

Ex: valid control experiment

Variable type	Identification	
Independent Variable	Temperature of oven regulated at 70°C, 80°C, 90°C, 100°C and 110°C	
Dependent Variable	Concentration of mineral using Hanna Instruments Agriculture Test Kit	
Controlled Variable	Soil samples were taken from the same site	 Ex: Methodology considers factor
	Duration of burning (24 hours)	that may influence reliability of dat
	Origin of soil from one area locally	,
	Ratio of water to soil in sedimentation (120:960)	
	Time allowed for soil particles to settle (24 hours)	
	Soil dried before use. The water content may affect the pH and mineral content in	
	measurement particularly of the unheated control.	
	Mass of soil used (120g)	
	Time for colour complex to development (30 seconds in mineral tests, 5 minutes for pH	
	tests).	
	Volume of extract used after sedimentation (2.5cm ³)	
	Amount of universal indicator (3-5 drops)	
	Mass of reagent used (pre-weighed sachets provided in the kits)	
Jncontrolled Variable	The amount of natural sunlight received both before the soil was obtained and during	Ev : Reliability of data considered
	the experiment. Soil areas exposed to sunlight may have more original minerals in soil,	
	as opposed to soil sourced from areas covered by shade because of greater plant	
	growth.	
	Original mineral content in soil This will vary with the area and deepness in soil thought	
	the sampling was consistent and the site carefully selected.	
	Porosity this may affect the efficiency of mineral transfer	
	Quantity of organic matter soil samples may have varied	

Control Experiment: The control experiment incorporated into the design is a condition whereby the soil is sourced directly from the ground and tested for pH, phosphorus, nitrogen and potassium without introducing the independent variable of heat. This control will allow for the comparison between normal soil content and subsequent mineral content to increased heat exposure.

Materials

 Table 2: Apparatus required for experiment with associated quantity

	Apparatus required for experiment with associated quantity	
Quantity	Apparatus	
1	Industrial Oven (110, 100, 90, 80, and 70 degrees C)	
1	Electronic Scales +/- 0.001	
1	Hanna Technology - HI 3895 Agriculture Test Kit (Chemicals listed in table below) Colorimetric reference cards, transfer pipette, volumetric reference card, 4 test tubes.	
1	Stopwatch	-
320	Alfisol Soil	Comm : units required
5	Large Crucibles	
1	500cm ³ beaker	
1	1000cm ³ beaker	
1	Protective oven mitts	
1	Sharpee© Labelling Marker	
1	Spatula	
1	Spotting glass	
1	5cm ³ micropipette	
1	Pipette pump	

Table 3: Chemicals required for experiment with associated quantity

Quantity	Chemical
5dm ³	Distilled Water
25	POTASSIUM test reagent - Tetrasodium salt dehydrate, lithium hydroxide monohydrate, sodium tetraphenyl borate
0.5	
25	NITROGEN test reagent - Citric acid monohydrate, barium sulphate
25	PHOSPHORUS test reagent - Mixture of crystalline powders
50cm ³	Universal Indicator
100g	Barium Sulphate _(s)

	SMENT and PRAC ORDER FORM	Ex: Safety factors considered
Method		Ex: The data is only semi-
Soil sample The soil was obtained from an unmanaged forested area. To iffected the mineral content, the sample was taken in a zone vas dug from the first 20cm of soil at least 20m from the trac ecent record of burning.	with no neighbouring farmland and the sample	quantitative. Ex: Relevant data. Comm :Clear, logical presentati Could be more concise. Comm: Method can be repeated
leating Preheat the industrial oven in practical set-up room by a		Ex: Outside factors controlled
temperature of 110 degrees C (or 100, 90, 80, 70°C)	· · ·	
Transfer measured mass of soil to individual crucible ar Wait until the oven has reached required temperature, t Wait 24 hours for the soil to burn		Ex: Sufficient range of temperatures
Remove the crucible and note qualitative observations	porosity, granularity etc).	
ve samples of soil were heated for each treatment. Three	samples were kept unheated as a control.	Ex: Sufficient data but the data semi-quantitative. This will
H measurement Sample 4g soil and add it to appropriate test tube from	agriculture test kit, fill with distilled water until the	weaken analysis
lower graduation mark is reached (2.5mL). Add the content of the packet labelled HI 3895-pH reag universal indicator and barium sulfate) to test tube from		
Replace the test tube cap and shake the tube gently measurement		
Allow the sample in test tube to stand for 5 minutes to e Compare the pH colour-card to the sample colour. Rec light filtered into laboratory is maximised, hold colour c	ord sample pH. For optimum measurement, ensure	
against a uniform white background.	· · · · · · · · · · · · · · · · · · ·	Ex: Good control
stimating N, P, K levels . Zero the balance and add 120g soil sample to the beak	ar	
Use the ratio of soil to distilled water (225:1800 reduced	l to 120:960).	
separation undisturbed (the clearer the extract become	s, the better the results of the mineral test)	
 Use a pipette to extract 2.5mL of clear general soil extra avoid agitation of soil sediment, squeeze the bulb of the 		
Add contents of specialised reagent for test (nitrogen - HI3895-K)		
 Replace and secure the test tube cap on the apparatus Allow the sample in test tube to stand for 30 seconds for 		
 Match the colour apparent in the sample to the reference under uniform lighting. 		
2. Record qualitative observations, record relevant data.		
nsure safety equipment is worn (laboratory overcoat and secorded for qualitative observation.	afety spectacles)Ensure photographic evidence is	
lodifications to the method ocks were removed from the measured mass in crucibles i ach trial. Rocks do not assist in transfer of minerals.	n order to maximise the amount of soil content in	
lore Universal indicator was added to some samples in ord	er to produce evidence pH of soil.	

The test tubes required shaking in order to react the reagents with the soil extract was approximate in the trials. This may have resulted in inconsistent measurements as reagents may not have associated with soil in these shaken more gently compared to those shaken more vigorously.

 Table 4: Tabulated raw data to show the levels of pH, nitrogen, phosphorus and potassium in soil after burning at varying temperatures - 70, 80, 90, 100, and 110°C

Temperatu	re / °C ±0.1°C		PH and mineral content of alfisol soil				
-		рН	Nitrogen	Phosphorus	Potassium		
Control	Trial 1	7.0	TRACE	TRACE	LOW		
	Trial 2	6.0	TRACE	TRACE	MEDIUM		
	Trial 3	7.0	TRACE	TRACE	MEDIUM		
70	Trial 1	7.0	TRACE	TRACE	LOW		
	Trial 2	7.0	TRACE	LOW	TRACE		
	Trial 3	7.0	TRACE	TRACE	LOW		
	Trial 4	7.0	TRACE	TRACE	TRACE		
	Trial 5	7.0	TRACE	TRACE	TRACE		
80	Trial 1	6.5	LOW	HIGH	HIGH		
	Trial 2	6.5	LOW	LOW	HIGH		
	Trial 3	6.5	LOW	LOW	HIGH		
	Trial 4	6.5	LOW	MEDIUM	HIGH		
	Trial 5	6.5	LOW	HIGH	MEDIUM		
90	Trial 1	6.5	MEDIUM	TRACE	LOW		
	Trial 2	6.5	MEDIUM	TRACE	LOW		
	Trial 3	6.5	MEDIUM	TRACE	LOW		
	Trial 4	6.5	MEDIUM	TRACE	LOW		
	Trial 5	6.5	MEDIUM	TRACE	LOW		
100	Trial 1	6.0	TRACE	LOW	MEDIUM		
	Trial 2	6.0	LOW	LOW	MEDIUM		
	Trial 3	6.0	LOW	TRACE	MEDIUM		
	Trial 4	6.0	TRACE	TRACE	MEDIUM		
	Trial 5	6.0	TRACE	TRACE	MEDIUM		
110	Trial 1	6.0	TRACE	TRACE	HIGH		
	Trial 2	6.0	TRACE	TRACE	HIGH		
	Trial 3	6.0	LOW	TRACE	HIGH		
	Trial 4	6.0	LOW	TRACE	HIGH		
	Trial 5	6.0	LOW	TRACE	HIGH		

Ev: limitations and sources of error discussed.

An: Selects and records raw data but will it permit coherent analysis?

Comm: Unambiguous presentation

Comm: Notation correct

Comm: Conventions respected except for precision of pH readings.

Table 5: Observations noted in the duration of the experimentation process with associated explanation and implication on results if applicable.

Observation	Explanation/ Implication
Grey soil after burning	Discolouration caused by absence of organic matter regulating soil
	content - suggests increase in silt and clay levels which limit the oxygen
	and carbon dioxide emission and absorption by soil.
Colour change was evident	The colour complex developed immediately in most trials - except
immediately - did not have to	nitrogen tests.
wait	
Sunlight changed content	During sedimentation, the 1000cm ³ beaker was placed in the sun for the
	90°C trial. This may have influenced the process of mineral separation in
	soil, as soil absorbs sunlight.
Different amounts of sediment	Vary levels of floating dead organic matter, and clay, silt etc on bottom of
	beaker. This may be the result of the source of soil - increased level of
	organic matter at levels of soil near the surface, however deeper down,
	the soil may contain more clay and rocks etc.
Some organic matter transferred	The method was to leave organic matter undisturbed when transferring
in trial	liquid soil extract, however, the micropipette had to penetrate the
	surface and matter adhered to/ transferred by the pipette to the test
	tube.

An: Qualitative observations made.

Comm: Unambiguous presentation

An : Averaging qualitative data has doubtful validity.

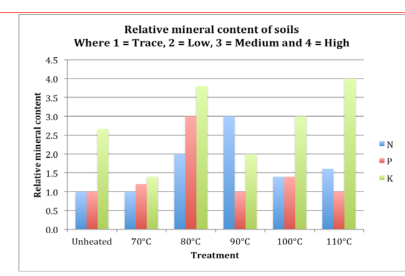
An: Uncertainties in the temperature presented but none for

the pH measurements.

Table 7: The averaged results of pH and the overall results of mineral tests at varying temperatures

Temperature / °C ±0.1	pH and minerals							
	рН	Nitrogen	Phosphorus	Potassium				
Unheated	6.7	TRACE	TRACE	MEDIUM				
70	7.0	TRACE	TRACE	TRACE				
80	6.5	LOW	HIGH	HIGH				
90	6.5	MEDIUM	TRACE	LOW				
100	6.0	TRACE	TRACE	MEDIUM				
110	6.0	LOW	TRACE	HIGH				

Graphs



An: Conversion of a qualitative scale into a numerical value is not scientifically valid. There has been no calibration of the colour charts other than ordering the results by relative importance.

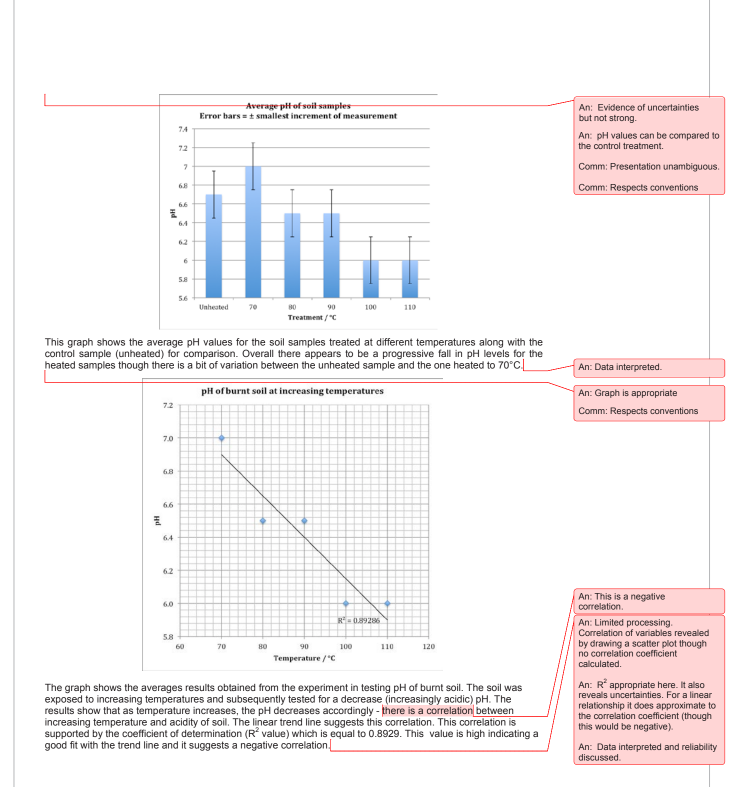
Comm: Nevertheless the bar chart is unambiguously presented and...

An: .. it permits a visual comparison between the values of the different treatments.

An: Uncertainties not relevant.

The graph shows the averaged results obtained from the experiment in testing for minerals - phosphorus, nitrogen and potassium. The graph depicts the mineral content as a result of increased temperature exposure of soil as well as the control (unheated). There appears to be a trend in increasing potassium levels as the temperature increases above 90°C, however there is no specific pattern for other tested minerals of nitrogen and phosphorus. It appears the optimum temperature to produce the highest content of phosphorus is 80°C, and the optimum for nitrogen is 90°C.

An: Interpretation is over optimistic.



Conclusion The data obtained supports the idea that the pH of the soil becomes more acidic at higher temperatures. Ev: Reasonable conclusion The data suggests that there is an increase in the mineral content with heating of alfisol soils. for which there is some evidence. Table 7, averages of the raw data, demonstrates the increase of temperature that is directly proportional to Ev: Dubious claim an increase in mineral content. At temperatures of 110°C, the potassium levels are highest. However, this trend is not present in all mineral compounds. Nitrate compounds in burnt soil are highest at 90°C. An: Over optimistic interpretation Phosphate molecules in soil are highest in at 80°C. This suggests that at certain temperatures, the mineral of the data content in soils for varying minerals changes, both in guantify and type. The general increase in mineral content is presumed to be a result of the microbe combustion in the soil. Microbes and organic matter store and regulate minerals such as nitrates and phosphates in the soil - in addition to absorption by plants and other organisms. When the soil is exposed to heat, the change in temperature combusts the organic matter, releasing the mineral and resulting in their increased levels. However, this does not suggest that plants are able to use these enriched resources. Naturally decomposer organisms, such as worms, are able to produce organic compounds and release minerals for plants. Without these decomposers to maintain the cycle, the plants will only have a certain store of minerals. Moreover, despite the increased availability of minerals, plants require other factors of soil such as porosity, which determines water retention. Soil that is subjected to extreme temperatures loses the ability to hold water for plants and organisms - which is why burnt lands are unable to sustain plant growth until revived naturally. However, an application of this is that burnt soil may be mixed in with mineral deficient soil in order to boost mineral levels in the soil for plants. This would enhance the soil chemical properties and aid growth of plants. Furthermore, alfisol soils are typically naturally deficient in minerals such as potassium and nitrogen in Australian regions. Ev: Good scientific context. The third graph depicts the change in pH as a result of increasing temperature exposure. There is a good correlation between increasing temperature and decreasing pH. This is supported by the high coefficient of determination, R² value = 0.8929. Nitrogen minerals exist as nitrates in the soil (N0₃). Since the increase in temperature increases the quantity of nitrates in the soil, the soil will acidify as a result. This is the same in phosphates (Holleman, A - Inorganic Chemistry 2001). It can be predicted that at temperatures Ev: Results compared to theory exceeding 110°C, it pH will continue to become increasingly acidic - demonstrated by the consistent linear trend line until all the organic matter is combusted. Ev: Extrapolates Solubility of minerals in soil varies with the pH of the soil. Most minerals are readily available at a pH of 6 to 7. Phosphorus is more available at pH 6.5 and below pH 5 aluminium, iron and manganese become more soluble. Aluminum is toxic for many plant species and it is particularly rich in alfisols. So the observed drop in pH with heating could result in some valuable minerals becoming more accessible to begin with but a strong drop in pH could make the soil less fertile. An: Data interpreted. Reliability The data collected is reliable with a number of minor omissions. The alfisol soil, which is typical soil family found in most areas of Australia, is known for a range of physical properties. The results cannot be generalised to all soils as the test conducted on the soil obtained may contain mineral levels different to those of an alfisol soil found in Western Australia. Moreover, it was unreliable to the extent the pH tests are variable in the controls. The water content of the controls may have influenced these results. In the heated samples the water evaporated off. Ev : Limitation of the methodology considered The original organic matter in the soil, which influence the mineral content, as well as the plant concentration in the area of source was not recorded. This would be an indication of the general nature of the results to areas containing similar quantities of organic matter. Moreover, the level of sunlight received by the areas where the soil was sampled was not controlled. The intensity of the sunlight will determine the rate of photosynthesis of plants. If the soil is in a particularly shady area, then the photosynthetic activity of plants would be lower than that of plants constantly receiving of sunlight. This would result in a lower biomass of plants and less organic matter in the soil. A decreased level of photosynthesis could result in the mineral content of the test soil being lower. Ev: Limitation of the methodology considered The experiment repeated trials for the samples which helps to reveal the reliability of the data in consistency between trials. The uncertainty associated with the reference cards means that the content of minerals in the soil solution could potentially be half a grade higher or half a grade lower of that identified. Ev: Limitation of the methodology Moreover, error in human visual analysis, is significant. The reference cards require an observation by the considered. experimenter to a standardised measurement, whereas a measuring probe like a colorimeter may be able to analyse the concentration more objectively. Ev: Suggested improvement Page 7

Evaluation of the method

There were a number of systematic and random errors associated with the uncertainty of the data which is a limitation to the results.

A systematic error associated with the data is in reference to the colorimetric indications cards. These apparatus are non-numerical - and rely on a comparison between the solution colour as a result of the reagent, and a standardised TRACE, LOW, MEDIUM or HIGH reference on the card. This is a limitation to the design whereby the data, excluding pH, cannot be converted into numerical data. Therefore the uncertainty related to the data is half of the smallest increment - for example between HIGH and MEDIUM. This is a large uncertainty as it suggests there may be a different concentration value of the mineral in the solution. A similar problem occurred with the pH tests. This method requires universal indicator in order to colour the soil and compare this subsequent colour to a pH reference chart.

A random error is that the method stated: "Allow soil to separate into components; organic matter, clay, silt, mineral content etc., for 24 hours to ensure maximum separation undisturbed (the clearer the extract becomes, the better the results of the mineral test)." However, each test was not allowed 24 hours (due to experimentation time restraints), which resulted in a yellow coloured extract, as opposed to a translucent one. This suggests that the results obtained may not have been the optimum indication of mineral content in the burnt soil. Moreover, the time for sedimentation was a maximum of 24 hours. The soil used does not give an indication of the duration of the sedimentation process needed, and therefore the range of time is 30 minutes to 24 hours. It was assumed the 24 hours would be suitable for a complete result.

The reagents used as indicators of the minerals were prepared in sachets that were opened and added to the soil aqueous extract. However, the entire quantity was not added in each case due to the difficulty of releasing the chemicals. This may result in a lighter colour of the solution, which would demonstrate a qualitative comparison that does not necessarily represent the actual quantity of mineral in the sample.

Another random error is that the oven, which controls the temperature of heating, was temporarily opened by other students and the temperature decreased from 110 to 97°C. This is an inconsistent heating process and the trials will require a repeat to ensure accurate results were opened. The temperature drop of 13°C temperature may have had and affect on the experiment.

Modifications to Experiment

To increase the accuracy of results, there is a professional version of the HI 3895 test kit used to test mineral content in the soil in this experiment. HI 3896 uses a similar methodology to the HI 3895 agriculture test kit, however uses colour compactors and assigns numerical values to the colour comparison. This will provide a significantly more measureable basis to the results.

Alternatively an ultraviolet-visible light spectrophotometer may be used to test the unknown concentration compared with a calibration curve created using standard solutions of diluted phosphate stock solution. The spectrophotometer will provide a more accurate measurement compared to the reference card. This may be applicable to other minerals such as potassium compounds and nitrate compounds.

Hanna instruments, the company that manufactured the agriculture test kit used to test the content of minerals and pH in the soil, additionally produces a product called a phosphate checker. The instrument uses a silicon photocell that works similar to a ultraviolet-visible spectrophotometer, however this tests for orthophosphates only using the reagents and no calibration is required to determine the concentration. The apparatus measurement ranges from 0.00 to 2.50ppm with an uncertainty of 4.0%. The apparatus is manufactured for nitrates and potassium compounds, in addition to phosphates.

In reference to the pH tests, a pH probe may be used to determine a more accurate measurement of the pH of the soil. The method for using this apparatus is extracting a specific mass of soil and adding distilled water. This solution may be filtered using filter paper and funnel. The pH probe, after calibrating in electrolytes and cleaning in distilled water, will measure the pH of the solution accurately.

A modification to the experiment could be the manipulating the independent variable as time rather than temperature. The experiment has demonstrated that temperature change will influence the mineral content in soil, however a derivative of the experiment could investigate the effect of time duration of burning. This would provide a more ecologically valid representation of bush fire examples - uncontrolled and controlled fires.

A further variation of the experiment may investigate the effect of lower temperatures, since the higher temperatures combusts all organic matter/large majority of organic matter, lower temperatures that will destroy only some of the organic matter.

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Ev: Reliability of data considered. Important weakness identified, but the candidate contradicts what was attempted in the analysis.

Ev: Sources of error identified.

Ev: Sources of error identified and impact considered.

Ev: Sources of error identified and impact considered.

Ev: Realistic improvements suggested

Ev: suggests feasible extension.

Table 9: Summary of modifications to experimental design

Modification	Explanation
UV-Vis Spectrophotometer	Accurate measure the concentration of phosphate ions in soil extracts
Phosphate checker	Works in similar effect to UV-Vis apparatus, does not require calibration curve - measures phosphate concentration directly.
HI 3896	Precise agriculture test kit compared to HI 3895 - assigns numerical values
Lower temperatures	Maintain higher content of minerals with regulation by remaining organic matter.
Duration of heating	Change the independent to time -> better reflects the ecological factor of fire and soil.
pH probe	Used to measure the pH of the solution - filter soil extract and measure pH
Control sample water content	Air dry the control soil at room temperature

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1	Electronic Scales 4	H- 0.00	1						
1	Hanna Technology								
1	Colorimetric refere Stopwatch	nce car	rds, transfer pip	bette, volumetri	c reference card, 4	test tube	15.		
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5	Large Crucibles								
1	500cm ³ beaker								
1	1000cm ³ beaker								
1	Protective oven mit	tts							
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Ex: Most safety factors considered except for the chemicals used which would be expected in this investigation.

BIOLOGICAL	Pathogenic		Infectious		Bite/sting	4	Do not ingest the soil, may risk ingesting	3
	Micro-organism		Blood		Airborne		micro-organisms or harmful compositions.	
	Bodily fluids		Penetrating		Venom			
	Contaminant		Poisonous		Spine/ prickle			
MANUAL HANDLING	Environment	\boxtimes	Repetition		Special needs	4	When using oven, maintain safe distance from	3
	Design		Force		Stability		oven so as to avoid burning.	
	Shape		Posture	\boxtimes	Temperature			
	U Weight		Personal Factors		Grip			
ELECTRICAL	Burns		Current		Shock	4	Oven reaches temperatures of 200°, therefore	3
	Cold	\boxtimes	Heat		Voltage		do not lough inside of oven. Apply cold water immediately if burn ensues.	
ENVIRONMENTAL	Moving water		Terrain		Weather	N/a	N/a	N/a
	Standing water							
MECHANICAL	🖾 Heat		Projectiles		Noise	4	Caution of hot surface in oven.	3
	Moving parts		Ventilation		Vibration			
	Vacuum		Pressure					
PENETRATING OBJECT	Blunt trauma	\boxtimes	Sharp trauma			4	Ensure if glassware breaks, do not attempt to clean up with without protection. If abrasion occurs, apply pressure and water.	3
PHYSICAL	Absorption		Exposure		Ingestion	4	Do not directly inhale dust from soil, do not	3
	Dehydration		Injury	\boxtimes	Inhalation		touch hot surfaces.	-
	Fatigue		Visual	\boxtimes	Temperature			
RADIATION	Arc flashes		Medical		Magnetic field	N/a	N/a	N/a
	Ionising		Laser		Radioactive decay			
	Microwave		Sunburn		UV			
	Infra Red		X-rays					
GAS	E Burns		Inhalation		Bunsen hose	N/a	N/a	N/a
	Bunsen hose co	ontact	with fiame		secured onto lurret			
EXPLOSION	Noise		Pressure		Exposure	N/a	N/a	N/a
	Fire		Explosion					
OTHER (specily)						N/a	N/a	N/a
L								

CHEMICAL REQUIRMENTS AND CHEMICAL RISK ASSESSMENT										
Chemical Name (include volume and concentration required)	Hazard Identification ²	Risk phrases (refer to MSDS)	Safety phrases (refer to MSDS)	Risk Score	What should I do? (Control measures/ Management Strategies)	New Risk Score				
N/a	N/a	N/a	N/a	N/a	N/a	N/a				

² Chemical Hazard Identification – refer to MSDS											
Hazardous	Initant	Taxic	Corrosive	Flammable	Spills	Body contact	Sensitisin	Mutagen	Environmental toxin	Asphyxiant	
Combustion	Explosive	Reactive	Harmful	Oxidising	Splashes	Bums	Carcinoge	Teratogenic	Hazardous by	Other	