	Hazard	Comment
2-hydroxybenzoic acid (Salicylic acid) solid	IRRITANT CORROSIVE	DANGER: causes serious eye damage; harmful if swallowed.
2-ethanoyloxybenzoic acid (Aspirin, o-acetylsalicylic acid) solid	IRRITANT	Warning: Causes skin and serious eye irritation; may cause respiratory irritation; harmful if swallowed. For a 15-minute exposure, the concentration in the atmosphere should not exceed 15 mg m ⁻³ . Aspirin (and other pain relief medicines) must not be sold in packs of more than 16, except in pharmacies where they can contain 32 tablets. It is illegal to sell more than 100 tablets or capsules in any one retail transaction. Used as a medication to treat pain, fever, and inflammation. Aspirin given shortly after a heart attack decreases the risk of death and is also used long-term to help prevent heart attacks, strokes, and blood clots, in people at high risk. Common side effects include an upset stomach. Aspirin can be extracted from the leaves of willow trees and has been used for its health effects for hundreds of years.
Phenyl 2-hydroxybenzoate (Salol, phenyl salicylate) solid	IRRITANT ENVIRONMENT	WARNING: Causes skin and serious eye irritation; may cause respiratory irritation; may also be labelled toxic to aquatic life with long-lasting effects. Has a relatively low melting point (41 °C) so releases a significant amount of vapour when heated. Insert a mineral wool plug in test tubes to minimise escape of vapour and ensure good ventilation.
Methyl 2-hydroxybenzoate (Oil of wintergreen, methyl salicylate) <i>liquid</i>	IRRITANT	WARNING: Causes skin and serious eye irritation; may cause respiratory irritation; harmful if swallowed. Used for pain relief, especially for muscles and joints where it is rubbed into the skin & flavouring agent in chewing gums, mints and mouth washes.

Typical control measures to reduce risk

- Use the smallest amount possible.
- Wear eye protection.
- Ensure good ventilation.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong? eg, specks of solid acid transferred into the eye, by rubbing with a contaminated finger.
- How serious would it be if something did go wrong?
- How can the risk(s) be controlled for this activity?

eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

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•	In the eye	Flood the eye with gently-running tap water for 10 minutes. Consult a medic.			
•	Swallowed	Do no more than wash out the mouth with drinking water. Do not induce vomiting. Consulte a medic.			
•	Spilt on the skin or clothing	Remove contaminated clothing. Then drench the skin with plenty of water.			
•	Spilt on the floor, bench, etc	Wipe up small amounts with a damp cloth and rinse it well. Brush up larger amounts, trying to avoid raising dust. Rinse with plenty of water.			

Ammonia (gas & solution)

also applies to Ammonium hydroxide

Substance	Hazard	Comment
Ammonia (Gas)	CORROSIVE TOXIC	DANGER: Causes severe skin burns and eye damage. Toxic if inhaled. Flammable gas. Very toxic to aquatic organisms. For a 15-minute exposure, the concentration in the atmosphere should not exceed 25 mg m ⁻³ . Effects of exposure develop or increase over some time. Inhalation
	ENVIRONMENT	may exacerbate the problems of those with asthma etc. However, the human nose can detect ammonia at well below danger levels.
Concentrated ammonia solution (If 3 M or more) Ammonium hydroxide solution; 35 % w/w (18 M) ammonia is commercially available, density 0.880 g cm ⁻³ , hence often called '880 ammonia'.	CORROSIVE IRRITANT ENVIRONMENT	Danger: Causes severe skin burns and eye damage. Respiratory irritant. Very toxic to aquatic organisms. Ammonia gas will be present and the pressure of gas builds up on hot days – open cautiously in a fume cupboard. 'Household' ammonia is about 6 M.
Moderately concentrated ammonia solution (If less than 3M but 1.8 M or more)	CORROSIVE	DANGER: Causes severe eye damage; irritates skin. Its odour can cause distress.
Moderately dilute ammonia solution (If less than 1.8 M but 0.6 M or more)	IRRITANT.	WARNING: irritates skin and eyes.
Dilute ammonia solution (If less than 0.6 M)	LOW HAZARD	It may still cause harm in eyes or in a cut.

Typical control measures to reduce risk

- Use the lowest concentration possible.
- Wear suitable eye protection including when making or disposing of solutions
- Use a fume cupboard for all but test tube amounts of the gas and more concentrated solutions (including opening bottles); ensure good laboratory ventilation.
- If smelling the gas, follow the safe technique for sniffing gases: use your hand to waft the gas towards your nose.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong?

eg, solution spurting out of test tubes when being heated; release of ammonia gas as a product of a chemical reaction; possibility of the ammonia gas concentration reaching dangerous levels.

- How serious would it be if something did go wrong? NB Alkali in the eye causes more damage than acid of equivalent concentration.
- How can the risk(s) be controlled for this activity?

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Ξ	mergency action	
•	In the eye	Flood the eye with gently-running tap water for at least 20 minutes. Consult a medic. If it is necessary to go to hospital, continue washing the eye during the journey in an ambulance.
•	Vapour breathed in	Remove the casualty to fresh air. Consult a medic if breathing is difficult.
•	Swallowed	Do no more than wash out the mouth with drinking water. Do not induce vomiting. Consult a medic.
•	Spilt on the skin or clothing	Remove contaminated clothing. Then drench the skin with plenty of water. If a large area is affected or blistering occurs, consult a medic .
•	Spilt on the floor, bench, etc	Consider the need to evacuate the laboratory and open windows if large amounts are spilt and especially for (moderately) concentrated solutions. Cover with mineral absorbent (eg, cat litter) and scoop into a bucket. Neutralise with citric acid. Rinse with plenty of water. Wipe up small amounts with a damp cloth and rinse it well.

Sodium hydroxide

also applies to Soda lime and Potassium hydroxide

Substance	Hazard	Comment
Sodium or potassium hydroxide Solid Also known as caustic soda and caustic potash. Soda lime contains about 5% sodium hydroxide, 1% potassium hydroxide, 0.2% silicon dioxide, 14 – 19% water and the remainder calcium hydroxide (it is used to absorb carbon dioxide). Carbosorb in addition contains an indicator.	CORROSIVE	DANGER: causes severe skin burns and eye damage. Potassium hydroxide is also harmful if swallowed. It gives out heat when added to water which can cause boiling or create a choking mist. It is used in the home for clearing drains.
Sodium or potassium hydroxide solution (Sodium hydroxide if 0.5 M or more; potassium hydroxide if 0.4 M or more)	CORROSIVE	DANGER: causes severe skin burns and eye damage. Potassium hydroxide is also harmful if swallowed if 3 M or more. Fehling's solution contains sodium hydroxide of this concentration. It is used in the home as an oven cleaner.
Dilute sodium or potassium hydroxide solution (Sodium hydroxide If less than 0.5 M but 0.125 M or more; potassium hydroxide if less than 0.4 M but 0.1 M or more)	IRRITANT	WARNING: irritating to the eyes and skin.
Very dilute sodium or potassium hydroxide solution (Sodium hydroxide If less than 0.125 M; potassium hydroxide if less than 0.1 M).	LOW HAZARD	It may still cause harm in the eyes or in a cut.

Typical control measures to reduce risk

- Use the lowest concentration possible; avoid using the solid if possible.
- Use the smallest amount possible.
- Wear eye protection, including when making or disposing of solutions. Goggles (or a face shield) rather than safety spectacles will be necessary if the chemical is classed as CORROSIVE at the concentration used.
- Wear protective gloves if the concentrated solution is handled in more than tiny amounts.
- If possible, use a safer alternative, eg, sodium carbonate when making salts or Benedict's solution rather than Fehling's solution for food tests.

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong? eg, solution spurting out of test tubes when being heated.
- How serious would it be if something did go wrong? NB Alkali in the eye causes more damage than acid of equivalent concentration.
- How can the risk(s) be controlled for this activity? eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

Ξ	Emergency action				
•	In the eye	Flood the eye with gently-running tap water for at least 20 minutes. Consult a medic. If it is necessary to go to hospital, continue washing the eye during the journey in an ambulance.			
•	Swallowed	Do no more than wash out the mouth with drinking water. Do not induce vomiting Consult a medic.			
•	Spilt on the skin or clothing	Remove contaminated clothing. Then drench the skin with plenty of water. If a large area is affected or blistering occurs, consult a medic.			
•	Spilt on the floor, bench, etc	Wipe up small amounts with a damp cloth and rinse it well. For larger amounts, and especially for (moderately) concentrated solutions, cover with mineral absorbent (eg, cat litter) and scoop into a bucket. Neutralise with citric acid. Rinse with plenty of water.			

Calcium hydroxide and oxide

including Slaked lime, Quick Lime and Limewater

Substance	Hazard	Comment
Calcium oxide Solid Quick lime Solid (Old samples of calcium oxide are mostly calcium hydroxide.)	CORROSIVE	DANGER: Causes serious eye damage and skin irritation. Reacts violently with water, becoming very hot - dust particles may shoot out. For a 15-minute exposure, the concentration of the powder in the atmosphere should not exceed 6 mg m ⁻³ . Used in making mortar, cement and concrete – careless handling of these has caused injuries in the building industry. Added to soil to neutralise the acidity.
Calcium hydroxide Solid Slaked lime, garden lime	CORROSIVE	DANGER: Causes serious eye damage and skin irritation. For a 15-minute exposure, the concentration of the powder in the atmosphere should not exceed 15 mg m ⁻³ . Used in making mortar, cement and concrete – careless handling of these has caused injuries in the building industry. Added to soil to neutralise the acidity.
Calcium hydroxide Solution Limewater Limewater is a saturated solution of calcium hydroxide, less than 0.02 M.	IRRITANT	Even a saturated solution of calcium hydroxide is so dilute that it is not classed as IRRITANT, despite pH = 12.4. However, limewater is usually made in schools by adding excess solid calcium hydroxide (or oxide) to water. Undissolved solid will remain and that is irritating to the eyes and skin (but any solid present might well be calcium carbonate, LOW HAZARD).

Typical control measures to reduce risk

- Wear eye protection when handling solids.
- Eye protection is advisable when using limewater, especially if blowing into it.

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong? eg, solid particles spitting when adding water, dust blowing around or liquid splashing into the eye when blowing into limewater.
- How serious would it be if something did go wrong? NB Alkali in the eye causes more damage than acid of equivalent concentration.
- How can the risk(s) be controlled for this activity? eq, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

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Ξ	mergency action	
•	In the eye	Flood the eye with gently-running tap water for at least 20 minutes. Consult a medic. If it is necessary to go to hospital, continue washing the eye during the journey in an ambulance. Limewater is unlikely to cause serious problems; flood the eye with gently-running tap water for at least 10 minutes. Consult a medic if there are any concerns.
•	Swallowed	Do no more than wash out the mouth with drinking water. Do not induce vomiting Limewater is unlikely to cause serious problems. Wash out the mouth. Consult a medic if there are any concerns.
•	Spilt on the skin or clothing	Brush off the solid. Remove contaminated clothing. Drench the skin with plenty of water. If a large area is affected or blistering occurs, consult a medic. For limewater, wash with water.
•	Spilt on the floor, bench, etc	Wipe up limewater or small amounts of solid with a damp cloth and rinse it well. For larger amounts of solid, scoop into a bucket, add water to the area followed by mineral absorbent (eg, cat litter).



Sodium, Potassium, Magnesium, Calcium **Carbonates**

including Hydrogencarbonates (bicarbonates)

Substance	Hazard	Comment
Sodium carbonate Hydrated or anhydrous solid Washing soda	IRRITANT	WARNING: cause serious eye irritation. Anhydrous solid presents a bigger risk because it is finely powdered, whereas the hydrate is crystalline (usually large crystals). Soda ash is an impure (industrial) form.
Potassium carbonate		Hydrated sodium carbonate is used as washing soda. They are approved food additives, E500 & E501.
Sodium & Potassium carbonate Concentrated solution (<i>If 0.8 M or more</i>)	IRRITANT	WARNING: cause serious eye irritation.
Sodium & Potassium carbonate Dilute solution (<i>If less than 0.8 M</i>)	LOW HAZARD	They are used as a dilute solution in bath salts.
Sodium & Potassium hydrogencarbonate (bicarbonate) Solid and solution Also Sodium sesquicarbonate	LOW HAZARD	They liberate carbon dioxide on gentle heating (or with acids). Sodium hydrogencarbonate is an approved food additive, E500, and is used as baking soda. 'Bicarb' or 'bicarbonate of soda' are old-fashioned names. Bath salts contain sodium sesquicarbonate, (sodium carbonate hydrogencarbonate), $Na_2CO_3.NaHCO_3.H_2O$.
Calcium & Magnesium carbonate Magnesium carbonate hydroxide ('light' and 'heavy' magnesium carbonate) Solid	LOW HAZARD	Calcium carbonate occurs naturally as calcite, marble, chalk and limestone. It decomposes if heated above 800 °C and gives carbon dioxide with acids (unless the calcium salt is insoluble in water). It is an approved food additive, E170. Blackboard chalk may be calcium carbonate but is more likely to be calcium sulfate. Magnesium carbonate also occurs naturally. It decomposes more easily than calcium carbonate and is also an approved food additive, E504.
Calcium & Magnesium hydrogencarbonate (bicarbonate) Solution	LOW HAZARD	The solid does not exist. The solution is the cause of temporary hardness of water. On warming, calcium (or magnesium) carbonate) is deposited as 'limescale' or 'fur', eg on kettles. The solution reacts with soap to produce insoluble calcium (or magnesium) salts ('scum').

Typical control measures to reduce risk

- Wear eye protection when handling irritant solids and avoid raising dust.
- Avoid contact with acids, especially in sealed containers, because large volumes of carbon dioxide will be formed.

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong? eg, specks of solid transferred into the eye, by rubbing with a contaminated finger.
- How serious would it be if something did go wrong?
- How can the risk(s) be controlled for this activity?
 - eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

E m	era	Δn	CV	cti	on

•	In the eye	Flood the eye with gently-running tap water for 10 minutes. Consult a medic.
•	Swallowed	Do no more than wash out the mouth with drinking water. Do not induce vomiting. If there are any concerns, consult a medic.
•	Spilt on the skin or clothing	Brush solid off contaminated clothing. Rinse clothing or skin as necessary.
•	Spilt on the floor, bench, etc	Brush up solid spills, trying to avoid raising dust, then wipe with a damp cloth. Wipe up solution spills with a cloth and rinse it well.

Sodium and **potassium salts**

Substance	Hazard	Comment	
Sodium and potassium chloride, bromide and iodide Solid & solution	LOW HAZARD	People have been killed through consuming very large amounts of salt. At 'normal' levels, can cause high blood pressure, hence heart disease. Adults should not eat more than 6 g/day, children less. Potassium chloride is approved food additive, E508, used as a 'low-salt' substitute.	
Sodium sulfate(VI) Also potassium sulfate(VI) Solid & solution	LOW HAZARD	Hydrated sodium sulfate(VI), $Na_2SO_4.10H_2O$ is known as Glauber's salt. Sodium sulfate(VI) is an approved food additive, E514, as is the potassium salt, E515.	
Sodium hydrogensulfate(VI) (bisulfate) Solid and concentrated solution (If 0.25 M or more)	CORROSIVE	DANGER: Causes serious eye damage. It is strongly acidic because of acidic hydrogen in NaHSO $_4$. It is used in some toilet cleaners.	
Sodium hydrogensulfate(VI) (bisulfate) Dilute solutions (less than 0.2M but more than 0.1M)	IRRITANT	WARNING: Causes serious eye irritation. It is strongly acidic because of acidic hydrogen in NaHSO ₄ .	
Sodium hydrogensulfate(VI) (bisulfate) Very dilute solutions (< 0.1M)	LOW HAZARD	It is strongly acidic because of acidic hydrogen in NaHSO ₄ .	
Sodium and potassium ethanedioate (oxalate) Solid	HARMFUL	WARNING: harmful if swallowed and in contact with skin.	
Sodium and potassium ethanedioate (oxalate) Solution	LOW HAZARD	Note that all sodium and potassium ethanedioate (oxalate) solutions are LOW HAZARD EXCEPT WARNING: potassium ethanedioate is harmful if swallowed if \geq 1.5 M.	
Sodium nitrate(V) Also potassium nitrate(V) Solid and solutions	OXID. IRRITANT	WARNING: Oxidiser Sodium nitrate(V) solid and solutions are harmful if swallowed (more than 3 M) and irritating to eyes and skin (more than 1 M).	
Sodium or potassium carbonate	and hydrogence	arbonate. See CLEAPSS Student Safety Sheet 33	
Sodium chlorate(I) (hypochlorite	e)	See CLEAPSS Student Safety Sheet 41.	
Sodium or potassium chromate(VI) or dichroma	te(VI) See CLEAPSS Student Safety Sheet 47.	
Sodium or potassium manganat	e(VII) (permang	anate) See CLEAPSS Student Safety Sheet 48.	

Typical control measures to reduce risk

Wear eye protection when handling hazardous solids and solutions and avoid raising dust.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong? eg, specks of solid transferred into the eye, by rubbing with a contaminated finger.
- How serious would it be if something did go wrong? eg, are there hazardous reaction products (such as chlorine from the electrolysis of sodium chloride)?
- How can the risk(s) be controlled for this activity? eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

- Flood the eye with gently-running tap water for 10 minutes. Consult a medic if pain In the eye persists. Do no more than wash out the mouth with drinking water. Do **not** induce vomiting. **Swallowed** Consult a medic. Brush solid off contaminated clothing. Rinse skin or clothing as necessary. Spilt on the skin or clothing
- Spilt on the floor, bench, etc Brush up solid spills, trying not to raise dust, then wipe with a damp cloth. Wipe up solutions spills and rinse well.

Sodium sulfites, thiosulfate & persulfate

including metabisulfite & potassium Salts

Substance	Hazard	Comment
Sodium & potassium sulfite [sulfate(IV)]; sodium & potassium metabisulfite [disulfate(IV)] Solid and concentrated solution (If 0.15 M or more)	HARMFUL CORROS.	DANGER: Harmful if swallowed, cause serious eye damage. With acids, produce sulfur dioxide (SO ₂) (a TOXIC gas, see <i>Sheet 52</i>); do not inhale. Smell of SO ₂ due to acidification by CO ₂ in air. Approved food additives: sodium sulphite E221, potassium sulphite E225, sodium metabisulfite E223, potassium metabisulfite E224 all used as preservatives. For a 15-minute exposure, concentration of metabisulfite in the atmosphere should not exceed 15 mg m ⁻³ .
Sodium & potassium sulfite [sulfate(IV)]; sodium & potassium metabisulfite [disulfate(IV)] Dilute solution (If less than 0.15 M)	LOW HAZARD	They smell of sulfur dioxide due to acidification by carbon dioxide in the air; do not inhale.
Sodium & potassium hydrogensulfite [hydrogensulfate(IV)] Concentrated solution (If 0.15 M or more)	HARMFUL CORROS.	DANGER: Harmful if swallowed, cause serious eye damage. With acids, produce sulfur dioxide (SO_2) (a TOXIC gas, see <i>Sheet 52</i>); do not inhale. Smell of SO_2 due to acidification by CO_2 in air. Sodium and potassium hydrogensulfites are approved food additives, E222 and E228, as preservatives. The pure solid NaHSO ₃ does not exist. Products sold as bisulfite contain metabisulfite.
Sodium & potassium metabisulfite [disulfate(IV)] Dilute solution (If less than 0.15 M)	LOW HAZARD	Although sodium metabisulfite solid is $Na_2S_2O_5$, it behaves as sodium hydrogensulfite, $NaHSO_3$, in solution.
Sodium & potassium thiosulfate Solid and solutions	LOW HAZARD	Produce sulfur (see CLEAPSS Student Safety Sheet 82) & sulfur dioxide (TOXIC gas, see Sheet 52) with acids, including carbon dioxide. Carbon dioxide may cause solutions to go cloudy.
Sodium & potassium persulfate [peroxodisulfate(VI)] Solid and most solutions (If 0.04 M or more)	OXIDISER IRRITANT HEALTH HAZARD	Danger: oxidisers; skin irritants; cause serious eye irritation; harmful if swallowed; may cause respiratory irritation, allergy or asthma. $Na_2S_2O_8$ is used, for bleaching hair, etching printed-circuit boards and to initiate polymerisation reactions. Solutions are low hazard if less than 0.04M.
Sodium sulfate(VI) and hydrogensulfa	ite(VI)	See CLEAPSS Student Safety Sheet 34.

Typical control measures to reduce risk

- Use the smallest quantity or concentration possible.
- Wear eye protection when handling hazardous solids and solutions.
- Take care not to inhale sulfur dioxide; asthmatics should be especially careful; use a fume cupboard to avoid exposure.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong? eg, accidental inhalation of sulfur dioxide when opening a bottle or dissolving a solid in water.
- How serious would it be if something did go wrong?
- How can the risk(s) be controlled for this activity?

eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

Emergency action

Flood eye with gently-running tap water for 10 minutes. Consult a medic if pain persists. In the eye Do no more than wash out the mouth with drinking water. Do **not** induce vomiting. **Swallowed** Consult a medic. Brush solid off contaminated clothing. Rinse skin or clothing as necessary.

Spilt on the skin or clothing

Spilt on the floor, bench, etc Brush up solid spills, trying not to raise dust, then wipe with a damp cloth. Wipe up solutions spills and rinse well.

Magnesium and Calcium salts

Substance	Hazard	Comment
Magnesium chloride	LOW HAZARD	It is an approved food additive, E511.
Calcium chloride Anhydrous / hydrated solid & conc. solution (if 0.9M or more)	IRRITANT	WARNING: causes skin and serious eye irritation and may cause respiratory irritation. Anhydrous calcium chloride can cause water to boil; often used as a drying agent. Approved food additive E509.
Calcium chloride Dilute solution (if less than 0.9 M)	LOW HAZARD	-
Magnesium sulfate(VI) Hydrated and anhydrous	LOW HAZARD	Hydrated magnesium sulfate, known as Epsom salts, is used as a medicine (a purgative). It is a cause of permanently hard water.
Calcium sulfate(VI) CaSO ₄ . ½H ₂ O, plaster of Paris CaSO ₄ .2H ₂ O, gypsum	LOW HAZARD	Plaster of Paris used for setting broken bones: CaSO ₄ .½H ₂ O absorbs water, becomes hot and expands slightly forming CaSO ₄ .2H ₂ O (gypsum). It is not safe to attempt to make a cast of the whole hand or encase any other part of the body. Unlike magnesium sulfate(VI), calcium sulfate(VI) is only sparingly soluble. It is commonly used as blackboard chalk. It is a cause of permanently-hard water. It is an approved food additive, E516.
Magnesium nitrate Solid	OXID. IRRIT.	DANGER: may intensify fire; causes skin and serious eye irritation and may cause respiratory irritation.
Magnesium nitrate Most solutions (if 0.5M or more)	IRRITANT	WARNING: irritating to eyes and skin.
Magnesium nitrate Dilute solutions (if less than 0.5M)	LOW HAZARD	-
Calcium nitrate Solid and most solutions (if 0.15M or more)	OXID. CORROS.	DANGER: the solid may intensify fire; solid and solutions cause serious eye damage.
Calcium nitrate Dilute solutions (if less than 0.15M but 0.05M or more)	IRRITANT	WARNING: irritating to eyes.
Calcium nitrate <i>Very dilute solution (if less than 0.05M)</i>	LOW HAZARD	-
Calcium (and magnesium) carbonate ar	nd hydrogenc a	
Calcium oxalate (ethanedioate)		See CLEAPSS Student Safety Sheet 25

Typical control measures to reduce risk

- Wear eye protection when handling hazardous solids and solutions.
- Avoid raising dust.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong? eg, specks of solid transferred into the eye, by rubbing with a contaminated finger.
- How serious would it be if something did go wrong?
- How can the risk(s) be controlled for this activity?

eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

Emergency action

In the eye Flood the eye with gently-running tap water for 10 minutes. Consult a medic if pain

persists.

Swallowed Do no more than wash out the mouth with drinking water. Do **not** induce vomiting.

Consult a medic.

Spilt on the skin or clothing Brush solid off contaminated clothing. Rinse clothing or skin as necessary.

Spilt on the floor, bench, etc Brush up solid spills, trying to avoid raising dust, then wipe with a damp cloth.

Wipe up solution spills with a cloth and rinse it well.

Ammonium salts

Substance	Hazard	Comment
Ammonium chloride		WARNING: harmful if swallowed, causes serious eye irritation.
Solid and concentrated solution	^	It decomposes on heating to form ammonia gas (see CLEAPSS
(if 1.8 M or more)	(!)	Student Safety Sheet 30) and hydrogen chloride gas (see CLEAPSS
,		Student Safety Sheet 20) but they recombine on cooling.
	HARMFUL	Warming with alkali will generate ammonia gas.
		Old name: sal ammoniac.
Ammonium chloride	LOW HAZARD	Warming with alkali will generate ammonia gas (see CLEAPSS
Dilute solution (if less than 1.8 M)	LOW HAZARD	Student Safety Sheet 30).
Ammonium sulfate(VI)		It decomposes on heating to form ammonia gas (see CLEAPSS
Solid and solutions		Student Safety Sheet 30) and sulfuric acid 'gas' (see CLEAPSS
	LOW HAZARD	Student Safety Sheet 22) but they recombine on cooling.
		Warming with alkali will generate ammonia gas.
		It is an approved food additive, E517.
Ammonium carbonate		WARNING: harmful if swallowed. The solid decomposes, even at
Solid and concentrated solution		room temperature, to ammonia (see CLEAPSS Student Safety Sheet
(if 1.0 M or more)	<!-- -->	30) and carbon dioxide (see CLEAPSS Student Safety Sheet 58).
,		Warming with alkali generates ammonia gas.
	HARMFUL	It is an approved food additive, E503.
		Old name: sal volatile; used as smelling salts.
Ammonium carbonate	LOW HAZARD	Warming with alkali generates ammonia gas (see CLEAPSS Student
Dilute solution (if less than 1.0 M)	LOW HAZARD	Safety Sheet 30).
Ammonium nitrate(V)		WARNING: oxidiser, causes serious eye irritation, may cause skin or
Solid		respiratory irritation. It may decompose explosively if heated or on
		grinding; many industrial accidents have occurred in this way.
	OXIDISER IRRITANT	Warming with alkali will generate ammonia gas (see CLEAPSS
		Student Safety Sheet 30).
Ammonium nitrate(V)		The solution does have oxidising properties.
Solution	LOW HAZARD	Do not heat the solution to dryness.
	LOW HAZARD	Warming with alkali will generate ammonia gas (see CLEAPSS
		Student Safety Sheet 30).
Ammonium dichromate(VI)		See CLEAPSS Student Safety Sheet 47.

Typical control measures to reduce risk

- Wear eye protection when handling hazardous solids and solutions.
- Do not heat solid ammonium nitrate(V) and do not heat ammonium nitrate(V) solution to dryness.
- Avoid exposure to hazardous decomposition products if ammonium carbonate, chloride or sulfate(VI) are heated, eg, by using a fume cupboard.
- Avoid exposure to ammonia gas when reacting ammonium salts with alkalis, eg, by using a fume cupboard.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong? eg, specks of solid transferred into the eye, by rubbing with a contaminated finger.
- How serious would it be if something did go wrong?
- How can the risk(s) be controlled for this activity?
 - eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

- In the eye Flood the eye with gently-running tap water for 10 minutes. Consult a medic if pain persists.
- **Swallowed** Do no more than wash out the mouth with drinking water. Do **not** induce vomiting. Consult a medic.
- **Spilt on the skin or clothing** Brush solid off contaminated clothing. Rinse clothing or skin as necessary.
- **Spilt on the floor, bench, etc** Brush up solid spills, trying to avoid raising dust, then wipe with a damp cloth.
 - Wipe up solution spills with a cloth and rinse it well.

Iron and its compounds

including Iron(II) and iron(III) oxides, carbonates, sulfates(VI), chlorides, bromide *Note:* iron(II) compounds are often called **ferrous** and iron(III) compounds **ferric**

Substance	Hazard	Comment
Iron metal (powder)	FLAMMABLE	WARNING: the powder is a flammable solid. Iron filings/powder in the eye are very painful because the iron oxidises rapidly in the saline environment. Samples of iron are often not very pure and on reacting with dilute acids may produce toxic hydrogen sulfide gas
Iron metal (filings, sheets or bars of metal)	LOW HAZARD	(smelling of bad eggs) (see CLEAPSS Student Safety Sheet 59). For reaction with sulfur, see CLEAPSS Student Safety Sheet 82. Iron often coated with zinc (galvanised) to protect it from corrosion.
Iron oxides <i>or</i> iron(II) carbonate	LOW HAZARD	Applies to all iron oxides: iron(III) oxide (haematite), iron(II) iron(III) oxide (magnetite or ferrosoferric oxide). Iron(II) carbonate is usually sold mixed with a sugar (saccharated), to slow down oxidation.
Iron(II) or iron(III) sulfate(VI) or ammonium iron(II) sulfate (Mohr's salt) or ammonium iron(III) sulfate (ferric alum) Solid or concentrated solutions (if 0.5 M or more for iron(II) or 0.3 M for iron(III)	IRRITANT	Warning: Causes skin and serious eye irritation. Iron(III) also harmful by ingestion if 0.6 M or more. Usually solutions are made up in dilute sulfuric acid (which may itself be hazardous – see CLEAPSS Student Safety Sheet 22) to slow down oxidation. Ammonium iron(II)/iron(III) solutions are more stable but are still made up in acid to limit oxidation.
Iron(II) or iron(III) sulfate(VI) or ammonium iron(II) or iron(III) sulfate Dilute solutions (if less than 0.5 M iron(II) or 0.3 M iron(III))	LOW HAZARD	Solutions which have been made up in sulfuric acid may be hazardous if the acid is more than 0.5 M (see CLEAPSS Student Safety Sheet 22).
Iron(II) or iron(III) chloride Hydrated or anhydrous solid or concentrated solutions (if 0.1 M or more).	HARMFUL CORROS.	DANGER: harmful if swallowed; causes skin irritation and serious eye damage. Some suppliers classify anhydrous solids and solutions more concentrated than 0.2M as corrosive. Usually solutions are made up in hydrochloric acid (see CLEAPSS Student Safety Sheet 20) to slow down oxidation. Solution (about 2 M) used for etching printed circuit boards.
Iron(II) or iron(III) chloride Dilute solutions (If less than 0.1M)	LOW HAZARD	Usually solutions are made up in hydrochloric acid (see CLEAPSS Student Safety Sheet 20) to slow down oxidation.

Typical control measures to reduce risk

- Use the lowest possible quantities and concentrations.
- Take care not to rub the eye with fingers contaminated with iron filings or powder.
- Wear eye protection.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong? eg, solutions spurting out of test tubes when heated or solutions heated to dryness and decomposing. .
- How serious would it be if something did go wrong?
- How can the risk(s) be controlled for this activity?

eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

Emergency action

In the eye Flood eye with gently-running tap water for 10 minutes. Consult a medic if pain persists. **Swallowed** Do no more than wash out the mouth with drinking water. Do **not** induce vomiting. Consult a medic.

Remove the casualty to fresh air. Consult a medic if breathing is difficult. Dust breathed in Spilt on the skin or clothing Brush solid off contaminated clothing. Rinse clothing or skin as necessary. Spilt on the floor, bench, Brush up solid spills, trying to avoid raising dust, then wipe with a damp cloth.

Wipe up small solution spills with a cloth and rinse it well. For larger spills use mineral

absorbent (eg, cat litter).

Boron compounds

including borax, boric acid, sodium perborate, sodium borohydride

Substance	Hazard Co	omment
Borax (sodium tetraborate,	^	DANGER: may damage fertility and the unborn child, although
disodiumtetraborate-10-water)		this classification is controversial.
Solid and concentrated solutions		It has been/is used in some laundry and cleaning products, as a
(if 80 g dm ⁻³ , ie 0.2 M or more)		fire retardant and as a food additive (E285, to aid food
,	HEALTH HAZARD	preservation and improve the texture).
Borax		The borax solution commonly used for making slime is usually
Dilute solutions	LOW HAZARD	either 80 g dm ⁻³ if using low molar mass PVA (< 85 000 g mol ⁻¹)
(if less than 80 g dm ⁻³ , less than 0.2 M)		or 40 g dm ⁻³ if using high molar mass PVA (> 85 000 g mol ⁻¹).
Boric acid (boracic acid)		DANGER: may damage fertility and the unborn child, although
Solid and concentrated solutions		this classification is controversial.
(if 0.9 M or more)		In solution used as a mild antiseptic. The powder is used as an
		insecticide and to treat wood that is rotten and as a food
	HEALTH HAZARD	additive (E284, to aid food preservation and improve the
		texture).
Boric acid (boracic acid)	LOW HAZARD	-
Dilute solutions (if less than 0.9 M)	2011111211112	
Sodium perborate (sodium	\wedge	DANGER: oxidiser; harmful if swallowed; causes serious damage
peroxoborate-4-water)		to eyes; may cause respiratory irritation; may damage unborn
	OXIDISER HARMFUL	child; suspected of damaging fertility.
	^	Used in the past in detergents, bleaches, cleaning products
		and for tooth-whitening but almost entirely replaced now.
		Releases oxygen if heated above 60°C, or in presence of
	CORROSIVE HEALTH	outury ou
Sodium borohydride (sodium		DANGER: Contact with water liberates flammable gases which
tetrahydridoborate(III))		may ignite spontaneously (hydrogen); toxic if swallowed;
	FLAMM. CORROSIVE	causes skin burns and eye damage; may damage fertility or the
		unborn child.
		Widely used in chemistry as a reducing agent.
	TOXIC	

Typical control measures to reduce risk

- Wear eye protection when handling hazardous solids & solutions;
- Wear gloves when handling these solids.
- Avoid the risk of inhaling dust from sodium tetraborate or boric acid, eg by weighing in a fume cupboard.
- Avoid naked flames when handling sodium borohydride.
- Slime made using sodium tetraborate should not be taken home; should only be handled wearing gloves.

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong?
- How serious would it be if something did go wrong?
- How can the risk(s) be controlled for this activity?
 - eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

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	mergency action	
•	In the eye	Flood the eye with gently-running tap water for 10 minutes. Consult a medic if pain
		persists.
•	Swallowed	Do no more than wash out the mouth with drinking water. Do not induce vomiting.
		Consult a medic.
•	Spilt on the skin or clothing	Brush solid off contaminated clothing. Rinse clothing or skin as necessary.
•	Spilt on the floor, bench, etc	Brush up solid spills, trying to avoid raising dust, then wipe with a damp cloth (EXCEPT
	-	sodium borohydride). Wipe up small solution spills with a cloth and rinse it well.



Copper and its compounds

including Copper oxides, carbonate, sulfate, chloride, nitrate

Substance	Hazard	Comment
Copper (metal)	LOW HAZARD	Sharp edges can present a risk of cuts.
Copper(I)/(II) oxides Cuprous / cupric oxide	CORR.* IRRITANT ENVIR.	DANGER. Copper(I) oxide: *causes serious eye damage; skin irritant; harmful if swallowed/inhaled; toxic to aquatic life. WARNING. Copper(II) oxide: causes serious eye irritation; skin; harmful if swallowed/ inhaled; toxic to aquatic life.
Copper(II) carbonate hydroxide Basic copper carbonate	IRRITANT ENVIRONMENT	WARNING. Copper(II) carbonate hydroxide: causes serious eye irritation; skin irritant; harmful if swallowed/inhaled, toxic to aquatic life. Also known as <i>malachite</i> .
Copper(II) sulfate and copper(II) nitrate solids and concentrated solutions	CORR. IRRITANT ENVIR*.	DANGER. Solids and solutions (≥ 1.0M sulfate, ≥ 1.3 M nitrate): skin irritant; cause serious eye damage; harmful if swallowed (especially saturated solutions for crystal-growing). Solid* only: very toxic to aquatic life. Water added to anhydrous solid copper(II) sulfate(VI) produces heat.
Copper(II) sulfate and copper(II) nitrate dilute solutions	CORROSIVE IRRITANT	DANGER. Sulfate/nitrate <1.0M/1.3M and ≥0.2M/0.15M: skin irritant; cause serious eye damage. WARNING Sulfate/nitrate <0.2M/0.15M and ≥0.02M/0.05M: irritating to skin and eyes. LOW HAZARD Sulfate/nitrate <0.02M/0.05M. Benedict's solution and Fehling's solution both contain dilute copper(II) sulfate(VI) but Fehling's solution has other hazards.
Copper(II) chloride solid	IRRITANT ENVIRONMENT	WARNING. Eyes; skin; harmful if swallowed; toxic to aquatic life.
Copper(II) chloride solution (if 0.8M or more)	IRRITANT	WARNING. Eyes; skin; toxic to aquatic life; harmful if swallowed (≥ 1.8M).
Copper(II) chloride solution (if less than 0.8M)	LOW HAZARD	

Typical control measures to reduce risk

- Use the lowest concentration possible.
- Avoid raising dust, eg by dampening powders
- Take care if evaporating solutions to dryness
- Wear eye protection when handling the solids and all but the most dilute solutions

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong?

eg, solutions spurting out of test tubes when heated or solutions decomposing to toxic products when heated to dryness

- How serious would it be if something did go wrong?
 - eg, are there hazardous reaction products (such as chlorine from the electrolysis of copper chloride)?
- How can the risk(s) be controlled for this activity?
 - eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

Emergency action

In the eye Flood the eye with gently-running tap water for 10 minutes. Consult a medic.

Swallowed Do no more than wash out the mouth with drinking water. Do not induce vomiting.

Consult a medic.

Dust breathed in Remove the casualty to fresh air. Consult a medic if breathing is difficult.

Spilt on the skin or clothing Remove contaminated clothing and rinse it. Wash off the skin with plenty of water

Spilt on the floor, bench, etc Scoop up solid (take care not to raise dust). Wipe up small solution spills or any traces of solid with cloth; for larger spills use mineral absorbent (eg, cat litter).

CLEAPSS

Student safety sheets

Sodium chlorate(I)

also known as Sodium hypochlorite

5	Substance					Haza	ra	C	omment	

Note that *solid* sodium chlorate(I) does not exist. Sodium chlorate(I) is NaOCI; do not confuse it with sodium chlorate(V), NaClO₃, or sodium chlorate(VII), NaClO₄.

$NaCiO_3$, or sodium chlorate(vii), $NaCiO_4$.				
Concentrated sodium chlorate(I)	^ ^	DANGER: causes severe burns and eye damage, similar to		
Solution	企图	sodium hydroxide solution. It is toxic to aquatic life.		
(if more than 0.7 M or more than 5% (w/v)		It produces a toxic gas (chlorine) with acids.		
available chlorine)	•	Pressure may build up in bottles during storage, due to		
,	CORROS. ENVIRON.	slow decomposition.		
		It removes the colour from many dyes.		
Moderately-dilute sodium chlorate(I)	^ ^	DANGER: causes severe eye damage; irritating to skin. It		
Solution	LE Y	is toxic to aquatic life.		
(if less than 0.7 M but 0.4 M or more; between		It produces a toxic gas (chlorine) with acids.		
5% and 3% (w/v) available chlorine)	CORROS. ENVIRON.	This includes most domestic bleach.		
	CONTROL ENVIRON.	It removes the colour from many dyes.		
Dilute sodium chlorate(I)		WARNING: irritating to eyes and skin.		
Solution		Microbiological spills can be dealt with using a 10%		
(if less than 0.4 M but 0.15 M or more;		solution diluted 100 times (ie, 0.1%), but it is quickly		
between 3% and 1% (w/v) available chlorine)	IRRITANT	made inactive by organic matter and so a 10 times		
		dilution (ie, 1%) is often preferred.		
Very dilute sodium chlorate(I)	LOW HAZARD	Microbiological spills can be dealt with using a 10%		
Solution		solution diluted 100 times (ie, 0.1%), but it is quickly		
(if less than 0.15 M; less than 1% (w/v)		made inactive by organic matter and so a 10 times		
available chlorine)		dilution (ie, 1%) is often preferred.		

Note: Available chlorine

Sodium chlorate(I) is normally made by reacting chlorine gas with sodium hydroxide solution. Sodium chloride is produced as a by-product and this is left mixed in the solution. So only part of the chlorine in the mixture (the Cl in the NaOCl but not the Cl in the NaCl) is available for oxidising or bleaching purposes. One gram of a 10% available chlorine bleach has the same bleaching power as 0.1 gram of pure chlorine.

Typical control measures to reduce risk

- Use the lowest concentration possible.
- Use the smallest volume possible.
- Wear eye protection, including when making or disposing of solutions.
- Wear protective gloves if the concentrated solution is handled on anything larger than a test-tube scale.
- Never mix domestic bleach with other household cleaners, because these could be acidic.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong?

 A paradous products of roaction (such as chloring age) are formed.

eg hazardous products of reaction (such as chlorine gas) are formed if sodium chlorate(I) is mixed with acid.

- How serious would it be if something did go wrong?
 - ${\it NB~Alkali~in~the~eye~causes~more~damage~than~acid~of~equivalent~concentration.}$
- How can the risk(s) be controlled for this activity?

eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

•	In the eye	Flood the eye with gently-running tap water for 20 minutes. Consult a medic. If a visit to hospital is necessary, continue washing the eye during the journey in an ambulance.
•	Chlorine breathed in	Remove the casualty to another room to rest. Consult a medic.
•	Swallowed	Do no more than wash out mouth with drinking water. Do not induce vomiting. Consult medic.
•	Spilt on the skin or clothing	Remove contaminated clothing. Drench the skin with plenty of water. If a large area is affected or blistering occurs, consult a medic. Rinse contaminated clothing with water.
•	Spilt on the floor, bench, etc	Wipe up small amounts with a damp cloth and rinse it well. For larger amounts, open the windows and, especially for quite-concentrated solutions, cover with mineral absorbent (eg, cat litter) and scoop into a bucket. Rinse with plenty of water.

Barium compounds

Substance	Hazard	Comment
Barium chloride		DANGER: toxic if swallowed, harmful if inhaled.
Solid	TOXIC	For a 15-minute exposure, the concentration of barium should not exceed 1.5 mg m ⁻³ .
Barium chloride Solution	!	WARNING: harmful if swallowed.
(if 0.4 M or more)	HARMFUL	
Barium chloride		-
Solution	LOW HAZARD	
(if less than 0.4 M)		
Barium nitrate(V)	^	DANGER: oxidiser; harmful if swallowed or inhaled.
and		For a 15-minute exposure, the concentration of barium should not
Barium peroxide	•	exceed 1.5 mg m ⁻³ .
Solids	OXIDISING HARMFUL	
Barium nitrate(V) Solution	LOW HAZARD	-
Barium sulfate(VI)		Unlike most barium compounds, barium sulfate(VI) is LOW HAZARD
Solid LOW HAZARD		because it does not dissolve in water or acids. Hence it is safe to eat a 'barium (sulfate) meal', before being X-rayed.

Typical control measures to reduce risk

- Use the lowest concentration possible.
- Use the smallest quantity possible.
- Wear eye protection.
- Wash hands after handling barium compounds.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong? eg, somebody drinking a toxic solution by mistake.
- How serious would it be if something did go wrong?
- How can the risk(s) be controlled for this activity?

eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

•	In the eye	Flood the eye with gently-running tap water for at least 10 minutes. Consult a medic.
•	Swallowed	Do no more than wash out the mouth with water. Do not induce vomiting. Consult a medic.
•	Spilt on the skin or	Brush off any solid. Remove contaminated clothing. Drench the skin with plenty of water
	clothing	Rinse contaminated clothing with water.
•	Spilt on the floor,	Scoop up any solid. Try to avoid raising dust. Rinse the area with water, diluting greatly.
	bench, etc	Solutions should be treated with mineral absorbent (eg, cat litter).

Lead and its compounds

including Lead oxides, bromide, nitrate(V), ethanoate (acetate) etc

Substance	Hazard	Comment
Lead (metal) and lead compounds (solid) ie, carbonates, oxides, sulfide, chloride, bromide, iodide, nitrate, sulfate, ethanoate (acetate), chromate. Lead nitrate, ethanoate (acetate) Concentrated solutions (if 1 M or more)	HARMFUL HEALTH HAZARD ENVIRONMENTAL HAZARD	DANGER: harmful if swallowed or inhaled; may damage organs through repeated or prolonged exposure; may damage the unborn child. Very toxic to aquatic life. Includes cerussite, white lead, litharge, massicot, red lead, galena. Lead chromate(VI) is used in the yellow lines on roads. Old paints often contained lead sulfate or other lead pigments. Young children may be at risk if they chew old objects painted with a lead-based paint. Tetraethyl lead was added to petrol to improve the combustion characteristics but leaded petrol has been phased out in most countries because when burnt in cars, it produces particles of lead compounds. Note that most lead compounds are insoluble in water or very nearly so. In soft-water areas, (old) lead pipes may very slowly dissolve exposing people to low levels of lead over very long periods of time.
Lead nitrate, ethanoate (acetate) Most solutions (if less than 1 M but 0.015 M or more)	HEALTH HAZARD	DANGER: harmful if swallowed or inhaled; may damage organs through repeated or prolonged exposure; may damage the unborn child. Very toxic to aquatic life.
Lead nitrate, ethanoate (acetate) Extremely dilute solutions (if less than 0.015 M)	LOW HAZARD	

Typical control measures to reduce risk

- Wear eye protection.
- Use the lowest possible amounts and concentrations
- Preferably, heat lead compounds in a fume cupboard; avoid raising dust (eg, by dampening powders).
- Less-volatile compounds (eg, oxides) may be heated in small amounts in a well-ventilated room (but not if those who are, or who might be, pregnant are present).
- Use lead nitrate rather than lead ethanoate (acetate) when a soluble lead salt is needed.
- Wash hands after using lead or its compounds.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong?

eg, solution spurts out of a test tube when heated or dust is breathed in.

- How serious would it be if something did go wrong?
 - eg, could anybody be exposed to dangerous lead levels for long periods of time?
- How can the risk(s) be controlled for this activity?

eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

•	In the eye	Flood the eye with gently-running tap water for at least 10 minutes. Consult a medic.
•	Swallowed	Do no more than wash out the mouth with water. Do not induce vomiting. Consult a medic.
•	Spilt on the skin or clothing	Brush off any solid. Remove contaminated clothing. Drench the skin with plenty of water. Rinse contaminated clothing with water.
•	Spilt on the floor,	Scoop up any solid. Try to avoid raising dust. Rinse the area with water, diluting greatly.
	bench, etc	Solutions should be treated with mineral absorbent (eg, cat litter).

Mercury and its compounds

including Mercury(I/II) oxides, chlorides, sulfides

Substance	Hazard	Comment
Mercury Metal	TOXIC HEALTH ENVIRONMENTAL HAZARD	DANGER: fatal if inhaled; causes damage to organs through prolonged or repeated exposure; may damage unborn child. Very toxic to aquatic life. Mercury is very dense - containers may be unexpectedly heavy - difficult to manipulate, eg in teat pipettes. Not trapped by filter fume cupboards. Clear up spills promptly, but do not panic. The main risk is from inhaling low concentrations of vapour over long periods of time from spills that were not noticed/cleared up. Wear gloves when handling mercury. It forms alloys with gold, silver, etc (eg, jewellery) - remove rings.
Mercury(II) chloride, mercury(I/II) oxides, sulfides Solids Mercury(II) chloride Solution (if 0.1 M or more)	TOXIC CORROSIVE HEALTH ENVIRO. HAZARD HAZARD	DANGER: fatal if swallowed or in contact with skin; causes severe burns and eye damage; suspected of causing genetic defects and damaging fertility; causes damage to organs through prolonged or repeated exposure. Very toxic to aquatic life. Mercury ('button') batteries contain mercury oxide and should be recycled. Mercury compounds used in making Victorian hats caused disease, as in the 'Mad hatter' of <i>Alice in Wonderland</i> .
Mercury(II) chloride Solution (if 0.01 M or more but less than 0.1 M)	TOXIC HEALTH	DANGER: fatal if swallowed or in contact with skin; suspected of causing genetic defects and damaging fertility; causes damage to organs through repeated or prolonged exposure. Very toxic to aquatic life.
Mercury(II) chloride Solution (if 0.002 M or more but less than 0.01 M) and Mercury(I) chloride, Mercury(II) sulfide Mercury(II) chloride	HARMFUL ENVIR.	WARNING: harmful if swallowed or in contact with skin; suspected of causing genetic defects and damaging fertility; causes damage to organs through repeated or prolonged exposure. Very toxic to aquatic life. Pollution by mercury compounds in a Japanese river in the 1950s caused serious poisoning of humans who ate river fish (Minimata disease). Note these solutions are extremely dilute.
Solution(if less than 0.002M)		

Typical control measures to reduce risk

- Wear eye protection and suitable gloves; use the lowest possible concentration.
- Avoid the use of mercury compounds where possible (eg, avoid Millon's reagent).
- Handle liquid mercury over a tray to contain spills; do not leave mercury surfaces exposed to the air.
- Avoid raising dust (eg, by dampening powder); work in a ducted (not filter) fume cupboard; clear up spills promptly.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong? eg, solution spurts out of a test tube when heated, mercury metal spills on the floor or a thermometer is broken.
- How serious would it be if something did go wrong? eg, could anybody be exposed to dangerous mercury levels for long periods of time?
- How can the risk(s) be controlled for this activity? eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

Emergency action

In the eye Flood the eye with gently-running tap water for at least 10 minutes. Consult a medic.

Swallowed There is little problem with mercury metal (but consult a medic). For compounds, do no more than wash out the mouth with water. Do **not** induce vomiting. Consult a medic.

Vapour Dangerous only if large amounts of vapour are breathed in over a short period of time (eg, when heating metal) or breathed in from long-term exposure. Consult a medic.

Spilt on the For mercury metal, remove contaminated clothing and wash off the skin. Check jewellery for damage. For mercury skin or clothing compounds, flood the area with large amounts of water. Remove and repeatedly rinse clothing. Consult a medic for large areas affected or if blistering occurs.

For metal, remove jewellery, collect mechanically (eg, with syringe). Mop up remainder with a hot paste of 1:1 Spilt on floor, bench, etc calcium oxide/sulfur in water. Spread same (dry) mixture over cracks etc. For compounds, scoop up solid. Rinse area with water, diluting greatly. For solutions, use mineral absorbent (eg, cat litter).

CLEAPSS

Student safety sheets

Aluminium and its compounds

including Aluminium oxide, hydroxide, sulfate(VI), chloride; also Potash alum

Substance	Hazard	Comment
Aluminium (metal)		Used in cooking utensils and generally considered safe. Suggestions at one
Solid (large pieces, sheets, etc)	LOW HAZARD	time it might cause Alzheimer's disease, now considered unlikely.
Aluminium (metal)		DANGER: flammable solid; in contact with water releases flammable gas
Fine powder	43	(hydrogen, see <i>CLEAPSS Student Safety Sheet 50</i>). A dust explosion is
	FLAMMABLE	possible if it is exposed to flame. Although difficult to ignite, it is difficult to
		extinguish. Used as a food additive, E173.
Aluminium oxide (alumina),	LOW HAZARD	Used in indigestion tablets.
or aluminium hydroxide	EOWITAZARD	
Aluminium potassium		Often used for crystal-growing in schools.
sulfate(VI) (potash alum)	LOW HAZARD	
Solid or solution		
Aluminium sulfate(VI)		DANGER: causes serious eye damage. Solutions are acidic. Used as a
Hydrated solid and most		mordant in dyeing. Added in small amounts to cloudy water in reservoirs
solutions (if 0.1 M or more)	CORROSIVE.	to coagulate clay particles. When large amounts were accidentally added
	00111001121	to a reservoir at Camelford in Cornwall, some ill effects were reported.
Aluminium sulfate(VI)		WARNING: irritating to eyes.
Dilute solutions (if less than 0.1	· · ·	
M but 0.03 M or more)	IRRITANT	
Aluminium sulfate(VI)	LOWILLAZADD	Note these solutions are extremely dilute.
Very dil. solns. (less than 0.03M)	LOW HAZARD	
Aluminium chloride		DANGER: causes severe skin burns and eye damage. Reacts exothermically
Anhydrous solid	₹ %	& violently with water to produce fumes of hydrogen chloride (see <i>Sheet</i>
	CORROSIVE	20). Pressure may build up in closed containers due to absorbed moisture.
Aluminium chloride	CORROSIVE	WARNING: irritating to eyes and skin (if ≥ 0.8 M) and respiratory system (if ≥
Hydrated solid or conc. solution	(!)	1.5 M).
(if 0.8 M or more)	· · ·	The solution is acidic.
	IRRITANT	
Aluminium chloride		In antiperspirants and deodorants, it is mixed with other substances and is
Dilute solution	LOW HAZARD	not classed as hazardous. However, it may be an IRRITANT to people with
(if less than 0.8 M)		sensitive skin.

Typical control measures to reduce risk

- Use the lowest possible concentration; wear eye protection.
- Avoid the use of aluminium powder or anhydrous aluminium chloride if possible.
- Avoid raising dust and keep aluminium powder away from naked flames.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong?
 eg, solution spurts out of a test tube when heated or a solution decomposes when heated to dryness.
- How serious would it be if something did go wrong?
 eg, are there hazardous reaction products, eg, hydrogen chloride gas from the action of water on anhydrous aluminium chloride?
- How can the risk(s) be controlled for this activity?
 eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

•	In the eye	Flood the eye with gently-running tap water for at least 10 minutes. Consult a medic.
•	Swallowed	Do no more than wash out the mouth with water. Do not induce vomiting. Consult a medic.
	Dust breathed in Spilt on the skin or clothing	Remove the casualty to fresh air. Consult a medic. if breathing is difficult. Remove and rinse contaminated clothing. Wash off the skin with plenty of water.
•	Spilt on floor, bench, etc	Scoop up the solid (take care not to raise dust). Wipe up small solution spills or any traces of solid with a cloth.
•	Fire	Powdered aluminium fires should be extinguished by smothering with clean dry sand

Silver and its compounds

including Silver bromide, chloride, iodide, nitrate(V) and oxide

Substance	Hazard	Comment
Silver (metal)		It is used in jewellery.
Solid	LOW HAZARD	It is an approved food additive, E174.
Silver halides,		Widely used in photographic emulsions. They are decomposed
ie, silver bromide, chloride		by light to give silver metal and the halogen (which then reacts
and iodide	LOW HAZARD	with other substances in the emulsion).
Solids		
Silver nitrate(V)	^	DANGER: oxidiser; causes severe skin burns and eye damage; very
Solid and fairly-concentrated		toxic to aquatic life.
solutions (If 0.3 M or more)		If swallowed, it may cause internal damage due to absorption
	OXIDISER CORROSIVE	into the blood, followed by deposition of silver in various
	*	tissues.
	32	The solid explodes dangerously with magnesium powder and a
	ENVIRONMENTAL HAZARD	drop of water. Accidents have caused many injuries and a very
		careful risk assessment is required before attempting this.
Silver nitrate(V)		DANGER: causes severe eye damage; irritating to skin.
Dilute solutions (if less than 0.3 M	<u>⟨</u> * &⟩	It may produce black stains on the skin, which, however, wear
but 0.18 M or more)	00000000	off in a few days.
Silver nitrate(V)	CORROSIVE	WARNING: irritating to eyes and skin.
Very dilute solutions (if less than		Very dilute solutions are adequate for most school work when
0.18 but 0.06 M or more).	V	testing for halides in solution.
0.10 but 0.00 W of more).	IRRITANT	
Silver nitrate(V)		
Extremely dilute solutions	LOW HAZARD	
(if less than 0.06 M)		
Silver nitrate(V) (ammoniacal)	\wedge	It is used for aldehyde tests and should be prepared only on a
ie, in ammonia solution	⟨♣⟩ ⟨!⟩	test-tube scale, when needed, and discarded into plenty of
(Tollen's Reagent)	EVELOUIVE IDDIT	water within ½ hour, otherwise explosives may form. Failure to
	EXPLOSIVE IRRITANT	do this has caused accidents.
Silver oxide	LOW HAZARD	It is used in some batteries, eg, button cells for watches and
Solid	LOTTINZARD	calculators.

Typical control measures to reduce risk

- Use the lowest possible concentration; wear eye protection.
- Avoid keeping solutions of silver compounds and ammonia for more than a few minutes.
- Avoid handling solid silver nitrate.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong?

Eg, Silver nitrate accidentally coming into contact with the skin.

- How serious would it be if something did go wrong?
 - Eg, Are there hazardous reaction products, eg, from solutions of silver compounds with ammonia?
- How can the risk(s) be controlled for this activity?

Eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

Emergency action

In the eye Flood the eye with gently-running tap water for at least 10 minutes. Consult a medic. **Swallowed** Do no more than wash out the mouth with water. Do not induce vomiting. Consult a medic. Spilt on the skin or Remove contaminated clothing and rinse it. Wash off the skin with plenty of water. If the clothing silver nitrate produces more than small burns, consult a medic . Spilt on floor, bench, etc Wear eye protection and gloves. Scoop up the solid. Rinse the area with water and wipe up,

rinsing repeatedly. Rinse the mop or cloth thoroughly.

Chromium & its compounds

including Chromium(III) salts, chrome alum, chromates and dichromates

Substance	Hazard	Comment
Chromium (metal) Solid	LOW HAZARD	Chromium plating gives a shiny, protective surface on steel.
Chromium(III) oxide Solid	LOW HAZARD	It is used as a green pigment, eg, in pottery.
Chromium(III) sulfate	LOW HAZARD	Solid & solutions. See below for chromium potassium sulfate.
Chromium(III) potassium sulfate(VI) (chrome alum) Solid and most solutions	IRRITANT	WARNING: irritating to eyes and skin (if 0.4 M or more). Often used for crystal growing. LOW HAZARD if less than 0.4 M.
Potassium chromate(VI) Solid and solutions	HEALTH IRRIT.	DANGER: skin & serious eye irritant; genetic effects, cancer by inhalation and allergic skin reaction. Vary toxic to aquatic life (if 0.9 M or more) DANGER: respiratory irritant (if 0.4 M or more) DANGER: skin sensitiser (if 0.04 M or more) DANGER: serious health hazard (if 0.01 M or more) LOW HAZARD if less than 0.01 M
Sodium chromate(VI) Solid and solutions	TOXIC CORRO. HEALTH ENVIR.	DANGER: toxic if swallowed; harmful in contact with skin; causes severe skin burns & eye damage; fatal if inhaled; may cause allergic skin reaction; may cause allergy or asthma if inhaled; may cause cancer or genetic defects; may damage fertility or the unborn child; causes damage to organs through prolonged or repeated exposure. Very toxic to aquatic life (if 0.8 M or more). DANGER: corrosive to skin and eyes; harmful by ingestion, respiratory irritant (if 0.2 M or more). DANGER: irritating to skin and eyes; skin & respiratory sensitiser; serious health hazard (if 0.01 M or more). Low HAZARD if less than 0.01 M
Ammonium, potassium, sodium dichromates(VI) Solid and solutions	OXID. TOXIC CORR. HEALTH ENVIRON.	DANGER: oxidiser; toxic if swallowed; harmful in contact with skin; causes severe skin burns & eye damage; fatal if inhaled; may cause allergic skin reaction; may cause allergy or asthma if inhaled; may cause cancer or genetic defects; may damage fertility or the unborn child; causes damage to organs through prolonged or repeated exposure. Very toxic to aquatic life (if 0.4 M or more). Solid ammonium dichromate decomposes if heated and will explode in confined spaces. It was used in indoor fireworks. DANGER: corrosive to skin and eyes; harmful by ingestion, respiratory irritant (if 0.1 M or more). DANGER: irritating to skin and eyes; skin & respiratory sensitiser; serious health hazard (if 0.004 M or more). Low HAZARD if less than 0.004 M
Lead chromate(VI)	•	See CLEAPSS Student Safety Sheet 43

Typical control measures to reduce risk

- Use the lowest possible concentration; wear eye protection; consider the use of gloves for chromates/dichromates.
- For volcano experiment, prevent exposure to dust by the use of a fume cupboard or mineral-wool plug in the vessel.
- Avoid inhaling chromate/dichromate dust or spray (eg, during electrolysis).

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong? eg, chromate or dichromate dust or solution is accidentally inhaled.
- How serious would it be if something did go wrong?
- How can the risk(s) be controlled for this activity?
 - eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

- In the eye Flood the eye with gently-running tap water for at least 10 minutes. Consult a medic.
- **Swallowed** Do no more than wash out the mouth with water. Do not induce vomiting. Consult a medic.
- Spilt on the Remove contaminated clothing and rinse until no colour remains. Wash off the skin with plenty of skin or clothing water. If skin contamination is more than small, consult a medic.
- Spilt on floor, Wear eye protection and gloves. Scoop up the solid. Rinse the area with water and wipe up, rinsing bench, etc repeatedly until no colour remains. Rinse the mop or cloth thoroughly.



Manganese and its compounds

including Manganese(IV) oxide (dioxide) & Potassium manganate(VII) (permanganate)

Substance	Hazard	Comment	
Manganese (metal) Solid	LOW HAZARD	Note that the powder would be highly flammable.	
Manganese(II) carbonate Solid	LOW HAZARD	Insoluble in water.	
Manganese(II) chloride Solid and concentrated solutions (if 1.5 M or more)	HARMFUL	WARNING: harmful if swallowed.	
Manganese(II) chloride Dilute solutions (if less than 1.5 M)	LOW HAZARD		
Manganese(II) sulfate Solid and concentrated solutions (if 0.7 M or more)	HEALTH ENVIRONMENT	WARNING: may cause damage to organs though prolonged or repeated exposure. Toxic to aquatic life with long-lasting effects.	
Manganese(II) sulfate Dilute solutions (if less than 0.7 M)	LOW HAZARD		
Manganese(IV) oxide (Manganese dioxide) Solid	HARMFUL	It is harmful by inhalation or if swallowed. It is often used as a fine powder. Many hazardous reactions occur with reducing agents or concentrated acids. It is used in dry cells (batteries). Insoluble in water.	
Potassium manganate(VII) (permanganate) Solid	OXID. HARMFUL ENVIR.	DANGER: oxidiser; harmful if swallowed; very toxic to aquatic life with long-lasting effects. Stains the hands and clothing. Many hazardous reactions occur with concentrated acids or reducing agents. On heating, releases a fine dust of potassium manganate(VI) [OXIDISER; IRRITANT] & oxygen gas.	
Potassium manganate(VII) (permanganate) Most solutions (if 0.1 M or more)	IRRITANT	WARNING: irritating to eyes and skin. Stains the hands and clothing.	
All manganates(VII) (permanganates) Very dilute solutions (if less than 0.1M)	LOW HAZARD	They stain hands and clothing.	

Typical control measures to reduce risk

- Wear eye protection.
- Avoid inhaling dusts.
- Avoid skin contact, especially with manganates(VII).
- Avoid contact between manganates(VII) or manganese(IV) oxide and concentrated acids or reducing agents.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong? eg, dust is accidentally inhaled.
- How serious would it be if something did go wrong? eg, are there hazardous reactions, eg, violent oxidations or decompositions?
- How can the risk(s) be controlled for this activity? eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

- In the eye Flood the eye with gently-running tap water for at least 10 minutes. Consult a medic.
- **Swallowed** Do no more than wash out the mouth with water. Do **not** induce vomiting. Consult a medic.
- Spilt on the Remove contaminated clothing and rinse it. Wash off the skin with plenty of water. [Manganate(VII) skin or clothing will give permanent stains to clothing but stains to the skin will wear off in a few days.] If skin contamination is more than small, consult a medic.
- Spilt on floor, Wear eye protection and gloves. Scoop up the solid. Rinse the area with water and wipe up, rinsing bench, etc repeatedly. [Manganate(VII) will give permanent stains.] Rinse the mop or cloth thoroughly.

Zinc and its compounds

including Zinc oxide, carbonate, sulfate(VI), chloride, bromide

Substance	Hazard	Comment
Zinc metal (granulated or sheets of metal)	LOW HAZARD	Pure zinc does not react readily with dilute acids, without a catalyst [usually copper(II) sulfate]. Iron or steel is often coated with zinc (galvanised) to protect it from rusting.
Zinc metal (powder or dust)	FLAM. ENVIR	DANGER: in contact with water releases flammable gases which ignite spontaneously; catches fire spontaneously if exposed to air; toxic to aquatic life with long-lasting effects. Reacts violently with iodine, sulfur and copper(II) oxide. Most school samples have a surface coating of zinc oxide, making reactions unpredictable.
Zinc oxide or carbonate	LOW HAZARD	The zinc oxide fumes ('philosopher's wool') formed when zinc dust burns in air are regarded as hazardous dust.
Zinc salts Solid or concentrated solutions Sulfate(VI) if 1.5 M or more Chloride & bromide if 1 M or more	CORR. HARM.	DANGER: Harmful if swallowed (especially saturated solutions for crystal-growing); causes serious eye damage (sulfate); causes severe skin burns and eye damage (chloride & bromide); toxic to aquatic life with long-lasting effects. When preparing zinc sulfate by reacting zinc and sulfuric acid, the reaction can be slow and is often incomplete.
Zinc salts Most solutions Sulfate(VI) if less than 1.5 M but 0.2 M or more Chloride if less than 1M but 0.2 M or more Bromide if less than 1 M but 0.1 M or more	CORR. IRRIT.	DANGER: corrosive to eyes (all) and to skin (chloride & bromide); respiratory irritant (chloride if more than 0.4 M, bromide if more than 0.2 M).
Zinc salts Dilute solutions Sulfate(VI) if less than 0.2 M but 0.06 M or more Chloride if less than 0.2 M but 0.1 M or more Bromide if less than 0.1 M but 0.05 M or more	IRRITANT	WARNING: irritating to eyes (all) and skin (chloride & bromide).
Zinc salts Very dilute solutions Sulfate(VI) if less than 0.06 M Chloride if less than 0.1 M Bromide if less than 0.05 M	LOW HAZARD	-

Typical control measures to reduce risk

- Use the lowest possible quantities and concentrations.
- Only electrolyse zinc chloride/bromide solutions briefly, unless in a fume cupboard (essential for molten compounds).
- Assume zinc powder/dust is fresh and not partially oxidised on the surface.
- When reacting zinc and acid, check no acid remains before evaporating solutions (pH should be 4 or higher).
- Wear eye protection.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong?
 - eg, solutions spurting out of test tubes when heated or solutions heated to dryness and decomposing.
- How serious would it be if something did go wrong?
 - eg, are there hazardous reaction products (such as chlorine from the electrolysis of zinc chloride)?
- How can the risk(s) be controlled for this activity?
 - eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

Emergency action

In the eye Flood the eye with gently-running tap water for at least 10 minutes. Consult a medic. **Swallowed** Do no more than wash out the mouth with water. Do not induce vomiting. Consult a

- Dust breathed in Remove the casualty to fresh air. Consult a medic if breathing is difficult.
- Spilt on the skin or clothing Remove contaminated clothing and rinse it. Wash off the skin with plenty of water.
- Spilt on floor, bench, etc Scoop up solid (take care not to raise dust). Wipe up small solution spills or any traces of
 - solid with cloth; for larger spills use mineral absorbent (eg, cat litter).



Hydrogen

Substance	Hazard	Comment
Hydrogen <i>Gas</i>	FLAMMABLE	Danger: extremely flammable gas; contains gas under pressure, may explode if heated (cylinders & canisters). It forms explosive mixtures with air and oxygen. Mixtures with air between 4% and 74% hydrogen by volume are explosive. Explosive mixtures will ignite below 500 °C and well below this temperature in the presence of catalysts such as transition metals and their oxides. The explosion with oxygen produces a very loud noise which can damage hearing.
	GAS CYLINDER	Mixtures of hydrogen and oxygen can arise when recharging a car battery (or model cells in schools); ensure good ventilation, avoid sparks and naked flames.

Typical control measures to reduce risk

- If preparing the gas in test-tube reactions, use the smallest possible amounts.
- Wear eye protection and stand well back.
- Use safety screens for all but test-tube amounts of the gas; ensure good laboratory ventilation.
- If preparing the gas on anything larger than a test-tube scale, make sure the apparatus has the smallest possible volume, so that only a little air has to be flushed out. If lighting the gas at a jet, test a sample to make sure that all the air has been flushed out and light from a distance.
- If possible, use a gas cylinder rather than generate your own hydrogen, because the cylinder produces a more-rapid flow which flushes air more quickly from the apparatus.
- When reducing, eg, metal oxides, consider alternative reducing agents such as methane or ammonia gas.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong? eg, could the hydrogen be ignited accidentally? How easy is it to make sure that all the air has been flushed out?
- How serious would it be if something did go wrong?
 - NB There are occasional reports of pupils being taken to hospital (for treatment to cuts or for splashes of chemicals) as a result of damage to apparatus in hydrogen explosions.
- How can the risk(s) be controlled for this activity? eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

Emergency action

Gas escape Open all the windows. Make sure there are no naked flames.

If there are cuts from flying glass, apply pressure using a clean pad of cloth. Do not attempt to **Explosion** remove large pieces of embedded glass, etc. If there is arterial bleeding, the casualty should be laid down and the injured limb raised up. Consult a medic.

Oxygen and ozone

Substance	Hazard	Comment
Oxygen (O ₂) Gas		DANGER: may cause or intensify fire; contains gas under pressure, may explode if heated (cylinders).
	(Q)	Air contains about 21% oxygen.
	OXIDISER	Combustible substances burn much more fiercely in air which has been only slightly enriched with oxygen (eg, 25%). Products of combustion are often strongly acidic or basic (alkaline) oxides.
	GAS CYLINDER	If the amount of oxygen in the air becomes too low, headaches, unconsciousness and death may result. First effects may be noticed when the percentage drops to 18%. Similar effects may be observed on high mountains due to lower pressure.
Ozone (O ₃) Gas	OXIDISER TOXIC IRRITANT	DANGER: may cause or intensify fire; causes serious eye irritation; fatal if inhaled. For a 15-minute exposure, the concentration in the atmosphere should not exceed 0.4 mg m ⁻³ . It is not normally made or used in school science. In the presence of sunlight, traces of hydrocarbons in the air react with nitrogen oxides (see <i>CLEAPSS Student Safety Sheet 53</i>), eg, from car exhausts, to form ozone. This causes photochemical smog in certain hot weather conditions. Small amounts of ozone are also formed in some photocopiers but this is only likely to be a problem in a small room with poor ventilation. Although dangerous if breathed in, ozone in the upper atmosphere performs a very important safety role, where it absorbs much of the ultraviolet radiation reaching the earth, thus preventing dangerous exposures (see <i>CLEAPSS Student Safety Sheet 12</i>). There is considerable concern that pollution by certain chlorinated hydrocarbons (see <i>CLEAPSS Student Safety Sheet 62</i>) is destroying the ozone layer.

Typical control measures to reduce risk

- Wear eye protection when preparing oxygen or burning substances in oxygen.
- Avoid looking directly at the very bright light from magnesium burning in oxygen, eg by using welding filters or averting eyes.
- Avoid inhaling products when non-metals or metals are burning in oxygen.
- Use safety screens when burning substances in oxygen on anything larger than a test-tube scale.
- If using cylinders of oxygen, do not lubricate controls with oil or grease as this might catch fire.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong? eg, could substances burn much more fiercely than expected?
- How serious would it be if something did go wrong? eg, would there be widespread health effects if the ozone layer is damaged by pollution?
- How can the risk(s) be controlled for this activity?

eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

Emergency action

Gas escape Open all the windows. Extinguish all naked flames. Smother flames on clothing or the skin with a fire blanket or other material. Cool any burnt skin with Clothing catches fire gently-running tap water for 10 minutes. Consult a medic if the area of burn is larger than a small coin. Other fires Allow fires in sinks etc to burn out. Fires at the top of test tubes, beakers, etc should be smothered with a damp cloth or heat-resistant mat.

Sulfur dioxide

Substance	Hazard	Comment
Sulfur dioxide Gas	CORROSIVE	DANGER: causes severe skin burns and eye damage; toxic if inhaled. Effects of exposure by inhalation may not be immediately apparent and can develop or increase over time. Inhalation by those with known breathing difficulties, eg asthma, may exacerbate such pre-existing conditions. Adverse health effects may be apparent even at very low levels, ~ 0.3 mg m ⁻³ (0.1 ppm). Previously, the HSE has said for a 15-minute exposure, the concentration of the gas in the atmosphere should not exceed 2.7 mg m ⁻³ . It is produced naturally in large amounts by volcanoes. Most fossil fuels contain traces of sulfur compounds. When burnt, these result in sulfur dioxide released into the atmosphere, causing acid rain. It may be produced in the laboratory by the action of heat or dilute acid on sulfites or thiosulfates. It is used in small amounts as a preservative in some foodstuffs and wines. Approved food additive, E220.
Sulfur dioxide Dilute solution in water	LOW HAZARD (but beware of TOXIC gas given off)	The gas is very soluble in water and may cause suck back. Alternatively, a sulfur dioxide solution can be prepared using sodium metabisulfite and adding sulfuric acid. The gas escapes easily from solution, especially if this is warmed, and should not be inhaled.

Typical control measures to reduce risk

- If preparing the gas in test-tube reactions, use the smallest amounts possible.
- Wear eye protection.
- Take steps to prevent suck back of water, eg, by the use of Bunsen valves.
- Use a fume cupboard for anything larger than test-tube amounts of gas; ensure good laboratory ventilation and quickly dispose of solutions containing sulfur dioxide.
- If testing for the gas by its smell, follow the safe technique for sniffing gases: use your hand to waft the gas towards vour nose.
- Do **not** expose asthmatics to the gas.
- Use fuels which are naturally low in sulfur, remove sulfur compounds before use or use scrubbers to absorb sulfur dioxide from the exhaust gases of coal- or oil-burning power stations.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong? eg, a leak of gas into the laboratory from apparatus or a warmed solution, or suck back.
- How serious would it be if something did go wrong?
- How can the risk(s) be controlled for this activity? eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

Emergency action

Flood the eye with gently-running tap water for 10 minutes. Consult a medic. In the eve Remove the casualty to fresh air. Consult a medic if breathing is difficult. Vapour breathed in Open all windows. If over 1 litre of gas is released, evacuate the laboratory. Gas escape in a laboratory

Nitrogen oxides

includes Nitrogen monoxide, Nitrogen dioxide, Dinitrogen tetroxide & Dinitrogen oxide

Substance	Hazard		Comment
Nitrogen monoxide (Nitric oxide; NO) Gas	OXIDISING	CORROSIVE	DANGER: May cause or intensify fire. Causes severe skin burns and eye damage; fatal if inhaled; may cause respiratory irritation; corrosive to the respiratory tract. Effects of exposure by inhalation may or may not be immediately apparent and can develop or increase over time. Inhalation by those with known breathing difficulties (eg, asthma) may
	то	OXIC	exacerbate such pre-existing conditions. In the past, HSE has said for 15-minute exposure, the concentration in the atmosphere should not exceed 1.4 mg m ⁻³ . It reacts with oxygen in the atmosphere to form nitrogen dioxide (see below).
			It may be formed by the reaction between oxygen and nitrogen in the air, especially in car engines. This is a major contributor to acid rain and photochemical smog. The mixture of NO and NO_2 formed in this way is often referred to as NO_X .
Nitrogen dioxide (NO₂), Dinitrogen tetroxide (N₂O₄) Gases	OXIDISING	CORROSIVE	DANGER: May cause or intensify fire. Causes severe skin burns and eye damage; fatal if inhaled; may cause respiratory irritation. Effects of exposure by inhalation may or may not be immediately apparent and can develop or increase over time. Inhalation by those with known breathing difficulties (eg, asthma) may exacerbate such pre-existing conditions. In the past, HSE has said for 15-minute exposure, the
	тс	OXIC	concentration in the atmosphere should not exceed 1.9 mg m ⁻³ . They are formed as air pollutants from nitrogen monoxide (see above). They are formed in the laboratory by the action of heat on many nitrates and by the reaction of nitric acid on some metals. They are very soluble in water; there is a risk of suck back.
Dinitrogen oxide (Nitrous oxide, N₂O) 'Laughing gas'	OXID	DISING	DANGER: May cause or intensify fire. For 15-minute exposure, the concentration in the atmosphere should not exceed 549 mg m ⁻³ . This is an anaesthetic in large amounts. It has been used as a general anaesthetic, eg, by dentists. An approved food additive, E942, where it is used as a propellant and foaming agent, eg for cream.

Typical control measures to reduce risk

- If preparing the gas in test-tube reactions, use the smallest amounts possible and take steps to avoid suck-back (eg, a Bunsen valve).
- Wear eye protection.
- Use a fume cupboard for anything larger than test-tube amounts of gas; ensure good laboratory ventilation.
- If testing for the gas by its smell, follow the safe technique for sniffing gases: use your hand to waft the gas towards your nose.
- Do **not** expose asthmatics to the gas.
- Use catalytic converters in car exhausts to reduce the amount of nitrogen oxides released into the air.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong? eg, a leak of a gas from apparatus into the laboratory atmosphere.
- How serious would it be if something did go wrong?
- How can the risk(s) be controlled for this activity? eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

Emergency action

Flood the eye with gently-running tap water for 10 minutes. Consult a medic.. In the eye Vapour breathed in Remove the casualty to fresh air. Consult a medic if breathing is difficult. Open all windows. If over 1 litre of gas is released, evacuate the laboratory. Gas escape in a laboratory

Chlorine includes Chlorine water

Substance	Hazard	Comment
Chlorine Gas OXIDISING TOXIC ENVIRONMENTAL HAZARE		DANGER: may cause or intensify fire; causes skin and serious eye irritation; toxic if inhaled; may cause respiratory irritation; very toxic to aquatic organisms. Effects of exposure by inhalation may or may not be immediately apparent and can develop or increase over time. Inhalation by those with known breathing difficulties (eg, asthma) may exacerbate such pre-existing conditions. For a 15-minute exposure, the concentration of the gas in the atmosphere should not exceed 1.5 mg m ⁻³ . It is used to kill microbes in public water supplies, at a concentration between about 0.1 and 1.0 mg per litre. It is also used to treat swimming pool water at a concentration between about 1.4 and 4.0 mg per litre. It may be formed in the laboratory by electrolysis and the oxidation of some chlorides. It may be formed in the laboratory, in the home or at work by the
Chlorine water Solution in water	LOW HAZARD (but beware of TOXIC gas given off)	action of acid on bleaches; see <i>CLEAPSS Student Safety Sheet 41</i> . It can be made in a fume cupboard by diluting a saturated solution with a little more than its own volume of water. Chlorine gas escapes easily from the solution, especially if it is warmed. For a 15-minute exposure, the concentration of chlorine gas in the atmosphere should not exceed 1.5 mg m ⁻³ and this could easily be approached in localised situations, eg just above open test tubes or bottles.

Typical control measures to reduce risk

- If preparing the gas in test-tube reactions, use the smallest possible amounts; where possible, absorb excess gas with a soda lime tube.
- Wear eye protection.
- Use a fume cupboard for anything larger than test-tube amounts of gas; ensure good laboratory ventilation.
- If testing for the gas by its smell, follow the safe technique for sniffing gases: use your hand to waft the gas towards
- Do not expose asthmatics to the gas; even with chlorine water, take care not to breathe in chlorine.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong?
- How serious would it be if something did go wrong?
- NB There are occasional reports of pupils being taken to hospital as a result of breathing in chlorine.
- How can the risk(s) be controlled for this activity?
- eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

Emergency action

In the eye Flood the eye with gently-running tap water for 10 minutes. Consult a medic.

Vapour breathed in Remove the casualty to fresh air. Consult a medic if breathing is even slightly affected.

Do no more than wash out the mouth with water. Do not induce vomiting. Consult a **Swallowed**

medic.

Spilt on the skin or clothing For chlorine water, remove contaminated clothing and rinse it. Then drench the skin with

plenty of water.

Spilt on the floor, bench, Open all windows. For a release of more than 1 litre of chlorine gas, evacuate the

laboratory. Mop up chlorine water and rinse with plenty of water.

Open all windows. If over 1 litre of gas is released, evacuate the laboratory. Gas escape in a laboratory

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Bromine *includes* **Bromine** water

Substance	Hazard	Comment
Bromine Liquid		DANGER: fatal if inhaled; causes severe skin burns and eye damage; very toxic to aquatic organisms.
Liquia		Effects of exposure by inhalation can increase over time. Inhalation by
	TOXIC CORROSIVE	
	TOXIC CORROSIVE	such pre-existing conditions.
	N. N	For a 15-minute exposure, the concentration of the vapour in the
	\\\\	atmosphere should not exceed 1.3 mg m ⁻³ .
		Keep the following next to liquid bromine at all times: At least 500 cm ³
	ENVIRONMENTAL HAZARD	of 1 M sodium carbonate and also a container of solid hydrated sodium
		carbonate (washing soda) for treating spills on hard surfaces.
Moderately		DANGER: causes severe eye damage; irritating to skin.
concentrated		Bromine rapidly diffuses out of solution, producing a vapour which is
	^	very toxic if breathed in. Effects of exposure by inhalation can increase
bromine solution,		over time. Inhalation by those with known breathing difficulties (eg,
in water or organic		asthma) may exacerbate such pre-existing conditions.
solvents	CORROSIVE	A saturated solution in water is about 0.25 M.
(if 0.2M or more)		There may also be hazards associated with the organic solvent; see
		relevant CLEAPSS Student Safety Sheets.
Moderately dilute		WARNING: irritating to eyes and skin.
bromine solution,	A	Bromine rapidly diffuses out of solution, producing a vapour which is
in water or organic		very toxic if breathed in. Effects of exposure by inhalation can increase
solvents		over time. Inhalation by those with known breathing difficulties (eg,
(if 0.06 M or more but		asthma) may exacerbate such pre-existing conditions.
less than 0.2 M)	IRRITANT	There may be hazards associated with the organic solvent; see relevant
1033 (11411 012 111)		CLEAPSS Student Safety Sheets
Very dilute bromine		There may be hazards associated with the organic solvent; see relevant
solution,	LOWILLAZADE	CLEAPSS Student Safety Sheets.
in water or organic	LOW HAZARD	0.002M is suitable for testing alkenes for unsaturation and for
solvents		halogen/halide displacement reactions.
(if less than 0.06 M)		

Typical control measures to reduce risk

- Use the lowest concentration and smallest volume possible.
- Wear eye protection and protective nitrile gloves for all but the most-dilute solutions.
- Avoid breathing the fumes from concentrated solutions, eg, by the use of a fume cupboard.
- When bromine liquid is in use, have plenty of 1M sodium carbonate solution available to deal with spills.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong?
- How serious would it be if something did go wrong?
- How can the risk(s) be controlled for this activity?

eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

•	In the eye	Flood the eye with gently-running tap water for 10 minutes. Consult a medic.
•	Vapour breathed in	Remove the casualty to fresh air. Consult a medic if breathing is even slightly affected.
•	Swallowed	Do no more than wash out the mouth with water. Do not induce vomiting. Consult a medic.
•	Spilt on the skin or clothing	Remove contaminated clothing, and then drench the skin with plenty of water. Consult a medic.
•	Spilt on the floor, bench,	For spills of all but a few drops of liquid bromine, open windows and evacuate the
	etc	laboratory. For small bromine spills, add solid hydrated sodium carbonate (or 1M
		sodium carbonate solution) and leave for 1 hour. Mop up and rinse with plenty of water.

Iodine includes Iodine solutions

Substance	Hazard	Comment
lodine Solid lodine solution, in potassium iodide / water or in organic solvent (if 1 M or more)	HARMFUL ENVIRONMENT HARMFUL ENVIRONMENT	Warning: harmful in contact with skin and if inhaled. Very toxic to aquatic life. It is easily vapourised if heated - the violet vapour is dangerous to the eyes. For a 15-minute exposure, the concentration in the atmosphere should not exceed 1.1 mg m ⁻³ . Warning: harmful in contact with skin and if inhaled. Very toxic to aquatic life. Even dilute solutions will stain the skin. There may be hazards associated with the organic solvent; see relevant CLEAPSS Student Safety Sheets. Ethanol is often used; it is HIGHLY FLAMMABLE.
Dilute iodine solution, in potassium iodide / water or in organic solvent (if less than 1 M)	LOW HAZARD	Even dilute solutions will stain the skin. Solutions 0.01 to 0.1 M are suitable for many activities, eg testing for starch, work with enzymes, etc. There may be hazards associated with the organic solvent; see relevant CLEAPSS Student Safety Sheets. Ethanol is often used; it is HIGHLY FLAMMABLE. 'Tincture of iodine', used as a mild antiseptic, is a dilute solution in ethanol (about 0.1 M).

Typical control measures to reduce risk

- Use the lowest concentration and smallest volume possible.
- Wear eye protection for all but the most-dilute solutions.
- Handle iodine solid using forceps or wear protective gloves especially for larger amounts.
- Avoid breathing iodine vapour, eg, by using a fume cupboard.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong?
- How serious would it be if something did go wrong?
 - NB Iodine can crystallise painfully on the eyeball.
- How can the risk(s) be controlled for this activity? eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

•	In the eye	Flood the eye with gently-running tap water for 10 minutes. Consult a medic.
•	Vapour breathed in	Remove the casualty to fresh air. Consult a medic if breathing is even slightly affected.
•	Swallowed	Do no more than wash out the mouth with water. Do not induce vomiting. Consult a medic.
•	Spilt on the skin or clothing	Brush off solid iodine and immerse in sodium thiosulfate solution (20%, 1 M). Remove contaminated clothing, soak it and drench the skin with plenty of water. Consult a medic if a large area is affected or blistering occurs.
•	Spilt on the floor, bench, etc	Scoop up any solid iodine, add sodium thiosulfate solution (20%, 1 M) to the remaining spill and leave for 1 hour. Mop up and rinse with plenty of water.

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Hydrogen peroxide

Substance	Hazard	Comment
Concentrated hydrogen peroxide solution [If less than 8.3M, ie, '100 volume strength' (30%) but 2.3 M or more, ie, over '28 volume strength' (8% w/v)]	CORROSIVE	DANGER: causes serious eye damage and may be harmful if swallowed. This is the most-concentrated solution found in schools. Take care to avoid skin contact. It decomposes slowly to produce oxygen gas (see <i>CLEAPSS Student Safety Sheet 51</i>); pressure may build up and care needs to be taken when opening a bottle. It should be stored in the dark. Decomposition is speeded up by catalysts such as some metal oxides and some enzymes. The oxygen formed will assist fires.
Dilute hydrogen peroxide solution [If less than 2.3 M but 1.5 M or more, ie, less than '28 volume strength' (8%) but '18 volume strength' (5% w/v) or more]	IRRITANT	Warning: irritating to the eyes (and to the skin, although not officially classified as such). The typical concentration used in school science practical work. It decomposes slowly to produce oxygen gas (see CLEAPSS Student Safety Sheet 51); pressure may build up and care needs to be taken when opening a bottle. It should be stored in the dark. Decomposition is speeded up by catalysts such as some metal oxides and some enzymes.
Very dilute hydrogen peroxide solution [If less than 1.5 M, ie, less than '18 volume strength' (5% w/v)]	LOW HAZARD	It is used for bleaching hair. It decomposes slowly to produce oxygen gas (see CLEAPSS Student Safety Sheet 51); pressure may build up and care needs to be taken when opening a bottle. It should be stored in the dark. Decomposition is speeded up by catalysts such as some metal oxides and some enzymes. Old stock may have insufficient peroxide molecules for the intended activity.

Additional information: Concentration of hydrogen peroxide solutions may be expressed in several different ways:

- Molarity: as with any chemical, the concentration may be given as x mol dm⁻³, or x M, ie x mol in 1 dm³ of solution
- Percentage, w/v: a y % w/v solution will contain y g hydrogen peroxide in 100 cm³ of solution
- Volume strength: 1 cm³ of z volume strength will give z cm³ of oxygen when it decomposes.

Typical control measures to reduce risk

- Use the lowest concentration and smallest volume possible.
- Wear eye protection for all but the most-dilute solutions.
- Store concentrated solutions away from heat and light, in bottles with special vented caps. Beware of a rapid release of pressure when opening a bottle.
- Avoid accidental contamination of solutions which may speed up the formation of oxygen and pressure build-up.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong?
- eg, could an impurity / catalyst cause rapid decomposition and frothing?
- How serious would it be if something did go wrong?
 - eg, if the solution splashes onto the skin, is it sufficiently concentrated to cause burns?
- How can the risk(s) be controlled for this activity?
- eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

•	In the eye	Flood the eye with gently-running tap water for 10 minutes. Consult a medic.
•	Swallowed	Do no more than wash out the mouth with water. Do not induce vomiting. Consult a medic.
•	Spilt on the skin or clothing	Flood the area with plenty of water. Remove contaminated clothing and soak it. If a large area is affected or blistering occurs, consult a medic.
•	Spilt on the floor, bench, etc	For large spills, and especially for (moderately) concentrated solutions, cover with mineral absorbent (eg, cat litter) and scoop into a bucket. Dilute with at least ten times its own volume of water. Rinse the floor etc with plenty of water. Wipe up small amounts with a damp cloth and rinse it well.

Carbon & its oxides

including carbon dioxide, carbon monoxide

Substance	Hazard	Comment
Carbon dioxide		WARNING (if supplied in cylinder) may explode if heated. Otherwise, not classed as
Gas		hazardous but it can cause asphyxiation if the proportion of carbon dioxide in the air
		becomes too high, eg, as a result of the rapid evaporation of the solid in a confined
	COMPRESSED	space or, in some African lakes, it is released from decaying organic matter. Because
	GAS	it is denser than air, it may build up in low areas, eg, in caves. For a 15-minute
		exposure, the concentration in the atmosphere should not exceed 27 000 mg m ⁻³ .
	otherwise	About 0.04% is present in normal air, as compared with about 0.03% fifty years ago.
		This increase is a result of burning fossil fuels in motor vehicles, power stations, etc.
	LOW HAZARD	This in turn is believed by almost all scientists to be contributing to a very gradual rise
		in the temperature of the Earth (global warming) as a result of the greenhouse effect.
Carbon dioxide		It causes frostbite (burns) and needs careful handling.
Solid	\ ``\\	If it evaporates rapidly in a closed vessel, it may cause an explosion or, in a confined
'dry ice'	COLD	space, it may cause asphyxiation as the air is forced out.
Carbon monoxide	^	DANGER: extremely flammable gas; toxic if inhaled; may damage the unborn child;
Gas	(M)	causes damage to organs through prolonged or repeated inhalation. As little as 0.01%
		can cause headaches. The gas has no taste or smell and is not trapped by the filters in
	FLAMMABLE	filter fume cupboards
		It is often formed when hydrocarbon fuels burn in a limited supply of air, eg, car
		engines especially in confined spaces, or gas-powered water heaters with poor
	TOXIC	ventilation. Every year, this causes many deaths in the home. Traces also occur in
		cigarette smoke and are implicated in heart and artery diseases. It also contributes to
		the greenhouse effect.
	HEALTH	For a 15-minute exposure, the concentration in the atmosphere should not exceed
	HAZARD	232 mg m ⁻³ . It forms explosive mixtures with air and oxygen. Mixtures with air
		between 12% and 74% carbon monoxide by volume are explosive.
Carbon		Applies to lampblack, charcoal, activated carbon, decolourising charcoal. Soot is also
Graphite, diamond,		mainly carbon but may be contaminated with carcinogenic chemicals. This was a
buckminsterfullerene	LOW HAZARD	cause of cancer amongst chimney sweeps in Victorian times. The hazards of
		buckminsterfullerene nanotubes are not fully known yet. Hot charcoal (carbon
Tamical control		blocks), even if not glowing red, can slowly combust and stay hot for many hours.

Typical control measures to reduce risk

- Wear protective thermal gloves or use tongs for handling solid carbon dioxide.
- Use energy-efficient vehicles and power stations and/or non-fossil fuels or wind or solar power to limit the amount of carbon dioxide or monoxide emitted.
- Use a ducted fume cupboard (not a recirculatory filter fume cupboard) for handling carbon monoxide.
- Have gas appliances serviced regularly and consider installing a carbon monoxide detector,
- Use catalytic converters in car exhausts to reduce the amount of carbon monoxide released into the air.
- After use, allow hot charcoal blocks to cool in air; store them in air-tight metal containers.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong?
- How serious would it be if something did go wrong?
 - eg, global warming causing rising sea level and resultant flooding of low-lying areas.
- How can the risk(s) be controlled for this activity?
 - eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

- Solid in the eye Flood eye with gently-running tap water for 10 minutes. Consult a medic if solid carbon dioxide. Brush off quickly and immerse the affected area in cold water. If there are any signs of burning Solid on the skin from solid carbon dioxide consult a medic. or clothing Vapour breathed For carbon monoxide, or carbon dioxide in larger quantities, remove the casualty to fresh air. Consult a medic if carbon monoxide was inhaled or breathing is difficult.
- Open all windows. For large amounts of carbon monoxide, evacuate the laboratory. Gas escape in lab.



Hydrogen sulfide & other sulfides

including carbon disulfide, ammonium sulfide and most metal sulfides

DANGER: extremely flammable gas; fatal if inhaled; very toxic to aquatic organisms. Mixtures of hydrogen sulfide with air containing between 4% and 45% hydrogen sulfide atmosphere should not exceed 14 mg m³. It has a very strong smell of rotten eggs and the human nose can detect as little as 0.01 mg m³. At higher concentrations it anaesthetises (deadens) the sense of smell and so the danger may not be realised. It may sometimes be found in coal mines (stinkdamp) and is produced by rotting seawed and is the commonest cause of death in sewer workers. Hydrogen sulfide Solution in water DANGER: hydrogen sulfide gas is acutely toxic by inhalation and diffuses readily from solutions. However, use of solutions of hydrogen sulfide is safer than using the gas. DANGER: highly flammable liquid & vapour; causes skin and serious eye irritation; suspected of damaging fertility and the unborn child; causes damage to organs through prolonged or repeated exposure. For a 15-minute exposure, the concentration of the gas in the atmosphere should not exceed 45 mg m³; flash point -30 °C. Very volatile. The demonstration involving the reaction with nitrogen monoxide (the 'barking dog') is very dangerous, especially if the oxide is not pure. DANGER (if more than 20%, "3M): flammable liquid & vapour; causes severe skin burns & eye damage; contact with acids liberates toxic gas; very toxic to aquatic organisms. DANGER (if more than 0.4 M but less than ~3M): causes severe skin burns & eye damage; contact with acids liberates toxic gas. Varning under than 0.1 M but less than 0.4M): harmful if swallowed; causes severe skin burns & eye damage; contact with acid liberates toxic gas; very toxic to aquatic organisms. DANGER (solution if more than 0.4M): harmful if swallowed; causes severe skin burns & eye damage; contact with acid liberates toxic gas. Warning (solution if more than 0.4M): harmful if swallowed; causes severe skin burns & eye damage; contact with acid liberates toxic gas. Warning (solution if more than 0.4M): harmful if swa	Substance	Hazard	Comment
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		ENVIRON. HA	the atmosphere, producing hydrogen samue.
sulfides Low HAZARD zinc sulfide. Dilute acid reacts to produce hydrogen sulfide. If heated strongly in air,	Most metal		
	sulfides	LOW HAZAR	· · · · · · · · · · · · · · · · · · ·
Insoluble solids may produce toxic sulfur dioxide (see CLEAPSS Student Safety Sheet 52).	Insoluble solids		may produce toxic sulfur dioxide (see CLEAPSS Student Safety Sheet 52).

Typical control measures to reduce risk

- If producing the gas, use the smallest amounts possible, eg no more than a few drops of acid on excess sulfide.
- Wear eye protection.
- Use a fume cupboard if adding more than a few drops of dilute acid to excess sulfide; ensure good laboratory ventilation and quickly dispose of solutions containing hydrogen sulfide.
- If smelling the gas, follow the safe technique: use your hand to waft the gas towards your nose.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong?
 eg, a leak of gas into the laboratory from apparatus or a warmed solution.
- How serious would it be if something did go wrong?
- How can the risk(s) be controlled for this activity?
 eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

- In the eye Flood the eye with gently-running tap water for 10 minutes. Consult a medic.
- Vapour breathed in Remove the casualty to fresh air. Consult a medic if breathing is difficult.
- Gas escape in lab. Open all windows. If over 1 litre of gas is released, evacuate the laboratory.



Ethanol

also applies to Denatured alcohol and Methylated spirit

Substance	Hazard	Comment
Ethanol		DANGER: highly flammable liquid & vapour.
Liquid		There is a serious risk of liquid catching fire; its vapour may catch fire
pure		above 13 °C. The vapour/air mixture is explosive (from 3.3 to 19%
	E3	ethanol). The concentration in the air should not exceed 5760 mgm ⁻³ .
	FLAMMABLE	Evaporation/cooling experiments: Alcohol-based hand gels may be used
		to experience the cooling effect of evaporation. Apply only a small
		quantity to the back of a hand. Do not use surgical spirits.
Industrial denatured		DANGER: highly flammable liquid & vapour; harmful if swallowed; may
alcohol (IDA)	^ ^	cause damage to organs. This is ethanol, containing 5% methanol (v/v). It
formerly		is often used in schools in place of pure ethanol (because it is cheaper)
Industrial		and usually labelled just as 'Ethanol' but it is more hazardous than pure
methylated spirit	FLAMM. HARM. HEALTH	ethanol because of the methanol. Commonly used as a solvent, eg, for
(IMS)		chlorophyll, for indicators (universal indicator, phenolphthalein).
Completely	^ ^	It contains methanol, propan-2-ol, methyl ethyl ketone, a bitter-tasting
denatured alcohol		compound and, sometimes, a purple dye. CDA is not suitable for use
(CDA)		indoors.
(CDA)	FLAMM. HARM. HEALTH	
Surgical spirit		It is prescribed for medical purposes, eg, foot infections. It must not be
Ethanol, with small		swallowed. Use only as prescribed. Do not use for skin evaporation
amounts of castor oil,		experiments.
methyl salicylate and	FLAMM. CORROS. ENVIR.	
diethyl phthalate		
Ethanol		Alcoholic drinks contain ethanol, typically 3 to 7% (v/v) (beers), 11 to 14%
Dilute solution in water		(v/v) (wines), 30 to 40% (v/v) (spirits).
	LOW HAZARD	Although chemical hazards are low, there may be considerable effects on
		the body leading to a loss of judgement, slower reaction times, etc.
		Consumption is dangerous if driving a vehicle or operating machinery.

Typical control measures to reduce risk

- Use the smallest volume possible; wear eye protection.
- Make sure the room is well ventilated.
- Check that equipment for extinguishing fires is nearby, eg, damp cloth, bench mat, fire blanket.
- Do not use near naked flames; if heating necessary, use an electrically-heated water bath or hot water from kettle.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong?
 - eg, does ethanol need to be heated? Could quantities of the vapour be breathed in? Might there be fooling around?
- How serious would it be if something did go wrong?
 - NB Some of the most serious accidents in school science have involved ethanol fires, including clothing fires and badly-burnt skin.
- How can the risk(s) be controlled for this activity?
- eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

- In the eye
 Flood the eye with gently-running tap water for 10 minutes. Consult a medic.
- **Swallowed** Do no more than wash out the mouth with water. Do **not** induce vomiting. Consult a medic. NB:
 - casualty may show signs of drunkenness.
- **Spilt on the skin** Remove contaminated clothing and rinse it. Wash the affected area and clothing with plenty of **or clothing** water.
- Clothing catches fire
 Smother flames on clothing or the skin with a fire blanket or other material. Cool any burnt skin with gently-running tap water for 10 minutes.
- Other ethanol Allow fires in sinks, etc to burn out. Fires at the top of test tubes, beakers, etc should be smothered with a damp cloth or heat-resistant mat.
- **Spilt on the** Extinguish all Bunsen flames. Wipe up small amounts with a cloth and rinse well. Open windows for **floor, bench, etc** larger amounts, cover with mineral absorbent (eg, cat litter), scoop into a bucket and add water.



Propanone

also known as Acetone

Substance

Hazard

Comment

Propanone (acetone)

Liquid

Used as nail polish (varnish) remover [but ethyl ethanoate (ethyl acetate) is more usual].





DANGER: highly flammable liquid & vapour. Causes serious eye irritation; may cause drowsiness or dizziness; repeated exposure may cause skin dryness and cracking

Its vapour may catch fire above -20°C.

For a 15-minute exposure, the concentration in the atmosphere should not exceed 3620 mg m⁻³. The smell can be detected by most people at about 47 mg m⁻³, well below the level which could cause harm.

Typical control measures to reduce risk

- Wear eye protection.
- Make sure the room is well ventilated or, in a laboratory, use a fume cupboard if possible.
- Check ways of putting out any fires.
- Do not use near naked flames; if heating is necessary, use an electrically-heated water bath or hot water from a kettle.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong? eg, does propanone need to be heated? Could there be high levels of vapour, perhaps as a result of chromatograms drying?
- How serious would it be if something did go wrong?
- How can the risk(s) be controlled for this activity?

eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

Emergency action

	9	
•	In the eye	Flood the eye with gently-running tap water for 10 minutes. Consult a medic.
•	Vapour breathed in	Remove the casualty to fresh air. Keep him/her warm. Consult a medic if breathing is difficult.
•	Swallowed	Do no more than wash out the mouth with water. Do not induce vomiting. Consult a medic.
•	Spilt on the skin or clothing	Remove contaminated clothing. If more than a test-tube amount was involved, wash the affected area and clothing with plenty of water.
•	Clothing catches fire	Smother flames on clothing or the skin with a fire blanket or other material. Cool any burnt skin with gently-running tap water for 10 minutes.
•	Other propanone fires	Allow fires in sinks, etc to burn out. Fires at the top of test tubes, beakers, etc should be smothered with a damp cloth or heat-resistant mat.
•	Spilt on the floor, bench, etc	Extinguish all Bunsen flames. Wipe up small amounts with a cloth and rinse well. Open windows for larger amounts, cover with mineral absorbent (eg, cat litter),

scoop into a bucket and add water.

Chlorinated hydrocarbons

Substance	Hazard	Comment
Dichloromethane	^ ^	DANGER: suspected of causing cancer; causes skin & serious eye irritation;
(Methylene		may cause respiratory irritation, drowsiness or dizziness and damage to
dichloride)	\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	organs through prolonged or repeated exposure. It is used in some paint
liquid	HEALTH HAZ. IRRITANT	strippers. For a 15-minute exposure, the concentration in the atmosphere
	HEALIH HAZ. IKKITANI	should not exceed 1060 mg m ⁻³ .
Trichloromethane		DANGER: harmful if swallowed; causes skin & serious eye irritation; toxic if
(Chloroform)		inhaled; may cause drowsiness or dizziness; suspected of causing cancer and
liquid		of damaging the unborn child; causes damage to organs through prolonged
•		or repeated exposure. For a 15-minute exposure, the concentration in the
	TOXIC HEALTH HAZ.	atmosphere should not exceed 30 mg m ⁻³ .
		Used in the past as an anaesthetic.
Tetrachloro-		DANGER: toxic if swallowed, inhaled or in contact with skin; may cause allergic
methane	^ ^	skin reaction; suspected of causing cancer; causes damage to organs through
(Carbon		prolonged or repeated exposure; harmful to aquatic life with long-lasting
tetrachloride)		effects; harms public health & the environment by destroying ozone in the
liquid	TOXIC HEALTH HAZ.	upper atmosphere. For a 15-minute exposure, the concentration in the
7	TOXIO TIEMETITIME	atmosphere should not exceed 39 mg m ⁻³ . It can no longer be legally bought
		and existing stocks should not be used in work with open test tubes.
1,1,1-		WARNING: harmful if inhaled; causes skin and serious eye irritation; harms
trichloroethane		public health & the environment by destroying ozone in the upper
(Methyl		atmosphere. For a 15-minute exposure, the concentration in the
chloroform)	HARMFUL	atmosphere should not exceed 1110 mg m ⁻³ . It can no longer be legally
liquid		bought; existing stocks should not be used in work with open test tubes.
Tetrachloroethene		WARNING: suspected of causing cancer; toxic to aquatic life with long-lasting
(Tetrachloro-	〈基〉〈對〉	effects; may cause skin and serious eye irritation. Used in dry cleaning. For
ethylene) liquid		15-minute exposure, conc. in atmosphere should not exceed 689 mg m ⁻³ .
carry rerrey require	HEALTH HAZ. ENVIRON.	
Trichloroethene	A A	DANGER: causes skin & serious eye irritation; may cause cancer, drowsiness or
(Trichloroethylene)		dizziness; suspected of causing genetic defects; harmful to aquatic life with
liquid		long-lasting effects. For a 15-minute exposure, the concentration in the
	HEALTH HAZ. IRRITANT	atmosphere should not exceed 820 mg m ⁻³ .
	TEXETITIVE: INIVITAINT	It was used in dry cleaning but has been replaced by tetrachloroethene.

Typical control measures to reduce risk

- Use the smallest volume possible and wear suitable eye protection.
- Use a fume cupboard for anything larger than test-tube amounts; ensure good laboratory ventilation.
- When choosing a solvent, pick the safest one with suitable properties cyclohexane, Volasils or Lotoxane are safer than chlorinated hydrocarbons and usually work satisfactorily.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong? How serious would it be if something did go wrong?
- How can the risk(s) be controlled for this activity?

eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

Emergency action

In the eye Flood the eye with gently-running tap water for 10 minutes. Consult a medic. Vapour breathed in Remove the casualty to fresh air. Consult a medic. **Swallowed** Do no more than wash out the mouth with water. Do **not** induce vomiting. Consult a medic. Spilt on the skin or Remove contaminated clothing. Wash the skin with soap and water. Take contaminated clothing outside for the solvent to evaporate. clothing Spilt on the floor, Open windows if large amounts are spilt. Consider the need to evacuate for large spills. Cover with mineral absorbent (eg, cat litter) and scoop into a bucket. Add washing-up liquid and work bench, etc

into an emulsion. Wash to waste with plenty of water.

Hydrocarbons

Substance	Hazard	Comment
Methane (natural gas); ethane; propane (Calor gas, camping gas); butane (lighter fuel); LPG is a mixture of propane & butane.	FLAMMABLE	DANGER: Extremely flammable gases; asphyxiants. Mixtures with air between 6% and 12% methane by volume are explosive, others similar. Mixtures may ignite below 650 °C. Butane is easily liquefied under pressure (it normally boils at 0 °C) and both it and propane are denser than air. For a 15-minute exposure, the concentration of butane in the atmosphere should not exceed 1810 mg m ⁻³ .
Pentane, hexane, heptane, etc; cyclo- hexane, cyclohexene; petrol (gasoline); paraffin (kerosine); benzene; methylbenzene (toluene); dimethylbenzene* (xylene) (* WARNING: flammable liquid & vapour, skin irritant, harmful if inhaled).	FLAMM. HEALTH HAZ. I ENVIRON.	DANGER: (highly) flammable liquid & vapour; may be fatal if swallowed and enters airways; may cause drowsiness or dizziness; (very) toxic to aquatic life with long-lasting effects. <i>Pentane</i> : repeated exposure may cause skin dryness/cracking. <i>Hexane</i> : causes skin irritation (also heptane, cyclohexane, paraffin, benzene); suspected of damaging fertility; may cause damage to organs through prolonged/repeated exposure (also benzene). Benzene: causes serious eye irritation; may cause genetic defects and cancer. <i>Petroleum spirits (ethers)</i> 40-60, 60-80, 80-100; 100-120 °C and petrol are mixtures of alkanes of variable composition – assume similar hazards if similar boiling points. Use of benzene is no longer banned in educational laboratories but is not recommended. For a 15-minute exposure, concentration of benzene in the atmosphere should not exceed 9.75 mg m ⁻³ .
Diesel fuel; engine oil Naphthalene	CORR. IRRIT. ENVIRON. HEALTH HARM. ENVIR.	DANGER: (Diesel fuel) causes serious eye damage & skin irritation; toxic to aquatic life with long-lasting effects. Some oils may contain substances which cause cancer. After oil has been used in car engines, it may have broken down into more hazardous products. WARNING: Harmful if swallowed; suspected of causing cancer by inhalation; very toxic to aquatic life with long-lasting effects. Used in moth balls. If heated, concentration of vapour increases considerably.
Waxes; oils	LOW HAZARD	Includes: paraffin wax, candle wax, petroleum jelly, Vaseline; medicinal paraffin, liquid paraffin, oil for oil baths.

Typical control measures to reduce risk

- Use smallest amount possible; wear eye protection; avoid skin contact; make sure room is well ventilated.
- Use fume cupboard or prevent escape of vapour, eg with mineral wool plug in test tube.
- Check gas supplies for leaks; store bottled gas in a cool place; use "spirit burners" with care.
- Check equipment to put out fires, eg damp cloth, bench mat, fire blanket.
- Do not use the highly flammable liquids near naked flames; if heating is necessary, use an electrically-heated water bath or hot water from a kettle.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong? eg, does hydrocarbon need to be heated? Could quantities of the vapour be breathed in?
- How serious would it be if something did go wrong?
- How can the risk(s) be controlled for this activity? eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

•	In the eye	Flood the eye with gently-running tap water for 10 minutes. Consult a medic.
•	Vapour breathed in	Remove the casualty to fresh air. Consult a medic.
•	Swallowed	Do no more than wash out the mouth with water. Do not induce vomiting. Consult a medic.
•	Spilt on the skin or	Remove contaminated clothing. Wash the skin with soap and water. Take contaminated
	clothing	clothing outside for the solvent to evaporate.
•	Spilt on the floor,	Open windows if large amounts are spilt. Consider the need to evacuate for large spills. Cover
	bench, etc	with mineral absorbent (eg, cat litter) and scoop into a bucket. Add washing-up liquid and work
		into an emulsion. Wash to waste with plenty of water.

Carbohydrates

Substance	Hazard	Comment
Sucrose Solid	LOW HAZARD	This is ordinary table sugar. It is cane sugar (also obtained from sugar beet). Acids formed by bacterial decomposition of sugar in the mouth cause tooth decay. Excess sugars in the diet, coupled with a lack of exercise can cause
		obesity and diabetes, and can lead to heart disease. Eating in laboratories is usually illegal under the COSHH Regulations because of the risk of contamination.
Glucose	LOW HAZARD	It is also known as dextrose. Acids formed by bacterial decomposition in the mouth cause tooth decay. Excess sugars in the diet, coupled with a lack of exercise can cause obesity and diabetes, and can lead to heart disease.
Fructose	LOW HAZARD	It is also known as laevulose or fruit sugar. Acids formed by bacterial decomposition in the mouth cause tooth decay. Excess sugars in the diet, coupled with a lack of exercise can cause obesity and diabetes, and can lead to heart disease.
Maltose	LOW HAZARD	It is also known as malt sugar.
Lactose	LOW HAZARD	It is also known as milk sugar.
Starch	LOW HAZARD	Starch is broken down by saliva and stomach acids into simple sugars.
Cellulose	LOW HAZARD	It is an approved food additive, E460. Cellulose is derived from the cell walls of fruit, vegetables and cereals. It is not digested. Dietary fibre (roughage) is an important part of the diet, helping prevent various diseases.
Food testing on ca	arbohydrates	See CLEAPSS Student Safety Sheet 4.

Typical control measures to reduce risk

- Wear eye protection when handling hazardous solids and solutions, eg, when food testing.
- Do not consume sugars or indeed any food or drink in laboratories; taste-testing investigations must be done outside laboratories unless scrupulous hygiene and no contamination can be ensured.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong?
- How serious would it be if something did go wrong? eg, small amounts of contaminants entering the mouth during taste-testing activities.
- How can the risk(s) be controlled for this activity? eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

Emera	ancv	action

	norgono, action	
•	In the eye	Flood the eye with gently-running tap water for 10 minutes. Consult a medic if pain persists.
		per 31313.
•	Swallowed	In small amounts, unlikely to be hazardous unless contaminated.
•	Spilt on the skin or clothing	Brush solid off contaminated clothing. Rinse clothing or the skin as necessary.
•	Spilt on the floor, bench, etc	Brush up solid spills, trying to avoid raising dust, then wipe with a damp cloth.
	•	Wipe up solution spills with a cloth and rinse it well.



Methanol Methyl alcohol

Substance	Hazard	Comment
Methanol Liquid Methyl alcohol, also known as wood alcohol	FLAMMABLE	Danger: highly flammable liquid and vapour; toxic if swallowed; toxic in contact with skin; toxic if inhaled; causes damage to organs if swallowed, through contact with skin or if inhaled. For a 15-minute exposure, the concentration in the atmosphere should not exceed 333 mg m ⁻³ . The flash point is 11°C, ie the liquid gives off sufficient vapour at 11°C to ignite if a flame or spark is applied. Methanol is often added deliberately to ethanol ('methylated spirit') to make it undrinkable. A purple dye and an unpleasant smelling chemical
	HEALTH HAZARD	(pyridine) make the product sold to the general public even less palatable. This is called denaturing. 'Industrial denatured alcohol' (IDA) lacks the purple dye and unpleasant smelling chemical. See <i>CLEAPSS Student Safety Sheet 60</i> .

Typical control measures to reduce risk

- Wear eye protection.
- Avoid skin contact, eg by wearing protective gloves.
- Make sure the room is well ventilated or, in a laboratory, use a fume cupboard if possible.
- Check availability of ways of putting out any fires.
- Do not use near naked flames; if heating is necessary, use an electrically-heated water bath or hot water from a kettle.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong? eg, does methanol need to be heated? Could there be high levels of vapour?
- How serious would it be if something did go wrong?
- How can the risk(s) be controlled for this activity?
 - ea, can it be done safely? Does the procedure need to be altered? Should aggales or safety spectacles be worn?

eg, can it be done sajely? Does t	ne procedure need to be ditered? Should goggles or sajety spectacies be worn?
Emergency action	
• In the eye	Flood the eye with gently-running tap water for 10 minutes. Consult a medic.
Vapour breathed in	Remove the casualty to fresh air. Keep him/her warm. Consult a medic if breathing is difficult.
Swallowed	Do no more than wash out the mouth with water. Do not induce vomiting. Consult a medic.
Clothing catches fire	Smother flames on clothing or the skin with a fire blanket or other material. Cool any burnt skin with gently-running tap water for 10 minutes.
Other methanol fires	Allow fires in sinks, etc to burn out. Fires at the top of test tubes, beakers, etc should be smothered with a damp cloth or heat-resistant mat.
• Spilt on the skin or clothing	Remove contaminated clothing. If more than a test-tube amount was involved, wash the affected area and clothing with plenty of water.

Higher alcohols

including Propanol, Butanol, Pentanol

Substance	Hazard	Comment
Propan-1-ol	\wedge	DANGER: highly flammable liquid & vapour; may cause drowsiness or dizziness;
(n-propanol),		causes serious eye damage (propan-1-ol)/irritation (propan-2-ol). For a 15-
Propan-2-ol	FLAMMABLE IRRITANT	minute exposure, the concentration of vapour in the atmosphere should not
(iso-propanol)	^	exceed 625 mg m ⁻³ (propan-1-ol), 1250 mg m ⁻³ (propan-2-ol). The liquid gives
Liquids		off sufficient vapour at 22°C (-1-ol)/12°C (-2-ol) to ignite if a flame or spark is
		applied.Propan-2-ol is sometimes used as 'rubbing alcohol', especially in the
	CORROSIVE	USA. Also used to clean optical surfaces, eg spectacle and camera lenses.
Butan-1-ol	\wedge	DANGER: flammable liquid & vapour; causes skin irritation & serious eye
(n-butanol),		damage; may cause respiratory irritation & drowsiness or dizziness; butan-1-ol
2-methylpropan-1-	FLAMMABLE IRRITANT	is harmful if swallowed.
ol	^	For a 15-minute exposure, the concentration of vapour in the atmosphere
(iso-butanol),	* B	should not exceed 154 mg m ⁻³ (butan-1-ol), 231 mg m ⁻³ (2-methylpropan-1-ol).
Liquids		The flash point of butan-1-ol is 37°C (2-methylpropan-1-ol 28°C), ie the liquid
	CORROSIVE	gives off sufficient vapour at 37°C/28°C to ignite if a flame or spark is applied.
2-methylpropan-2-	\wedge	DANGER: highly flammable liquid & vapour; causes serious eye irritation;
ol (tert-butyl alcohol)	⟨७⟩⟨!⟩	harmful if inhaled; may cause respiratory irritation. For a 15-minute exposure,
Liquid but may be		the concentration of vapour in the atmosphere should not exceed 462 mg m ⁻³ .
solid on cold day	FLAMMABLE IRRITANT	Liquid gives off sufficient vapour at 11°C to ignite if a flame or spark is applied.
Butan-2-ol	A A	Warning: flammable liquid & vapour; causes serious eye irritation; may cause
(sec-butyl alcohol)		respiratory irritation & drowsiness or dizziness. For a 15-minute exposure, the
Liquid		concentration of vapour in the atmosphere should not exceed 462 mg m ⁻³ . The
	FLAMMABLE IRRITANT	flash point of is 24°C, ie the liquid gives off sufficient vapour at 24°C to ignite if
		a flame or spark is applied.
Pentan-1-ol	\wedge	Warning: flammable liquid & vapour; may cause skin, respiratory & serious
(n-amyl alcohol),	〈** 〉〈!〉	eye irritation; may be harmful if swallowed. For a 15-minute exposure, the
3-methylbutan-1-ol	\vee	concentration of 3-methylbutan-1-ol in the atmosphere should not exceed
(iso-amyl alcohol)Liqs	FLAMM. HARMFUL	458 mg m ⁻³ .

Typical control measures to reduce risk

- Wear eye protection.
- Make sure the room is well ventilated or, in a laboratory, use a fume cupboard if possible.
- Check availability of ways of putting out any fires.
- Do not use near naked flames; if heating necessary, use an electrically-heated water bath or hot water from kettle.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong? eg, does alcohol need to be heated? Could there be high levels of vapour?
- How serious would it be if something did go wrong?
- How can the risk(s) be controlled for this activity?

eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

Emergency action

ш	nergency action	
•	In the eye	Flood the eye with gently-running tap water for 10 minutes. Consult a medic.
•	Vapour breathed in	Remove the casualty to fresh air. Keep him/her warm. Consult a medic if breathing is difficult.
•	Swallowed	Do no more than wash out the mouth with water. Do not induce vomiting. Consult a medic.
•	Clothing catches fire	Smother flames on clothing or the skin with a fire blanket or other material. Cool any burnt skin
		with gently-running tap water for 10 minutes.
•	Other fires	Allow fires in sinks, etc to burn out. Fires at the top of test tubes, beakers, etc should be
		smothered with a damp cloth or heat-resistant mat.
•	Spilt on the skin or	Remove contaminated clothing. If more than a test-tube amount was involved, wash the
	clothing	affected area and clothing with plenty of water.
•	Spilt on the floor,	Put out Bunsen flames. Wipe up small amounts with cloth and rinse it well. For larger amounts,
	bench, etc	open windows, cover with mineral absorbent (eg, cat litter), scoop into a bucket and add water.

Methanal

including formaldehye, formalin

Substance	Hazard	Comment
Methanal (formalaldehyde) <i>Gas</i>	HEALTH TOXIC	DANGER: toxic if swallowed, inhaled or in contact with skin; causes severe skin burns and eye damage; may cause cancer or allergic skin react ion; suspected of causing genetic defects. For a 15-minute exposure, concentration should not exceed 2.5 mg m ⁻³ . Vapour may arise when handling solution. Commonly supplied for laboratory use as a saturated solution, about 13 M (40%).
Very concentrated methanal solution (formalin) If 10M (30%) or more	CORROSIVE	Methanal is used to make polymers, eg for cavity wall insulation – there have been suggestions that small amounts may be released into houses if polymerisation is incomplete. A methanal resin is also used as a binding agent in MDF and machining it may cause decomposition and release methanal.
Concentrated methanal solution (formalin) If less than 10M (30%) but 8M (25%) or more	HEALTH HARM.	DANGER: causes severe skin burns and eye damage; harmful by skin contact or if swallowed; irritating to eyes, skin & respiratory system; may cause cancer or allergic skin reaction; suspected of causing genetic defects.
Moderately dilute methanal solution (formalin) If less than 8M (25%) but more than 0.07 M (0.2%)	HEALTH HAZARD HARMFUL	DANGER: Harmful by skin contact (if \geq 5M (15%)). Harmful if swallowed; irritating to skin, eyes & respiratory system (if \geq 1.7 M (5%)). Suspected of causing genetic effects (if \geq 0.3M (1%)). May cause cancer or allergic skin reaction. A 1.3 M (4%) solution was commonly used in the past as a preservative for biological specimens. A solution of concentration between 0.07 M (0.2%) and 0.3 M (1%) was used in the [past for extracting earthworms from soil but there are safer alternatives which are also less damaging to the worms.
Dilute methanal solution (formalin) If less than 0.07 M (0.2%) but more than 0.03 M (0.1%)	HEALTH HAZARD	DANGER: may cause cancer.
Very dilute methanal if less than 0.03M (0.1%)	LOW HAZARD	-

Typical control measures to reduce risk

- Wear appropriate eye protection (depending on the concentration) and, for all except dilute solutions of methanal, wear protective gloves (preferably nitrile).
- Open bottles of all except dilute solutions of methanal in a fume cupboard.
- Do not use near naked flames; if heating is necessary, use an electrically-heated water bath a kettle.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong?
- How serious would it be if something did go wrong?
- How can the risk(s) be controlled for this activity?
 - eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

•	In the eye	Immediately flood the eye with gently-running tap water for 10 minutes. Consult a medic.
•	Vapour breathed in	Remove the casualty to fresh air. Keep him/her warm. Consult a medic if breathing is difficult.
•	Swallowed	Do no more than wash out the mouth with water. Do not induce vomiting. Consult a medic.
•	Spilt on the skin/clothing	Remove contaminated clothing. Wash the affected area and clothing with plenty of water.
•	Spilt on the floor, bench, etc	Put out all Bunsen flames. Wipe up small amounts with a cloth and rinse well. For larger amounts, open windows, if fumes are not too strong cover with mineral absorbent (eg, cat litter), scoop into a bucket and add water.

Ethanal and higher aldehydes

including propanal, butanal

Substance	Hazard	Comment
Ethanal		DANGER: extremely flammable liquid and vapour; causes serious eye irritation;
(acetaldehyde)		may cause respiratory irritation; suspected of causing cancer.
Liquid with boiling		For a 15-minute exposure, the concentration in the atmosphere should not
point close to room	FLAMM. IRRITANT	exceed 92 mg m ⁻³ . The flash point is -39°C, ie the liquid gives off sufficient
temperature (21 °C)		vapour at -39°C to ignite if a flame or spark is applied.
		May boil spontaneously during storage or on opening. May spurt out of the
		bottle as the pressure is released when it is opened. Tiny traces of impurity
	HEALTH HAZARD	can cause spontaneous boiling. Cool in an ice bath before opening in a fume
		cupboard.
Ethanal tetramer		WARNING: flammable solid; harmful if swallowed.
(metaldehyde,		Sometimes used as solid fuel for model steam engines but it may spit as it
2,4,6,8-tetramethyl-		burns. Hexamethylene is possibly safer.
1,3,5,7-	FLAMM. HARMFUL	
tetraoxacyclooctane)		
Solid		
Propanal		DANGER: highly flammable liquid and vapour; causes skin and serious eye
(propionaldehyde)		irritation; may cause respiratory irritation.
Liquid		The flash point is -30°C, ie the liquid gives off sufficient vapour at -30°C to
	FLAMM. HARMFUL	ignite if a flame or spark is applied.
		Because of its higher boiling point (48 °C), it is much safer to use than ethanal.
Butanal		DANGER: highly flammable liquid and vapour.
(butyraldehyde)	<u> </u>	The flash point is -22°C, ie the liquid gives off sufficient vapour at -22°C to
Liquid	•	ignite if a flame or spark is applied.
	FLAMMABLE	Because of its higher boiling (75 °C), it is much safer to use than ethanal.

Typical control measures to reduce risk

- Wear eye protection and protective gloves (preferably nitrile).
- Open ethanal bottles very cautiously in a fume cupboard, after cooling in an ice bath.
- Avoid using ethanal if at all possible; use propanal or butanal instead.
- Do not use near naked flames; if heating is necessary, use an electrically-heated water bath or hot water from a kettle.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong?
- How serious would it be if something did go wrong?
- How can the risk(s) be controlled for this activity?

eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

=	mergency action	
•	In the eye	Immediately flood the eye with gently-running tap water for 10 minutes. Consult a medic.
•	Vapour breathed in	Remove the casualty to fresh air. Keep him/her warm. Consult a medic if breathing is difficult.
•	Swallowed	Do no more than wash out the mouth with water. Do not induce vomiting. Consult a medic.
•	Spilt on the skin/clothing	Remove contaminated clothing. Wash the affected area and clothing with plenty of water.
•	Spilt on the floor, bench, etc	Put out all Bunsen flames. Wipe up small amounts with a cloth and rinse well. For larger amounts, open windows, if fumes are not too strong cover with mineral absorbent (eg, cat litter), scoop into a bucket and add water.

Dyes, stains & indicators

Substance	Hazard	Comment
Solid dyes, stains & indicators including:	\wedge	DANGER: May include one or more of the following
Acridine orange, Congo Red (Direct dye 28), Crystal violet		statements: fatal/toxic if swallowed/in contact
(methyl violet, Gentian Violet, Gram's stain), Ethidium	TOXIC HEALTH	with skin/ if inhaled; causes severe skin burns &
bromide, Malachite green (solvent green 1), Methyl		eye damage/ serious eye damage; may cause
orange, Nigrosin, Phenolphthalein, Rosaniline, Safranin	\vee	allergy or asthma symptoms or breathing
	CORR. IRRIT.	difficulties if inhaled; may cause genetic defects/
	¥	cancer/damage fertility or the unborn child;
	<u>~</u>	causes damages to organs/through prolonged or
	ENVIRONMENT	repeated exposure.
Solid dyes, stains & indicators including Alizarin (1,2-		WARNING: May include one or more of the
dihydroxyanthraquinone), Alizarin Red S, Aluminon (tri-		following statements: harmful if swallowed/in
ammonium aurine tricarboxylate), Aniline Blue (cotton /		contact with skin/if inhaled; causes skin/serious
spirit blue), Brilliant yellow, Cresol Red, DCPIP (2,6-dichl-		eye irritation; may cause allergic skin reaction;
orophenolindophenol, phenolindo-2,6-dichlorophenol,	HEALTH	suspected of causing genetic
PIDCP), Direct Red 23, Disperse Yellow 7, Dithizone (di-		defects/cancer/damaging fertility or the unborn
phenylthiocarbazone), Eosin (Eosin Y), Eriochrome Black T	<u>~</u>	child; may cause damage to organs/respiratory
(Solochrome black), Fluorescein (& disodium salt), Haem-	HARMFUL	irritation/drowsiness or dizziness/damage to
atoxylin, HHSNNA (Patton & Reeder's indicator), Indigo,	《鉴》	organs through prolonged or repeated exposure.
Magenta (basic Fuchsin), May-Grunwald stain, Methyl-	ENVIRON.	
ene blue, Methyl green, Orcein, Phenol Red, Procion		
dyes, Pyronin, Resazurin, Sudan I/II/IV dyes, Sudan black		
(Solvent Black 3), Thymol blue, Xylene cyanol FF		Some dyes may contain hazardous impurities and
Solid dyes, stains & indicators including Acid blue 40, Blue dextran, Bromocresol green,		many have not been well researched. Suppliers
Bromophenol blue, Carmine (cochineal, Natural Red 4),		often vary in the hazard classifications they use.
Diazine Green (Janus Green B), Indigo carmine, Litmus,	LOW HAZARD	So although the substances listed here are not
Methyl blue, Methyl red, Murexide (ammonium	LOWINZARD	usually classified as hazardous they should be
purpurate), Neutral red, Rhodizonic acid, Sudan III,		used with caution. Skin contamination should be
Thymolphthalein, Toluidine blue, Xylenol orange		avoided.
Dilute solutions of the above	DEPENDS ON	Aqueous solutions low risk but with non-aqueous
Dyes, stains & indicators, usually less than 1% in water or	SOLVENT	solvents risks greater, eg, ethanol [FLAMMABLE if
non-aqueous solvents		more than 25% (v/v)] or ethanoic acid [CORROSIVE].
non-aqueous solvents		more than 25% (V/V)] or ethanoic acid [CORROSIVE].

Typical control measures to reduce risk

- Use the lowest concentration possible and wear eye protection for all but the most-dilute solutions.
- Reduce the risk of skin contact by wearing disposable gloves.
- Avoid powdered dyes & indicators escaping into the air; use a fume cupboard when handling more hazardous ones.
- Avoid naked flames if using flammable solvents.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong? Eg, could dust from a dye or indicator be breathed in?
- How serious would it be if something did go wrong? Eq, would it be more serious than the skin being stained for a few days?
- How can the risk(s) be controlled for this activity?
 - eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

- In the eye Immediately flood the eye with gently-running tap water for 10 minutes. Consult a medic. Do no more than wash out the mouth with water. Do **not** induce vomiting. Consult a medic. **Swallowed Dust breathed in** Remove the casualty to fresh air. Consult a medic if breathing difficult.
- Remove contaminated clothing. Wash off the skin with soap and plenty of water. Rinse contaminated Spilt on skin or clothing. clothing
- Spilt on floor, Scoop up solids (take care not to raise dust). Wipe up solution spills or any traces of solid with a damp cloth and rinse it well. bench, etc



Sharps

including scalpels, knives, syringe needles, seekers, etc

Source	Hazard	Comment
Scalpels, knives and		Cuts and puncture wounds can lead to infection, especially if the blade or point
other blades	DANGER	is contaminated by contact with living or once-living material. Careless use and handling of scalpels, syringes with needles, seekers and other sharps can lead to cuts and puncture wounds.
Syringe needles		Sharp scalpels are safer to use than blunt ones because there is less risk of them slipping as less force needs to be used.
Seekers and other	DANGER	Carrying scalpels, syringes with needles, seekers and other sharps especially in crowded rooms, can present a hazard to the user and others.
sharps	DANGER	Carelessly-disposed sharps can present a hazard to waste handlers and others.

Typical control measures to reduce risk

- Find out from a reliable source if there any health & safety issues relating to to the material being dissected.
- Cut in a direction away from yourself and where possible cut using a cutting board, dissection tray, a pad or similar.
- Wear eye protection when changing scalpel blades or cutting material likely to "flick" (eg, cartilage or bone).
- Cooperate with any requirement to count sharps at the beginning and end of a lesson.
- Carry sharps with the blade or point protected, eg in a shallow tray, and do not carry them at all if you are likely to be
- Dispose of used sharps in a proper, safe container, eg a sturdy box, clearly labelled, and sealed and wrapped before disposal.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong? eg, could the user or somebody else be cut or stabbed by accident?
- How serious would it be if something did go wrong?
- How can the risk(s) be controlled for this activity? eg, can it be done safely? Does the procedure need to be altered?

- Wash the wound. Get the casualty to apply a small, sterilised dressing. Minor cuts
- Severe cuts Lower the casualty to the floor. Raise the wound as high as possible. If feasible, ask the casualty to apply pressure on or as close to the cut as possible, using fingers, a pad of cloth or, better, a sterile dressing (adding further layers as necessary). If the casualty is unable to do so, apply pressure yourself, protecting your skin and clothes from contamination by blood if possible. Leave any embedded large bodies and press around them. Consult a medic.

Animals (dead) and animal parts

See also CLEAPSS Student Safety Sheet 71: Sharps.

Source	Hazard	Comment
Animals (whole) (fresh or recently- defrosted from frozen), eg whole rats, mice, fish and		Whole animals obtained from a reputable biological supplier should be safe to use; but road kill, for example, might be infected. Items intended for human consumption, available from butchers, abattoirs and fishmongers, should also be safe. Because of the risk of BSE it would only be legal to supply cattle eyes (or brain tissue) from animals slaughtered at less than 12 months or eyes of sheep and goats from animals slaughtered at less than 12 months or with at least one erupted incisor.
organs such as eyes, hearts, kidneys, lungs, and chicken's feet/legs and bones.	BIOHAZARD	If only bones are to be studied, and especially if they are to be kept, use a knife to remove as much flesh as possible. Place the bones in a saucepan of water to which sodium carbonate is added and <i>simmer</i> until the remaining flesh can easily be removed, using an old brush. Return to the pan for more simmering until the bones are cleaned.
bones.		Some people have cultural or religious objections to handling particular species.
		Some people object to killing animals, whether for food, medical research or dissection in schools. Fewer people object to the use of material intended for human consumption available from butchers, abattoirs and fishmongers.
Animal parts (preserved)		In the past, specimens were preserved in 1.3 M (4%) methanal solution (formalin) (see CLEAPSS Student Safety Sheet 67). Formalin-free preservatives are mostly used now, but formalin will still have been used professionally as a fixing agent.
	HEALTH HAZARD	Sometimes 70% ethanol or propanol solutions are used (see CLEAPSS Student Safety Sheets 60 and 66).
	HARMFUL	Preservative should always be rinsed off before use, preferably soaking for an hour in water. However, preservative may still remain in body cavities, which must be rinsed as soon as they are exposed.
Dissection		See CLEAPSS Student Safety Sheet 71: Sharps.
	DANGER	

Typical control measures to reduce risk

- Use material from reliable sources.
- Wash preserved material prior to dissection and rinse it if body cavities are exposed during dissection.
- Wear eye protection when cutting bone or cartilage, using preserved material or changing scalpel blades.
- Count sharps at the beginning and end of the lesson and carry around in a safe manner.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong?
- How serious would it be if something did go wrong?
- How can the risk(s) be controlled for this activity? eg, can it be done safely? Does the procedure need to be altered?

Emergency action

 Animal material in the eye Flood the eye with gently-running tap water for 10 minutes. If discomfort persists, consult a medic.

Animals (living)

See also CLEAPSS Student Safety Sheet 75: Fieldwork

Source	Hazard	Comment
Vertebrates including		It is illegal to treat vertebrates in a manner causing pain, suffering, distress or lasting harm. Such cruelty might result from some experiments, poor handling, unsuitable housing or inadequate feeding.
small mammals, fish, birds, reptiles, amphibia	BIOHAZARD	Some animals may bite and some people may be allergic to hairs, skin, scales, feathers, droppings, etc. Some animals can present a health hazard, although the risk of diseases being passed to humans is usually low but may be higher for farm animals. Wild animals can harbour diseases and parasites, especially if injured. Obtaining animals from reputable sources, preventing contact with wild species and adopting good hygiene practices will usually make the risk insignificant. A few species present higher risks and should be avoided. Bees and farm animals need special facilities and specialist knowledge. Some native species are protected and must not be brought in from the
		wild. It is illegal to release any non-native species into the wild.
Invertebrates including		Although cruelty to invertebrates is not an offence, they should still be treated humanely, handled carefully and housed and fed in a suitable manner.
insects, snails, worms, brine shrimps, water fleas,	BIOHAZARD	Any garden 'minibeasts' brought into school for study should be returned to the environment from which they came as quickly as possible. It is illegal to release any non-native species into the wild.
etc		Some animals may sting and some people may be allergic to hairs, skin, scales, droppings, etc.

After use, healthy animals taken from the wild should be returned to the place from which they were taken.

Typical control measures to reduce risk

- Check detailed guidance on suitability, handling, housing and feeding of individual species.
- Use material from reliable sources; avoid species presenting higher risks; check that mammals are docile.
- Ensure laboratory mammals cannot come into contact with wild rodents.
- Before handling animals, cover cuts and abrasions on exposed hands and arms. Gauntlets may be necessary when handling some animals. If animals of any sort are to be handled regularly, it is a good idea to have up-to-date antitetanus vaccination.
- Wash hands in warm soapy water before and soon after handling animals, or coming into contact with their bedding, water, droppings or housing.
- Wear gloves when cleaning cages or handling soiled bedding; dispose of such bedding by incineration or in a sealed plastic sack with normal refuse.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong?
- How serious would it be if something did go wrong?
- How can the risk(s) be controlled for this activity? eg, can it be done safely? Does the procedure need to be altered?

Emergency action

Animal material in the eye Flood the eye with gently-running tap water for 10 minutes. Do not attempt to remove any embedded object. If discomfort persists, consult a medic.

Wash the wound with warm soapy water; raise and support the wound, cover with a **Animal bites**

sterile dressing. Consult a medic.

Plants, fungi and seeds

See also CLEAPSS Student Safety Sheet 75: Fieldwork.

Source	Hazard	Comment
Plants and fungi	TOXIC	Some plants, eg cacti, have spines which can cause deep puncture wounds and other spines may cause skin irritation. Other plants, eg <i>Euphorbia</i> have irritant or poisonous sap. Nettles are well known but hyacinth bulbs can also cause dermatitis (skin rash), as can bulbs of other members of the liliaceae family such as daffodils, tulips and primulas. Many common garden and hedgerow plants are poisonous or have toxic parts. Some <i>Umbellifers</i> , eg giant hogweed, can cause photosensitisation after skin contact, resulting in blisters on exposure to light. Some plants used as common food sources may have poisonous parts, eg rhubarb leaves and the green parts of the potato (including the tuber) and tomato. Some fungi are very poisonous and easily mistaken for the common (field) mushroom. Pollen from some plants can produce an allergic reaction in susceptible individuals, commonly known as hay fever. Plant material collected from farming areas may have been contaminated with pesticides, although most rapidly break down after spraying. Wild plants are protected by law. It is illegal to uproot, pick or destroy certain highly protected plants and it is illegal to uproot any plant without the landowner's permission.
Seeds	TOXIC	Some seeds are very poisonous, eg castor oil, laburnum. French beans or red kidney beans are poisonous unless cooked. Seeds bought from commercial suppliers are often treated with a pesticide to prevent attack by insects or fungi. Seeds from health food stores will not have been treated. The effects of radiation on plant growth are often investigated using seeds that have been exposed to gamma-radiation. Such seeds are NOT radioactive.

Typical control measures to reduce risk

- Check reliable sources to find if there are hazards in any plants in your garden, in the neighbourhood or in school.
- Where possible, use seeds known to be pesticide free or rinse the seeds in running water for several hours.
- Treat unknown plants with caution; never taste or eat a plant unless it is definitely known to be non-poisonous.
- Wash hands in warm soapy water after handling seeds and other plant material.
- Wear gloves when handling plants likely to cause irritation or seeds treated with pesticides.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong?
- How serious would it be if something did go wrong?
- How can the risk(s) be controlled for this activity?

eq, can it be done safely? Does the procedure need to be altered?

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•	Skin rash from irritant plants	Cool the affected area with cold water (if available) and cover with a sterile dressing. You might self-medicate using a traditional remedy by rubbing a nettle sting (acidic) with a dock leaf (alkaline).
•	Poisonous seeds swallowed	Do no more than wash out the mouth with drinking water. Do NOT induce vomiting. Consult a medic.



Fieldwork

including any science work outside the laboratory

See also CLEAPSS Student Safety Sheets 73, 74.

Source	Hazard	Comment
Site-specific hazards	DANGER BIO- HAZARD	Schools will normally have identified any significant hazards in school grounds, as will other sites which regularly welcome schools for fieldwork but broken glass and animal faeces are always possible. Working outdoors, eg at the seaside, in the countryside or near water will introduce hazards specific to the location, eg drowning, trapping by tides, falls from a height, slips and trips, plants with poisonous parts, pesticides, animal bites/stings and animal-borne infections eg toxocariasis from soil contaminated with cat or dog faeces, Weil's disease from water contaminated with rat urine and Lyme disease from ticks. Working outdoors can lead to exposure to weather conditions which present hazards, eg
		strong sunlight (near ultraviolet, see CLEAPSS Student Safety Sheet 12), freezing cold, rain.
Transport hazards	DANGER	Walking to the site may result in exposure to traffic hazards. Public transport, hired coaches, the school minibus or private cars introduce a variety of different hazards.
Child abuse &/or disappearance	DANGER	Children in unfamiliar settings may wander off. If children are in contact with members of the public, there is a small possibility of child abuse.

Typical control measures to reduce risk

- As far as possible only visits sites you know to be safe and follow the Code of Practice for that type of fieldwork.
- Ensure that others know where you are and make sure you use safe equipment and transport arrangements.
- When working in shoreline ecologies check tide tables and beware of the risk of slipping on seaweed.
- Ensure shoes and clothing are suitable for the terrain and for varying weather conditions; eg, avoid bare legs and arms in areas known to be infested with ticks or in sunny conditions, wear sun hats (and use sun cream).
- Wear gloves when handling plants likely to cause irritation or seeds treated with pesticides; wear hard hats when working in quarries and eye protection when hammering rocks.
- Avoid contact with water with growths of blue-green algae on the surface; cover cuts and abrasions with water-proof dressings when working in and around water which may be contaminated by rat urine.
- Treat unknown plants with caution; never taste or eat a plant unless it is definitely known to be non-poisonous.
- Approach vertebrates with caution, especially those with young.
- Ensure easy access to hand-washing facilities before eating/drinking; if soap/water unavailable use alcohol gel.
- Beware of the possibility of hay fever, etc. Carry a first-aid kit and ensure at least one person is trained to use it.
- Ensure there is a workable emergency contact system (there may be no mobile phone signal in some areas).

Assessing the risks

- What are the details of the activity to be undertaken? Hazards?
- What is the chance of something going wrong?
- How serious would it be if something did go wrong?
- How can the risk(s) be controlled for this activity? eg, can it be done safely? Does the procedure need to be altered?

- Insect bites & If sting is visible and can be displaced easily, brush off sideways with the edge of a credit card or finger nail, avoiding further damage. Raise affected body part and if possible apply cold compress for 10+ min. stings
- Remove the tick with forceps by grasping as close to the skin as possible and pulling gently without **Tick bites** twisting or jerking. Keep the tick for identification and consult a medic.
- Animal bites Wash the wound with warm soapy water or, if unavailable, alcohol-free cleansing wipes. Raise and support the wound, cover with a sterile dressing. Consult a medic.
- Wash the wound or, if water is unavailable, use cleansing wipes. Get the casualty to apply a small, sterile Minor cuts dressing...

Bioreactors and **fermenters**

See also CLEAPSS Student Safety Sheet 1, Microorganisms

Source	Hazard	Comment
Micro- organisms	BIOHAZARD	Bioreactors and fermenters present greater risks than agar plates because of the much greater volume of medium involved and the possibility of liquid spills and aerosol formation. Schools should restrict work to yeasts or bacteria with unusual growing conditions, eg, low pH, high salt concentration or specialist media. Generation of biogas from silage or pond mud is acceptable but the use of animal manure as an inoculum should be avoided because it may introduce pathogens and/or medication administered to animals.
Electrical equipment	ELECTRIC SHOCK	The proximity of large volumes of liquid and mains electrical equipment (eg, from heaters, aerators, sensors, etc) presents a hazard. Commercially-designed equipment from a reliable source should prevent access to live conductors (check the integrity before each use) but d-i-y equipment should not exceed 25 V.
Gases	EXPLOSION HIGHLY FLAMMABLE	Depending on the reactions taking place, large volumes of gas may be produced, usually carbon dioxide or methane (biogas). The vessel must be vented to avoid the build-up of pressure but it is important to prevent the entry of external microorganisms or the release of aerosols. Care must be taken to ensure no naked flames are near the bioreactor if a highly flammable gas such as methane is being produced.
Sterilisation	DANGER	Steam sterilisation may be impossible if the bioreactor is too large to fit into an autoclave, so chemical disinfection will be necessary. Some possible disinfectants are hazardous. <i>Virkon</i> is the preferred disinfectant.

Typical control measures to reduce risk

- Use only safe microorganisms from safe sources.
- Do not seal bioreactors but prevent the entry of external microorganisms and the release of aerosols.
- Sterilise all equipment before and after use and sterilise the culture before disposal.
- When withdrawing samples, take care to avoid aerosol production and place a pad of tissues moistened with disinfectant underneath to catch drips.
- Keep electrical leads tidy and site mains equipment as far away from the reactor as possible.
- Always wash hands after handling cultures.
- Wear a clean lab. coat or overall to protect cultures from microbes on the skin, clothing, etc.

Assessing the risks

- What are the details of the activity to be undertaken? Hazards?
- What is the chance of something going wrong? eg, could a culture become contaminated? Could microorganisms escape?
- How serious would it be if something did go wrong? eg, could there be an explosion resulting from a pressure build-up or ignition of a flammable product?
- How can the risk(s) be controlled for this activity? Eg, can it be done safely? Does the procedure need to be altered?

Emergency action

For spills of cultures, place paper towels over the spill, pour disinfectant (eg, Virkon) on top and Spilt on the floor. bench, etc leave for at least 15 minutes.

Working with DNA

See also CLEAPSS Student Safety Sheet 78: Genetic modification

Source	Hazard	Comment
'Naked' DNA (ie DNA not incorporated into a living cell)	LOW HAZARD	DNA only functions when it is inserted into a living cell, hence work with DNA itself is generally low hazard, although there may be other hazards associated with the process, eg chemical, microbiological or electrical hazards (electrophoresis).
Extraction of DNA from human tissue	BIOHAZARD	Extraction of DNA from human tissue, eg cheek cells, prior to amplification by the polymerase chain reaction (PCR), could result in the transfer of infective material between participants. See CLEAPSS Student Safety Sheet 3.
DNA from laboratory suppliers	BIOHAZARD	DNA from sources such as bacteriophage lambda and salmon sperm is generally safe but DNA from mammalian sources may be contaminated with viruses.
Gel electrophoresis	ELECTRIC SHOCK	Electrophoresis can be very slow unless moderately high voltages are used, giving a risk of electric shock, especially because of the high conductivity of the buffer solutions. If voltages in excess of 30V AC or DC are used it must be impossible to touch a live conductor accidentally or to open the tank if a current is flowing. Some commercial tanks, especially if imported from the USA, may not satisfy this requirement.
Chemicals used	TOXIC	Polyacrylamide gels are too toxic to make or cast in schools. Some stains, eg ethidium bromide are also unsuitable. Others may be used with care. See CLEAPSS Student Safety Sheet 70, Dyes, stains and indicators.

Typical control measures to reduce risk

- If extracting DNA from human tissue, you should only handle your own.
- Avoid using DNA obtained by laboratory suppliers from mammalian sources.
- Carry out electrophoresis at voltages below 30V AC or DC unless the design of tank is such that it is impossible to open the tank when a current is flowing or accidentally touch a live conductor.
- Use agarose gels, but if polyacrylamide gels are used, buy ready-made ones.
- Use safe stains such as methylene blue, Azure A or B or Nile blue sulfate; avoid ethidium bromide.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong?

eg, could a culture become contaminated? Could microorganisms escape?

- How serious would it be if something did go wrong?
 - eg, could somebody receive an electric shock from damaged or unsuitable equipment?
- How can the risk(s) be controlled for this activity?

eg, can it be done safely? Does the procedure need to be altered?

•	Spilt on the floor,	For spills of DNA extracts, place paper towels over the spill, pour disinfectant (eg, Virkon) on
	bench, etc	top and leave for at least 15 minutes.



Genetic modification

See also CLEAPSS Student Safety Sheets 1, Microbiology and 77, Working with DNA

Source	Hazard	Comment
Genetic modification	DANGER	In law, genetic modification is the alteration of genetic material (DNA or RNA) by means that could not occur naturally by mating and/or recombination. Throughout most of the world, the use of genetically-modified organisms (GMOs) is controlled by law. In the UK separate regulations control work with GMOs in the laboratory and their deliberate release into the environment. Before genetic modification (other than 'self-cloning') is undertaken the premises must be registered and approved by the HSE and other procedures put in place. Some procedures used in schools in the USA, or found on the internet, would be illegal in the UK.
'Self-cloning' (plasmid transfer) (Plasmids are small rings of DNA, comprising just a few genes)	DANGER	Returning genetic material to a species in which it could occur naturally is called self-cloning; cloning here means making copies of plasmid DNA within an organism, even if the DNA has been modified by enzymes, chemicals, etc. Providing the resulting organism is unlikely to cause disease in humans, other animals or plants no HSE registration is needed. It is still illegal to release the GMO into the environment without approval and so it must be contained. The law requires GMOs to be inactivated after use by validated means, ie by steam sterilisation.
Incubation	BIOHAZARD	Although schools normally avoid incubating at 37°C because of the risk of promoting the growth of pathogens, the particular strains of <i>E. coli</i> used for cloning work will not grow quickly or reliably at other temperatures. Hence very strict adherence to good microbiological practice is essential.

Typical control measures to reduce risk

- Do not seal cultures completely before incubation (otherwise hazardous anaerobic bacteria may be encouraged) but make sure they cannot be opened accidentally.
- Use sterile equipment and aseptic technique (eg, by flaming loops and mouths of bottles, etc).
- Avoid draughts (from open windows and doors) which could contaminate cultures.
- Work near Bunsen-burner flames so that the updraught helps to prevent contamination of cultures.
- After work is complete, treat surfaces using a suitable disinfectant, for a sufficient length of time e.g. 1% for at least
- Dispose of cultures by sterilisation in an autoclave (pressure cooker).
- Always wash hands after handling cultures.
- Wear a clean lab coat or overall to protect cultures from microbes on the skin, clothing, etc.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong? eg, could a culture become contaminated? Could microorganisms escape?
- How serious would it be if something did go wrong?
- How can the risk(s) be controlled for this activity? eg, can it be done safely? Does the procedure need to be altered?

•	Spilt on the floor,	For spills of cultures, place paper towels over the spill, pour disinfectant (eg, 1% Virkon) on top
	bench, etc	and leave for at least 15 minutes.



Alkali metals

includes lithium, sodium, potassium

Substance	Hazard	Comment
Lithium solid		DANGER: It causes severe skin burns and eye damage; in contact with water releases flammable gases which may ignite spontaneously; reacts violently with water.
	FLAMM. CORROSIVE	With water produces hydrogen, an extremely flammable gas (see CLEAPSS <i>Student Safety Sheet 50</i>). Although difficult to ignite, once lit, it burns readily in air and is difficult to extinguish.
		It reacts violently with many substances.
Sodium solid		DANGER: It causes severe skin burns and eye damage; in contact with water releases flammable gases which may ignite spontaneously; reacts violently with water.
	FLAMM. CORROSIVE	With water produces hydrogen, an extremely flammable gas (see CLEAPSS <i>Student Safety Sheet 50</i>). It burns vigorously and is difficult to extinguish. Contact with moisture produces sodium hydroxide which is corrosive (see CLEAPSS <i>Student Safety Sheet 31</i>).
		It reacts violently with many substances.
Potassium solid		DANGER: It causes severe skin burns and eye damage; in contact with water releases flammable gases which may ignite spontaneously; reacts violently with water.
	FLAMM. CORROSIVE	With water produces hydrogen, an extremely flammable gas (see CLEAPSS Student Safety Sheet 50). It burns vigorously and is difficult to extinguish. Contact with moisture produces potassium hydroxide which is corrosive (see CLEAPSS Student Safety Sheet 31). It reacts violently with many substances.
		Over a period of years, it may develop a coating of yellow superoxide. Under slight pressure, eg, from a knife blade, this may explode.

Typical control measures to reduce risk

- Store alkali metals under liquid paraffin (mineral oil) and check there is sufficient liquid.
- Check potassium samples regularly for signs of custard yellow coating; if found dispose of sample safely.
- Handle sample using forceps, wear eye protection and use safety screens.
- Conduct all investigations on a small scale generally use a rice grain-sized piece.
- Make sure everybody involved (eg, technicians clearing away) understands the hazards.
- Take steps to prevent theft.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong? eg, could molten, corrosive metal spit out of a container?
- How serious would it be if something did go wrong?
 - NB There are occasional reports of pupils being taken to hospital (for treatment to cuts or as a result of chemical splashes) as a result of explosions of apparatus involving sodium.
- How can the risk(s) be controlled for this activity?
 - eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

Emergency action

ㅁ	Emergency action			
•	In the eye	Flood the eye with gently-running tap water for 20 minutes. Consult a medic. If it is necessary to go to hospital, continue washing the eye during the journey in the ambulance.		
•	Swallowed	Do no more than wash out the mouth with drinking water. Do not induce vomiting. Consult a medic.		
•	Spilt on the skin or clothing	Remove any pieces of solid with forceps. Then drench the skin with plenty of water. If a large area is affected or blistering occurs, consult a medic.		
•	Metal catches fire	For sodium and potassium, smother with clean, dry sand. For lithium, smother with dry sodium chloride, <i>not</i> sand.		
•	Spilt on the floor, bench, etc	Scoop up as much metal as possible into a dry container. Cover the area with dry sand or anhydrous sodium carbonate (or, for lithium, sodium chloride) and scoop into a dry bucket for further treatment. Rinse the area with plenty of water and mop.		



Group II metals

includes magnesium & calcium

Substance	Hazard	Comment
Magnesium Solid powder, turnings, ribbon	FLAMMABLE	DANGER: (powder, turnings) flammable solid, self-heating in large (kilogram) quantities; may catch fire; contact with water releases flammable gases. Ribbon – classification varies – may be as powder/turnings or may be low hazard! It is moderately difficult to ignite but, once burning, it does so very vigorously and is difficult to extinguish. Ordinary fire-fighting methods are not suitable, but dry sand may be used. The flame is very bright and may damage eye sight. View through shade 9 welding filter or narrow gap between fingers.
		It reacts readily with acids to produce hydrogen, an extremely flammable gas. (See CLEAPSS Student Safety Sheet 50.) .
Calcium	^	DANGER: In contact with water releases flammable gases.
solid		It reacts readily with water (or acids) to produce hydrogen, an extremely flammable gas. (See CLEAPSS <i>Student Safety Sheet 50</i> .)
	FLAMMABLE	Contact with moisture forms calcium oxide or hydroxide which are CORROSIVE to eyes IRRITANT to skin. (See CLEAPSS <i>Student Safety Sheet 32</i>).
		It is difficult to ignite but, once burning, does so vigorously.

Typical control measures to reduce risk

- Conduct all experiments on a small scale.
- Keep careful control of stocks to prevent theft.
- Wear eye protection.
- Avoid looking directly at the flame from burning magnesium: use welding filters shade 9 or narrow gap between fingers.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong? eg, Is there the possibility of theft or foolish behaviour?
- How serious would it be if something did go wrong?
- How can the risk(s) be controlled for this activity?

eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

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•	In the eye	If magnesium powder or calcium contaminates the eyes, flood with gently-running tap water for 10 minutes. Consult a medic.
•	Swallowed	Do no more than wash out the mouth with drinking water. Consult a medic.
•	Skin burnt by burning metal or by moist calcium	Remove any pieces of solid with forceps. Then drench the skin with plenty of water. If a large area is affected or blistering occurs, consult a medic.
•	Metal catches fire	Smother with clean, dry sand.
•	Spilt on the floor, bench, etc	Scoop up as much metal as possible into a dry container. Wipe the area with a damp cloth which (for calcium) should then be placed in a bucket of water.

Sulfur & Phosphorus

Substance	Hazard	Comment
Sulfur Solid	!	WARNING: Causes skin irritation. Some suppliers may also classify it as a flammable solid. Under the <i>Explosives Regulations</i> it is illegal to make mixtures with potassium chlorate(V) or other chlorates, without the prior approval of the Health & Safety Executive.
	IRRITANT	Yellow crystals of sulfur occur in volcanic regions. In Victorian times, children were fed a mixture of brimstone (sulfur) and treacle, to do them good! Sulfur burns to form sulfur dioxide gas (TOXIC). See CLEAPSS <i>Student Safety Sheet 52</i> . Asthmatics are particularly vulnerable.
		When melting sulfur or heating it, eg, with iron, insert a plug of mineral wool in the mouth of the test tube to prevent sulfur vapour escaping and igniting.
Phosphorus (red) solid	FLAMMABLE	DANGER: Flammable solid; harmful to aquatic life with long-lasting effects. Under the <i>Explosives Regulations</i> it is illegal to make mixtures with potassium chlorate(V) or other chlorates, without the prior approval of the Health & Safety Executive. May be explosive when mixed with oxidising substances. It has been used in the heads of some matches.
Phosphorus (yellow/white) solid	FLAMMABLE TOXIC CORROSIVE ENVIR.	DANGER: catches fire spontaneously if exposed to air; fatal if swallowed or inhaled; causes severe skin burns and eye damage; very toxic to aquatic life. For a 15-minute exposure, the concentration in the atmosphere should not exceed 0.3 mg m ⁻³ . Under the <i>Explosives Regulations</i> it is illegal to make mixtures with potassium chlorate(V) or other chlorates, without the prior approval of the Health & Safety Executive. It has a long history of causing poisoning, eg, amongst workers using phosphorus to make matches. When handling it, have copper(II) sulfate(VI) solution (0.2 M to 0.5 M) available to remove specks on the skin, clothing, bench, etc.
		It is used in incendiary bombs. When it burns, corrosive fumes are formed. Phosphorus fires are difficult to extinguish; smother with dry sand. It must be stored under water (or under an inert gas). When cutting phosphorus, do this under water, otherwise friction ignites it. It is hard to cut; do this in a strong container, eg, a mortar or plastic bowl – <i>not</i> glass, which is too fragile.

Typical control measures to reduce risk

- Wear eye protection when handling phosphorus or when heating or burning sulfur. Use small amounts.
- Avoid breathing fumes of sulfur dioxide, eg, use a fume cupboard or prevent sulfur vapour from igniting by using a mineral-wool plug in the mouth of a test tube.
- Wear protective gloves if handling yellow/white phosphorus; store and handle it in the absence of air.

Assessing the risks

- What are the details of the activity to be undertaken? What are the hazards?
- What is the chance of something going wrong? eg, could sulfur vapour form? Could it ignite?
- How serious would it be if something did go wrong? eg, could people be exposed to sulfur dioxide gas?
- How can the risk(s) be controlled for this activity? eg, can it be done safely? Does the procedure need to be altered? Should goggles or safety spectacles be worn?

Emera	ency	/ action
~		

•	In the eye	Flood the eye with gently running tap water for 10 minutes. Consult a medic unless only a small amount of sulfur is involved.
•	Vapour breathed in	Remove the casualty to fresh air. Consult a medic if breathing is even slightly affected.
•	Swallowed	Do no more than wash out the mouth with drinking water. Do not induce vomiting. Consult a medic.
•	Spilt on the floor, bench, etc	For sulfur or red phosphorus, brush up. For yellow/white phosphorus, cover with sand to prevent ignition. Soak in copper sulfate solution until there is no further reaction, then brush up.

M

Student safety sheets

Vocabulary

Very tiny droplets of liquid floating in a gas (usually air). Diseases are often spread by coughing **Aerosol**

and sneezing, which results in an aerosol containing microorganisms.

Some people are allergic to particular substances. Their bodies' immune system reacts to these Allergy

substances to an unusual extent, resulting in skin rashes, runny noses, wheezing or even shock.

Asthma One type of allergy, which results in breathing difficulties.

The temperature at or above which vapour from a liquid will inflame spontaneously in the **Auto-ignition**

presence of air. point

May cause disease or harm in humans. Biohazard

Carcinogen A substance which may cause cancer, if breathed in, swallowed or absorbed via the skin.

Corrosive A substance which may destroy living tissue, causing burns.

COSHH Control of Substances Hazardous to Health Regulations which aim to limit the exposure of

employees and others in the workplace to hazardous substances which may damage their health.

EHT Extra High Tension. See HT.

Explosive Substances that may explode as a result of heat, fire, friction or shock.

This includes safety spectacles, goggles and face shields. Goggles (or face shields) give better Eye protection

protection against chemical splashes and should always be used for anything which is CORROSIVE or

Flammables Substances which burn easily. There are three groups: EXTREMELY FLAMMABLE, HIGHLY FLAMMABLE and

FLAMMABLE, depending on how easily a liquid vapourises.

Flash point The lowest temperature at which a liquid gives off vapour at the surface in sufficient quantity to

ignite with air when a spark or flame is applied.

GHS Globally harmonised system (of hazard symbols and safety information). See CLEAPSS Student

Safety Sheet 91.

Harmful Similar to the effects of toxic substances, but larger quantities are needed to produce an effect. High Tension (high voltage); power packs used in schools may deliver more than 30 V at a current HT

of more than 5 mA. These are less safe than EHT power packs, which have an output of up to 5 or

6 kV but the current is limited to 5 mA or less.

Radiation which, when absorbed by a substance, including living tissue, causes some of the **Ionising radiation**

molecules to turn into ions.

Irritant A substance (including a dust) which behaves in a similar way to corrosives but, instead of

> destroying living tissue, causes significant inflammation (reddening) through immediate, prolonged or repeated contact with the skin or a mucous membrane (eg, eyes, lungs, etc). Molarity - a way of measuring concentration. More correctly written as mol dm⁻³ or moles per

litre. An ammonia solution which is 2 M has twice as many molecules of ammonia dissolved in the

water as a solution which is 1 M.

mg m⁻³ Concentrations of pollutant gases in air are sometimes measured as x milligrams of pollutant per

cubic metre of air. An alternative unit is ppm, parts per million.

When clearing up spills of chemicals it is often useful to soak them up on something inert Mineral

(unreactive). Sand can be used, but other substances soak up more, eg, clay in the form of some absorbent

types of grey 'cat litter' (familiar to those who keep cats as pets).

Mutagen A substance which may cause genetic defects.

Oxidising agent A substance which helps other substances to burn or explode.

An organism which causes disease. **Pathogen**

A further exposure to a sensitising substance will produce an unusually severe reaction, even Sensitising

when the dose or the exposure time is less than the first exposure.

Toxic A substance which, in very small quantities, may cause death or damage to health when breathed

in, swallowed or absorbed via the skin.

A teratogen is any medication, chemical, infectious disease or environmental agent that might **Teratogen**

interfere with the normal development of a fetus and result in the loss of a pregnancy, a birth

defect or a pregnancy complication.

v/v A crude measure of concentration. A 10% v/v methanol solution contains 10 ml of methanol in

w/v A crude measure of concentration. A 10% w/v sodium chlorate(I) solution contains 10 grams of

sodium chlorate(I) in 100 ml water.

Chemical safety symbols

Containers of hazardous chemicals must be labelled according to the 'globally harmonized system' (GHS). The diamond-shaped symbol will be accompanied by

- a signal word (DANGER, WARNING or nothing),
- up to 6 hazard statements (H-statements, explaining the nature of the hazard), and
- a number of precautionary statements (giving advice on handling the chemical).

Symbol	Official name	Meaning	Some examples you might find in schools
	GHS01	EXPLOSIVE	Tollen's Reagent (ammoniacal silver nitrate) if allowed to stand.
	GHS02	FLAMMABLE	Zinc and aluminium dust, hydrogen, ethanol, methanol, propanone, sodium.
	GHS03	OXIDISING	Potassium manganate(VII) solid, ammonium nitrate solid, oxygen gas, nitrogen monoxide/dioxide, chlorine.
\Diamond	GHS04	GAS UNDER PRESSURE	Hydrogen, oxygen.
	GHS05	CORROSIVE	Concentrated acids, some dilute acids (depending on concentration), concentrated alkalies, some dilute alkalies (depending on the concentration), sulfur dioxide gas, nitrogen monoxide/dioxide, sodium.
	GHS06	(ACUTELY) TOXIC	Solid barium chloride, most mercury compounds, sulfur dioxide gas, nitrogen monoxide/dioxide, chlorine, methanol.
	GHS07	MODERATE HAZARD (eg, harmful if inhaled or in contact with skin, causes eye irritation)	Some dilute acids or alkalies (depending on concentration), iodine solid and concentrated solutions, propanone.
	GHS08	HEALTH HAZARD (eg, sensitisers, carcinogens)	Most lead compounds and their solutions, most chromates and dichromates, dichloromethane, methanol.
	GHS09	ENVIRONMENTAL HAZARD	Most copper, mercury and lead compounds, and chromates and dichromates.

The chemical hazard symbols below are no longer used but may still be found on some old containers.



Note: some of these symbols (eg TOXIC), inside a yellow triangle, can be found where chemical containers are not involved. See CLEAPSS Students Safety Sheets 91b.



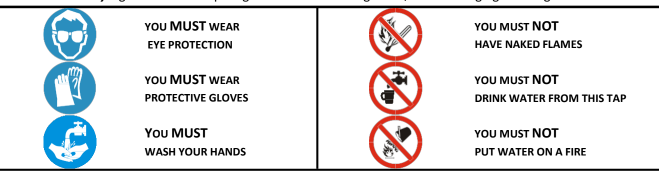
91b

Non-chemical safety signs and symbols

The Health & Safety (Safety Signs & Signals) Regulations define what the law requires. There is some additional information in the British/International Standard, BS ISO 7010:2011, Graphical symbols – Safety colours & safety signs - Registered Safety Signs.

Safety signs which are circular are mandatory (you MUST) or prohibitory (you MUST NOT).

- **Mandatory** signs have a white pictogram on a blue background;
- Prohibitory signs have a black pictogram on a white background, with red edging and diagonal line.



If there is no standard sign available you can design your own but it must conform to the general principles and must have a sign board that explains it in words.



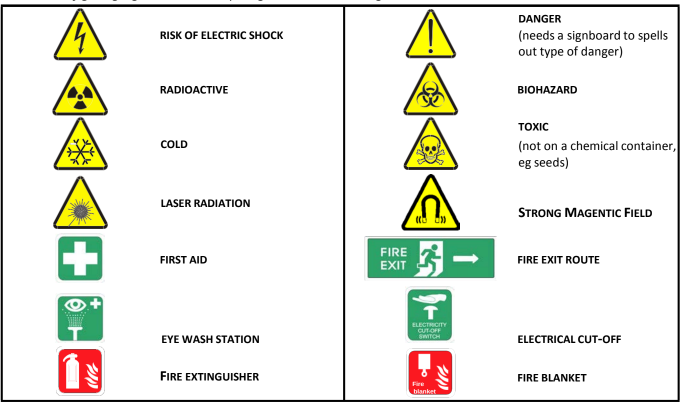
You MUST do this ...



You must NOT do this ...

Other signs give advice or information.

- Hazard information signs are triangular and have a yellow background
- **Emergency** and **first aid** signs have a white pictogram on a *green* rectangle
- Fire-fighting signs have a white pictogram on a red rectangle



CLEAPSS

Student safety sheets

Using a Bunsen burner

For heating non-flammable liquids and solids, see *CLEAPSS Student Safety Sheet 93*.

For heating flammable liquids and solids, see *CLEAPSS Student Safety Sheet 94*.

For handling hot liquids in beakers, see *CLEAPSS Student Safety Sheet 95*.

Using a Bunsen burner

- Tie any long hair back to avoid it catching fire.
- Make sure your clothing does not get in the way of the flame and don't lean over a flame to reach other apparatus.
- Wear eye protection.
- Place the Bunsen burner on a heat-resistant mat 30 to 40 cm from the edge of the bench.
- Make sure the air hole is closed (*unless* you are lighting the gas with a piezo-electric lighter, in which case air hole should be half open).
- Attach the gas tubing to the gas tap.
- Light a match or a wooden splint from a central flame. (Do not walk around the laboratory with wooden splint alight.)
- Turn on the gas tap.
- Keeping your head well away from the Bunsen burner, use the match or lighted splint to light the gas coming up the Bunsen burner chimney.
- Do not allow solids to drop into the Bunsen burner to clog the gas jet or the collar at the top of the chimney.

When you have finished:

- Make sure the air is closed and the flame is yellow.
- Switch the gas off.
- Remove the tubing by putting your fingers on the tubing around the gas tap nozzle and pulling. Don't pull the tubing off by stretching it.

Types of flame from a Bunsen burner

Type of flame	Gas tap	Air hole	Appearance	Hazards	When used
Yellow flame	Fully (or partly) open	Closed		This flame is still hot if you put your hand into it.	To light the burner and when it is not being used to heat anything because is easy to see and will not readily set fire to clothing etc. This flame is unsuitable for heating as it coats surfaces with soot (carbon).
Gentle flame Yellow just disappears; pale lilac, almost invisible.	Half open	Partly closed		It is difficult to see the flame in bright sunlight. Some risk of blowing out.	For general heating.
Medium flame Yellow just disappears; pale lilac, almost invisible.	Fully open	Partly closed		It is difficult to see the flame in bright sunlight. Some risk of blowing out.	Start with the gentle flame and then open the gas tap fully to give the medium flame.
Roaring flame Inner cone blue, outer cone lilac.	Fully open	Fully open		It is difficult to see the flame in bright sunlight. Some risk of blowing out. Do not partly close the gas tap if the air hole is fully open as this will extinguish the flame.	For very strong heating (which is not very often needed). The hottest section of the flame is just above the blue cone of unburnt gas.

Heating non-flammable liquids and solids in test tubes

For using Bunsen burners and especially use of different flames, see CLEAPSS Student Safety Sheet 92. For heating flammable liquids and solids, see CLEAPSS Student Safety Sheet 94. For handling hot liquids in beakers, see CLEAPSS Student Safety Sheet 95.

Test tube holders

- Test tube holders (if of the metal wire design shown right) must have the O-ring in the correct place, so that the clamping ends cannot slip apart.
 - (The test tube holder shown on the right does not have the O-ring in the correct position and should not be used).
- Wooden peg-type test tube holders are also suitable if they are not badly burnt.
- Tongs should not be used for holding test tubes, they are intended for crucibles.



Heating non-flammable solids in test tubes

- Wear eye protection.
- Do not have the test tube more than one-fifth full of solid.
- If the solid is a powder, shake it so that it is sloped in the test tube.
- Using a suitable holder, which is in good condition, keep the test tube pointing just up from the horizontal.
- Take care that the test tube is not pointing directly at yourself or anybody else.
- Hold the test tube so that the bottom is just in the tip of the flame and the top is well clear of the flame.
- Start with a gentle flame (see CLEAPSS Student Safety Sheet 92), increase to a medium flame and then a roaring flame, if necessary.

Heating non-flammable liquids in test tubes

- Wear eye protection.
- If possible, use a wide-diameter test tube (usually called a boiling tube).
- Do not have the test tube more than one-tenth full (it is much less likely to boil over).
- Add an anti-bumping granule ('boiling chip') before starting to heat (but never add one when it is already warm, which might result in it frothing up).
- Using a suitable holder, which is in good condition, keep the test tube at
- Take care that the test tube is not pointing directly at yourself or anybody else.
- Hold the test tube so that the bottom is just at the tip of the flame and the top is well clear of the flame.
- Start with a gentle flame (see CLEAPSS Student Safety Sheet 92), and increase it only if necessary.
- Flick your wrist continuously but gently, so that the liquid is shaken all the time it is being heated.





Heating flammable liquids and solids in test

For using Bunsen burners and especially use of different flames, see CLEAPSS Student Safety Sheet 92. For heating non-flammable liquids and solids, see CLEAPSS Student Safety Sheet 93. For handling hot liquids in beakers, see CLEAPSS Student Safety Sheet 95.

Heating flammable liquids

For example, when extracting chlorophyll from leaves.

- Wear eye protection.
- If possible, use a wide-diameter test tube (boiling tube).
- Add an anti-bumping granule ('boiling chip') before starting to heat.
- Do not heat directly over a naked flame. Instead, stand the test tube in a beaker halffilled with freshly-boiled water from a kettle.
- (If necessary, boil the water in the beaker using a roaring flame; then extinguish the flame before the flammable liquid is put into the hot water. Also, see below)
- It is always better to allow the apparatus to cool down before removing the test tube but this can take a long time.
- If necessary, use a test tube holder or a piece of folded paper carefully placed around the top of the test tube, and slowly and carefully lift the test tube out of the hot water into a test tube rack.



Heating flammable (organic) solids

For example, heating wax or salol for cooling curves.

Do not heat the wax or salol directly over a naked flame. The tube will be heated quickly and locally to a very high temperature. If a 'spirit' thermometer is used, the spirit vaporises and the glass explodes under the increase in pressure. In addition, the boiling tube may break and the contents catch fire.

- Wear eye protection.
- Use a test tube or boiling tube. Test tubes contain less solid and so take a shorter time for the contents to melt and cool. (It took 6 -7 minutes for the wax to melt in the boiling tube on the right.)
- Do not have the test tube more than half full of wax or salol.
- Fill a 250 ml beaker half full with hot water from a kettle.
- (If necessary, boil the water in the beaker using a roaring flame; then extinguish the flame before the flammable solid is put into the hot water. Also see below.)
- After the activity is completed allow the equipment to cool as much as possible.
- If necessary, use a test tube holder or a piece of folded paper carefully placed around the top of the test tube, and slowly and carefully lift the test tube out of the hot water into a test tube rack.



If you have to heat the water with a Bunsen burner (ie, not a kettle)

- At the end of the activity, first get the beaker onto the heatresistant mat, (see CLEAPSS Student Safety Sheet 95).
- Then remove the boiling tube using a good test tube holder, as described above.

This method of heating takes about the same time to melt the wax as using water boiled in a kettle and so offers no real advantage.

Note that a thermometer standing in a beaker on a tripod has a high risk of being knocked over.





Handling hot liquids in beakers

For using Bunsen burners and especially use of different flames, see CLEAPSS Student Safety Sheet 92. For heating non-flammable liquids and solids, see CLEAPSS Student Safety Sheet 93. For heating flammable liquids and solids in test tubes, see CLEAPSS Student Safety Sheet 94.

Heating liquids in beakers

- Wear eye protection.
- Place the Bunsen burner well back from the edge of the bench.
- Do not fill the beaker more than half full.
- If available, add an anti-bumping granule ('boiling chip') to the liquid before heating. Do not add an anti-bumping granule once the liquid is boiling.
- Place a tripod and gauze on a heat resistant mat.
- Place the Bunsen burner underneath and attach it to the gas.
- Place the beaker of water on the gauze.
- Light the Bunsen burner and open the air hole.
- Once the liquid begins to boil close the air hole to half open and turn the gas tap down.
- At the end it is always better to allow the apparatus to cool down naturally, but this can take a long time. If time is short use one of the procedures described below.
- The tripod will be very hot. Take care.

Removing hot beakers

- To remove very hot beakers of liquid from a tripod the teacher (or a technician) should wear thermal protection gloves and go around the laboratory moving the beakers for each group of students.
- If this is not possible, beakers are best left where they are until cooler.



For less hot beakers, skilled students can use the procedure illustrated below

- Switch off the gas.
- Place a commercial 'J-cloth' loosely around the hot beaker.
- Carefully tighten the cloth.
- Lift the beaker onto the heat resistant mat.









Risk assessment

What is risk assessment?

A risk assessment is a judgment of how likely it is that someone (anyone) might come to harm if a planned action is carried out. The law requires the likelihood of harm to be reduced to as low as is reasonably practicable. Risk assessments, although an excellent idea for all of us, are only *legally* required for actions which take place at work. The *significant findings* of risk assessment must be recorded. (You must show the answer, you don't have to show your workings).

You carry out risk assessments all the time, for example, when riding a bike or crossing the road. When riding a bike in the UK you can choose whether or not to wear a cycling helmet. However, because risk assessments are required at work, paid cycling couriers will wear helmets. The risk of them being knocked off their bike is quite high but a helmet reduces the likelihood of head injury.

Who is responsible for risk assessment?

In law, risk assessment is the responsibility of the employer. The employer can ask employees to assess risks, as long as they have been trained, but the employer must then check from time to time that it is being carried out correctly. Students can be asked to draw up a risk assessment as part of their education, but it must always be checked by teachers before being put into effect. Everybody in a work situation must, by law, take care for their own safety and that of other people and employees must do what their employer requires on health & safety matters.

Model risk assessments

In schools and colleges the employer usually makes use of model (or general) risk assessments written for them by national organisations (such as CLEAPSS). Model risk assessments give sufficient details of a procedure to enable it to be carried out safely – equipment, amounts, safety precautions, etc. However teachers (or other employees) must consider whether the model assessment needs to be adapted slightly to the particular circumstances of their own situation, eg the nature of the building or equipment used, the proximity of other students, etc. An activity considered suitable, in the model risk assessment, to be carried out in the open laboratory might not be suitable in a laboratory with poor ventilation.

How do you 'do' a risk assessment?

To make a risk assessment you need to know the hazards and the risk of them causing harm in the planned activity.

A **hazard** is anything which could cause harm. For example some chemicals, electricity at high enough currents, glass (if it breaks) and even you running in the corridor are all hazards because they can all cause harm. Although sometimes you can use your common sense to identify a hazard, often you will need specialist information, eg as provided on *CLEAPSS Student Safety Sheets* or on chemical suppliers' *Safety Data Sheets*.

The risk is the likelihood that a hazard would cause significant harm. It is a matter of judgment and depends on:

- how likely it is that something would go wrong with this hazard;
- how serious any resulting injuries would be; and
- how many people would be affected.

To reduce the risks to an acceptable level, we put in place relevant **control measures**. These are the safety precautions used to reduce the risk of harm. In science we often wear safety spectacles, or use fume cupboards. We also minimise the quantities of materials used and the concentration of hazardous solutions.

What should you do when making a risk assessment?

When making a risk assessment, go through the following process.

- 1. Consider what materials you are working with and what procedures you are you following. You could list them on the CLEAPSS *Student Form for Assessing Risk*. Think about microorganisms, heavy weights, electricity, chemicals (how much of each, what concentration of solutions), hot objects. You should also try to find out if there are any hazardous materials produced by your procedure you may need to ask your teacher!
- 2. For each of the materials and procedures, ask what are the hazards? Add them to your list. What could possibly go wrong? Look up the materials and procedures in reliable and relevant sources, eg, *CLEAPSS Student Safety Sheets*.
- 3. How many people could be affected if it went wrong? Who would they be?
- 4. What control measures (safety precautions) would you adopt? Check relevant CLEAPSS Student Safety Sheets.
- 5. Make sure you record anything important and especially the control measures.
- 6. Have the result of the risk assessment checked by your teacher before you carry on.

Science Safety Certificate

Name	For	m
I can carry out the following with d	ue regard to safety.	
Behave sensibly during practical sessions	Teacher's signature	Date
Wear safety spectacles or goggles when appropriate		
Recognise and understand the hazard symbols		
Control a Bunsen burner		
Heat a liquid in a boiling tube		
Heat a liquid in a beaker using a tripod and gauze		
Locate the position of the eye wash in the laboratory		
Measure and pour dilute acids		
Deal with an acid spill		
Check that mains plugs have the correct fuse & are correctly wired		
Explain why each of the Lab Safety Rules is needed		



Checked by:

Student safety sheets

Student form for assessing risks

Date:

Name / anti-		_			
Class / set:	••••••	Da	Date:		
dazardous chemical or nicroorganism being used or nade, or hazardous procedure or equipment	Nature of the hazard(s)	Sources(s) of information	Control measures to reduce risk		

Student form for assessing risks

Class / set:	Date:
Name(s) of pupil(s) completing form:	
Proposed practical activity: Making copper sulfate crystals from copper o	xide or copper carbonate

Hazardous chemical or microorganism being used or made, or hazardous procedure or equipment	Nature of the hazard(s)	Sources(s) of information	Control measures to reduce risk
(1) Sulfuric acid	(1) Acid is corrosive if 1.5 M or more; irritant if 0.5 M or more.	(1) Bottle label; CLEAPSS Student Safety Sheets.	(1) Use lowest possible concentration, 0.5 M; wear eye protection.
(2) Copper carbonate	(2) (a) The solid is harmful if swallowed and dust irritates lungs and eyes.(b) When the reaction takes place, tiny bubbles of carbon dioxide are formed which may produce a spray of sulphuric acid as they burst.	(2) (a) Bottle label; CLEAPSS Student Safety Sheets. (b) Text book; teacher.	(2) (a) Avoid raising dust; wear eye protection. (b) Keep face well away from reaction; wear eye protection.
(3) Copper oxide	(3) (a) The solid is harmful if swallowed and dust irritates lungs and causes serious damage to eyes. (b) Unlike copper carbonate, copper oxide needs to be heated so mixture may boil over, spill hot acid, etc. (c) Hot tripods, etc.	(3) (a) Bottle label, CLEAPSS Student Safety Sheets. (b) Teacher; text book. (c) Teacher; past experience.	(3) (a) Wear eye protection, (b) Control Bunsen-burner flame; stir to speed dissolving; stand up throughout process. (c) Pay attention.
(4) Copper sulfate	(4) Solid and solutions more concentrated than 1 M are irritant and cause serious eye damage. The solid is harmful if swallowed.	(4) CLEAPSS Student Safety Sheets.	(4) Wash hands after activity; when solution is standing to crystallise label it carefully.
(5) Evaporating solution to form saturated solution	(5) (a) Solution may boil over, or start spitting when nearly saturated.(b) Hot tripods, etc.(c) Process is slow, leading to rushing at end of lesson and accidents.	(5) (a) Teacher; past experience. (b) Teacher; past experience. (c) Teacher; past experience.	5) (a) Keep careful watch over Bunsen burner. Do not evaporate too much - allow to crystallise slowly. Wear eye protection. (b) Pay attention. (c) Use small volume, so it is quicker

Checked by:	Date:
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Checked by:

Student safety sheets

Student form for assessing risks

Date:

Froposed practical activity.				
Name(s) of pupil(s) completing form:				
Class / set:	••••	Date:		
A hazard is anything which could cau	se harm, eg, a hot tripod, a cluttere	ed floor.		
A risk is the likelihood of harm actual				
		e labels on the bottles to fill in this form.		
Hazardous chemical or procedure	Type of hazard	Control measures to reduce the risk		
Hazardous Chemical or procedure	туре ој пагага	Control measures to reduce the risk		



Student form for assessing risks

Proposed practical activity:					
	Name(s) of pupil(s) completing form:				
Hazardous chemical or microorganism being used or made, or hazardous procedure or equipment	Nature of the hazard(s)	Source(s) of information	Control measures to reduce the risks	Emergency procedure	
Chacked by:					

Transferring (handling) solid chemicals

Why 'transferring' and not 'handling'

It is better to use the word 'transferring' as opposed to 'handling' because 'handling', if taken literally, means 'using your hands' (or fingers). Many chemicals are toxic, corrosive or irritant to the skin so directly 'handling' such chemicals is never a good idea.

Should I wear gloves?

The use of chemically resistant gloves should always be considered but wearing gloves reduces manual dexterity, gives rise to an environmental issue (because they do not degrade very quickly in the waste) and are expensive for the school.

More importantly, if a chemical is on the gloves, wearers may not realise it is there and so may wipe that chemical on other parts of the body (eg, by rubbing their eyes).

However, if there are cuts which cannot be covered, or other skin issues on the hand, then gloves should be worn.

For a small number of chemicals which are corrosive but do not immediately produce a sensation of burning (eg, phenol) they should also be worn.

Transferring solids

When transferring solid chemicals, the main risk is of spilling a hazardous solid.

A spatula is the correct tool for transferring solids from one container to another. Always check that the spatula is clean, and do not use the same spatula for different solids, unless cleaned in water and dried. Spatulas come in various shapes and sizes so beware of instructions that simply say "use a spatula-full".

Nuffield spatula

This is the most common spatula used by students in schools, about 14 cm long. But with either end available for use, care must be taken in using the same end. The spatula must be cleaned and dried before changing to another chemical. The 'curved' end is the most suitable to use.



Chattaway spatula

This is available in different sizes. Students should use the small size version, about 10 cm long; teachers and technicians may use larger versions, about 20 cm long, in the preparation of solutions. There are also micro-sized versions.



Trulla (trowel spatula) and Spoon Spatulas

This is usually only used by technicians or teachers for transferring larger quantities of solids.



DIY spatula - use a wooden splint

If you have used a wooden splint to transfer one solid chemical, the end can then be cut off with scissors and another chemical can be transferred with the same splint. It can be cut to a point to transfer tiny amounts of solids.

Mixing solids

The solids to be mixed should each be placed on separate pieces of paper (or in plastic weighing boats) and the solids then poured gently from one onto the other, back and forth. Repeat the pouring action about 10 times so that there is thorough mixing. This is the only safe way of preparing explosive mixtures (which might explode if stirred with a metal spatula) and is good practice for all solid mixtures.

Transferring (handling) liquid chemicals

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Should I wear gloves?

The use of chemically resistant gloves should always be considered but wearing gloves reduces manual dexterity, gives rise to an environmental issue (because they do not degrade very quickly in the waste) and are expensive for the school.

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However, if there are cuts which cannot be covered, or other skin issues on the hand, then gloves should be worn.

For a small number of chemicals which are corrosive but do not immediately produce a sensation of burning (eg, phenol) they should also be worn.

Transferring liquids

When transferring chemicals which are liquid, the main risk is of spilling or splashing a hazardous liquid or a hazardous aqueous solution.

Spills occur when people attempt to pour liquids from a large container (bottle) into a small container such as a test tube. There is a risk of liquids dribbling down the side of the bottle or measuring cylinders, possibly damaging labels, making the bottle unsafe to pick up for the unwary, or producing fumes in the store as the liquid evaporates. It is better to pour from large bottles into (labelled) beakers first.

Pouring from bottles, measuring cylinders and beakers

One way of avoiding dribbles is to pour down a glass rod into a container via a funnel.

Small volumes of liquid (up to 3 ml)

Use plastic teat pipettes or dropping bottles.





Using automatic or volumetric pipettes

More sophisticated pipettes are available for other purposes.

Mixing liquids

Stirring

Spatulas should not be used for stirring. Stirring rods made of glass or plastic should be used. Over enthusiastic stirring can cause a glass stirrer or container to break or the liquid to splash out. Some laboratories now have magnetic stirrers.

Filling a test tube

If using a test tube, do not fill it more than one-fifth full. To mix the contents, 'waggle' the test tube from side to side. Do not shake it up and down, especially not with a thumb over the end.

Waste disposal

Avoid waste

If you carry out a practical activity in a school or college you may end up with waste. Waste is anything you don't want. The best way of dealing with waste is not to create it in the first place. Take care not to contaminate a large stock bottle, for example, by putting a dirty spatula or teat pipette into it. Plan the activity so as to minimise the amount of waste you need to deal with. If there are two alternative methods, which one generates the least waste? Microscale chemistry uses much smaller quantities than traditional methods, so generates much less waste.

Waste disposal in general

Waste disposal is tightly, and expensively, regulated. Legislation distinguishes between many different categories of waste. This sheet refers mainly to the waste that might be generated in student practical work.

Recycle or re-use

Before disposing of waste, consider whether it can be recycled or re-used. Can the product you have made in this chemical reaction, eg copper sulfate, be used for some further purpose? Can an impure solvent be re-distilled, safely, to purify it?

Solids, including solid non-hazardous chemicals

Solids which are not hazardous, and not separately classified, including non-hazardous solid chemicals, eg calcium carbonate, can be placed in the solid refuse collection. You have a duty of care to those handling the waste later on, so, for example, broken glass (if there is no separate glass collection) should be wrapped and labelled to minimise the risk of injury.

Water-soluble chemicals

If a chemical dissolves in water, and is not classed as hazardous, eg sodium chloride, it can be dissolved in water and poured down the drain as effluent. Small amounts of some hazardous chemicals can be disposed of in the same way, providing the concentration is below a threshold – the limit depends on the nature of the hazard. For some chemicals, eg copper sulfate, a 10% solution is OK (ie, not more than 10 g of the chemical in 100 cm³ of the solution), but for others, eg potassium dichromate, the maximum concentration is 0.1% and some are completely prohibited, eg mercury compounds. It is considered prudent to react acids or alkalis so that the effluent is roughly neutral. Similarly, react oxidising agents with reducing agents before flushing away.

Hazardous solid chemicals

Hazardous chemicals which do not dissolve in water, eg lead oxide, or which do dissolve but are too hazardous to dispose of in this way, eg many pesticides, must be collected by a Licensed Waste Carrier for safe processing.

Non-aqueous liquid chemicals

Chemicals which are liquids which do not mix with water, eg paraffin or solvents for cleaning paint brushes, must be collected by a Licensed Waste Carrier for safe processing. Cooking oil, even although it is not regarded as hazardous, must **not** go down the drain because it can result in fatbergs which block sewers.

Biological waste

Much biological waste (ie, plant materials) can be treated as solid waste, although ideally it should be composted. Left-overs from dissections must NOT enter the food waste system. It should be wrapped in newspaper and placed in the main non-recyclable waste.

Microbiological waste

Microbiological waste needs to be sterilized before disposal, usually by autoclaving. Very occasionally suitable disinfectants may also be acceptable.

Gases

Discharging gases to the atmosphere is regarded as pollution but small amounts of most gases can be discharged from fume cupboards to the atmosphere as long as they are not in quantities which would allow them to become a risk or a nuisance (eg, causing an unpleasant smell) to neighbours.

Cells and batteries

Dead cells and batteries of all types must not be disposed of as solid waste. Collection points can be found at civic amenity sites and at shops selling batteries. May have the 'do not discard' symbol (right).

Waste electrical and electronic equipment

No electrical and electronic equipment - from vacuum cleaners to light bulbs, from mobile phones to ammeters - can be disposed of as solid waste but must be collected separately, either by licensed waste carriers (for businesses, including schools) or taken to a civic amenity site (usually only domestic users). Modern items should have the 'do not discard' symbol (right).



Radioactive substances

Some low-level radioactive materials can be disposed of as solid waste or effluent, as above, but legislation is very detailed and a licensed waste carrier is usually needed.

CLEAPSS

Student safety sheets

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