SAFETY DATA SHEET



Regular Unleaded petrol with 10% ethanol

Section 1. Identification

GHS product identifier Regular Unleaded petrol with 10% ethanol

Other means of identification

E10, E10M

 Product code
 0000002889

 SDS no.
 0000002889

Relevant identified uses of the substance or mixture and uses advised against

Use of the substance/

mixture

Use only as a motor fuel for spark ignition engines. NOT for aviation use. Should

NOT be used as a solvent nor cleaning agent.

Manufacturer

Supplier BP Australia Pty Ltd

Level 17, 717 Bourke Street Docklands, Victoria 3008 ABN 53 004 085 616

www.bp.com.au

Technical Helpline Number: 1300 139 700

EMERGENCY TELEPHONE

NUMBER

1800 638 556

Section 2. Hazard(s) identification

Classification of the AMMABLE LIQUIDS - Category 1

substance or mixture SKIN CORROSION/IRRITATION - Category 2

SERIOUS EYE DAMAGE/EYE IRRITATION - Category 2A

GERM CELL MUTAGENICITY - Category 1B

CARCINOGENICITY - Category 1B

REPRODUCTIVE TOXICITY (Unborn child) - Category 2

SPECIFIC TARGET ORGAN TOXICITY - SINGLE EXPOSURE (Narcotic effects) -

Category 3

ASPIRATION HAZARD - Category 1

GHS label elements

Hazard pictograms







Signal word DANGER

Hazard statements H224 - Extremely flammable liquid and vapour.

H319 - Causes serious eye irritation. H315 - Causes skin irritation. H340 - May cause genetic defects.

H350 - May cause cancer.

H361 - Suspected of damaging the unborn child. H304 - May be fatal if swallowed and enters airways.

H336 - May cause drowsiness or dizziness.

Precautionary statements

General P103 - Read label before use.

P102 - Keep out of reach of children.

P101 - If medical advice is needed, have product container or label at hand.

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Section 2. Hazard(s) identification

Prevention

₱201 - Obtain special instructions before use.

P202 - Do not handle until all safety precautions have been read and understood.

P261 - Avoid breathing vapour.

P280 - Wear protective gloves. Wear eye or face protection. Wear protective

clothing.

 $\mbox{P210}$ - Keep away from heat, hot surfaces, sparks, open flames and other ignition sources. No smoking.

P241 - Use explosion-proof electrical, ventilating, lighting and all material-handling

equipment.

P273 - Avoid release to the environment.

Response P304 + P340 - IF INHALED: Remove victim to fresh air and keep at rest in a

position comfortable for breathing.

P301 + P310 + P331 - IF SWALLOWED: Immediately call a POISON CENTER or

physician. Do NOT induce vomiting.

P303 + P361 + P353 - IF ON SKIN (or hair): Take off immediately all contaminated

clothing. Rinse skin with water or shower.

Storage P403 + P233 - Store in a well-ventilated place. Keep container tightly closed.

P235 - Keep cool. P405 - Store locked up.

Disposal P501 - Dispose of contents and container in accordance with all local, regional,

national and international regulations.

Supplemental label

elements

Not applicable.

Other hazards which do not result in classification

Contains Benzene. Prolonged or repeated exposure to benzene can cause anaemia and other blood diseases, including leukaemia.

Section 3. Composition and ingredient information

Substance/mixture

Mixture

Complex mixture of volatile hydrocarbons containing paraffins, naphthenes, olefins and aromatics with carbon numbers predominantly between C4 and C12. May contain oxygenates. May also contain small quantities of proprietary performance additives. Contains: ethanol.

Ingredient name	% (w/w)	CAS number
© asoline	>90	86290-81-5
Ethanol	<10	64-17-5
Contains:		
Benzene	<1	71-43-2
tert-butyl methyl ether	<1	1634-04-4
2-methylpropan-2-ol	<1	75-65-0
diisopropyl ether	<1	108-20-3
Polycyclic aromatic hydrocarbons (PAHs)	<1	mixture

There are no additional ingredients present which, within the current knowledge of the supplier and in the concentrations applicable, are classified as hazardous to health or the environment and hence require reporting in this section.

Occupational exposure limits, if available, are listed in Section 8.

Section 4. First aid measures

Description of necessary first aid measures

Eye contact In case of contact, immediately flush eyes with plenty of water for at least 15

minutes. Check for and remove any contact lenses. Get medical attention if

irritation occurs.

or fumes causes drowsiness, headache, blurred vision or irritation of the eyes, nose or throat, remove immediately to fresh air. Keep patient warm and at rest. If any

symptoms persist obtain medical advice.

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Section 4. First aid measures

Skin contact rcase of contact, immediately flush skin with plenty of water for at least 15 minutes

while removing contaminated clothing and shoes. Drench contaminated clothing with water before removing. This is necessary to avoid the risk of sparks from static electricity that could ignite contaminated clothing. Contaminated clothing is a fire hazard. Contaminated leather, particularly footwear, must be discarded. Clean

shoes thoroughly before reuse. Get medical attention.

Do not induce vomiting. Never give anything by mouth to an unconscious person. If Ingestion

unconscious, place in recovery position and get medical attention immediately. Aspiration hazard if swallowed. Can enter lungs and cause damage. Get medical

attention immediately.

Most important symptoms/effects, acute and delayed

See Section 11 for more detailed information on health effects and symptoms.

Indication of immediate medical attention and special treatment needed, if necessary

Notes to physician Treatment should in general be symptomatic and directed to relieving any effects.

> Product can be aspirated on swallowing or following regurgitation of stomach contents, and can cause severe and potentially fatal chemical pneumonitis, which will require urgent treatment. Because of the risk of aspiration, induction of vomiting and gastric lavage should be avoided. Gastric lavage should be undertaken only

after endotracheal intubation. Monitor for cardiac dysrhythmias.

Specific treatments No specific treatment.

Protection of first-aiders No action shall be taken involving any personal risk or without suitable training. If it

is suspected that fumes are still present, the rescuer should wear an appropriate mask or self-contained breathing apparatus. It may be dangerous to the person providing aid to give mouth-to-mouth resuscitation. Wash contaminated clothing

thoroughly with water before removing it, or wear gloves.

Section 5. Firefighting measures

Extinguishing media

Suitable extinguishing media

Unsuitable extinguishing

media

In case of fire, use water fog, foam, dry chemical or carbon dioxide extinguisher or spray.

Do not use water jet.

Specific hazards arising from the chemical

Extremely flammable liquid and vapour. Fire water contaminated with this material must be contained and prevented from being discharged to any waterway, sewer or drain. In a fire or if heated, a pressure increase will occur and the container may burst, with the risk of a subsequent explosion. Runoff to sewer may create fire or explosion hazard. This product is a poor conductor of electricity and can become electrostatically charged. If sufficient charge is accumulated, ignition of flammable mixtures can occur. To reduce potential for static discharge, use proper bonding and grounding procedures. This liquid may accumulate static electricity when filling properly-grounded containers. Static accumulation may be significantly increased by the presence of small quantities of water or other contaminants. Liquid will float and may reignite on surface of water.

Hazardous thermal decomposition products combustion products may include the following:

carbon oxides (CO, CO₂) (carbon monoxide, carbon dioxide)

Special protective actions for fire-fighters

No action shall be taken involving any personal risk or without suitable training. Promptly isolate the scene by removing all persons from the vicinity of the incident if there is a fire. Move containers from fire area if this can be done without risk. Use water spray to keep fire-exposed containers cool.

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Special protective equipment for fire-fighters

Hazchem code

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Fire-fighters should wear positive pressure self-contained breathing apparatus (SCBA) and full turnout gear.

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Section 6. Accidental release measures

Personal precautions, protective equipment and emergency procedures

For non-emergency personnel

Immediately contact emergency personnel. No action shall be taken involving any personal risk or without suitable training. Evacuate surrounding areas. Keep unnecessary and unprotected personnel from entering. Do not touch or walk through spilt material. No flares, smoking or flames in hazard area. Avoid breathing vapour or mist. Provide adequate ventilation. Put on appropriate personal protective equipment. Floors may be slippery; use care to avoid falling. Eliminate all ignition sources.

For emergency responders

Entry into a confined space or poorly ventilated area contaminated with vapour, mist or fume is extremely hazardous without the correct respiratory protective equipment and a safe system of work. Wear self-contained breathing apparatus. Wear a suitable chemical protective suit. Chemical resistant boots. See also the information in "For non-emergency personnel".

Environmental precautions

Avoid dispersal of spilt material and runoff and contact with soil, waterways, drains and sewers. Inform the relevant authorities if the product has caused environmental pollution (sewers, waterways, soil or air). Water polluting material. May be harmful to the environment if released in large quantities. In case of small spillages in closed waters (i.e. ports), contain product with floating barriers or other equipment. Collect spilled product by absorbing with specific floating absorbents. If possible, large spillages in open waters should be contained with floating barriers or other mechanical means. If this is not possible, control the spreading of the spillage, and collect the product by skimming or other suitable mechanical means. The use of dispersants should be advised by an expert, and, if required, approved by local authorities. Collect recovered product and other contaminated materials in suitable tanks or containers for recycle, recovery or safe disposal.

Methods and material for containment and cleaning up

Small spill

Eliminate all ignition sources. Stop leak if without risk. Move containers from spill area. Absorb with an inert material and place in an appropriate waste disposal container. Use spark-proof tools and explosion-proof equipment. Dispose of via a licensed waste disposal contractor. The method and equipment used must be in conformance with appropriate regulations and industry practice on explosive atmospheres.

Large spill

Eliminate all ignition sources. Stop leak if without risk. Move containers from spill area. Approach the release from upwind. Prevent entry into sewers, water courses, basements or confined areas. Dike spill area and do not allow product to reach sewage system and surface or ground water. Contain and collect spillage with noncombustible, absorbent material e.g. sand, earth, vermiculite or diatomaceous earth and place in container for disposal according to local regulations. Use spark-proof tools and explosion-proof equipment. Contaminated absorbent material may pose the same hazard as the spilt product. The method and equipment used must be in conformance with appropriate regulations and industry practice on explosive atmospheres. Dispose of via a licensed waste disposal contractor.

Section 7. Handling and storage

Precautions for safe handling

Protective measures

Put on appropriate personal protective equipment (see Section 8). Avoid exposure obtain special instructions before use. Avoid exposure during pregnancy. Do not handle until all safety precautions have been read and understood. Do not get in eyes or on skin or clothing. Do not swallow. Aspiration hazard if swallowed. Can enter lungs and cause damage. Never siphon by mouth. Avoid breathing vapour or mist. Use only with adequate ventilation. Wear appropriate respirator when ventilation is inadequate. Keep in the original container or an approved alternative made from a compatible material, kept tightly closed when not in use. Store and use away from heat, sparks, open flame or any other ignition source. Use explosionproof electrical (ventilating, lighting and material handling) equipment. Use only nonsparking tools. Take precautionary measures against electrostatic discharges.

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Section 7. Handling and storage

Empty containers retain product residue and can be hazardous. Do not reuse container. Avoid contact of spilt material and runoff with soil and surface waterways. Handling operations that can promote accumulation of static charges include but are not limited to: mixing, filtering, pumping at high flow rates, splash filling, creating mists or sprays, tank and container filling, tank cleaning, sampling, gauging, switch loading, vacuum truck operations. Restrict flow velocity according to API 2003 (2008), NFPA 77 (2007), and Laurence Britton, "Avoiding Static Ignition Hazards in Chemical Operations". To reduce potential for static discharge, ensure that all equipment is properly grounded and bonded and meets appropriate electrical classification requirements.

Advice on general occupational hygiene Do not fill container while it is in or on a vehicle. Place container on ground when filling and keep nozzle in contact with container.

Eating, drinking and smoking should be prohibited in areas where this material is handled, stored and processed. Wash thoroughly after handling. Remove contaminated clothing and protective equipment before entering eating areas. See also Section 8 for additional information on hygiene measures.

Conditions for safe storage, including any incompatibilities

Store in accordance with local regulations. Store in a segregated and approved area. Store in original container protected from direct sunlight in a dry, cool and wellventilated area, away from incompatible materials (see Section 10) and food and drink. Store locked up. Eliminate all ignition sources. Separate from oxidising materials. Keep container tightly closed and sealed until ready for use. Store and use only in equipment/containers designed for use with this product. Containers that have been opened must be carefully resealed and kept upright to prevent leakage. Do not store in unlabelled containers. Use appropriate containment to avoid environmental contamination.

Light hydrocarbon vapours can build up in the headspace of tanks. These can cause flammability/explosion hazards even at temperatures below the normal flash point (note: flash point must not be regarded as a reliable indicator of the potential flammability of vapour in tank headspaces). Tank headspaces should always be regarded as potentially flammable and care should be taken to avoid static electrical discharge and all ignition sources during filling, ullaging and sampling from storage tanks. Do not enter storage tanks. If entry to vessels is necessary, follow permit to work procedures. Entry into a confined space or poorly ventilated area contaminated with vapour, mist or fume is extremely hazardous without the correct respiratory protective equipment and a safe system of work. When the product is pumped (e.g. during filling, discharge or ullaging) and when sampling, there is a risk of static discharge. Ensure equipment used is properly earthed or bonded to the tank structure. Electrical equipment should not be used unless it is intrinsically safe (i.e. will not produce sparks). Explosive air/vapour mixtures may form at ambient temperature. If product comes into contact with hot surfaces, or leaks occur from pressurised fuel pipes, the vapour or mists generated will create a flammability or explosion hazard. Product contaminated rags, paper or material used to absorb spillages, represent a fire hazard, and should not be allowed to accumulate. Dispose of safely immediately after use.

Section 8. Exposure controls and personal protection

Control parameters Occupational exposure limits

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Section 8. Exposure controls and personal protection

Ingredient name	Exposure limits
Sasoline	ACGIH TLV (United States). TWA: 300 ppm 8 hours. Issued/Revised: 5/1996 TWA: 890 mg/m³ 8 hours. Issued/Revised: 5/1996 STEL: 500 ppm 15 minutes. Issued/ Revised: 5/1996 STEL: 1480 mg/m³ 15 minutes. Issued/ Revised: 5/1996
Ethanol	Safe Work Australia (Australia). TWA: 1880 mg/m³ 8 hours. Issued/Revised: 5/1995 TWA: 1000 ppm 8 hours. Issued/Revised: 5/1995
Benzene	Safe Work Australia (Australia). TWA: 3.2 mg/m³ 8 hours. Issued/Revised: 4/2003 TWA: 1 ppm 8 hours. Issued/Revised: 4/2003
tert-butyl methyl ether	Safe Work Australia (Australia). STEL: 275 mg/m³ 15 minutes. Issued/ Revised: 4/2002 STEL: 75 ppm 15 minutes. Issued/Revised: 4/2002 TWA: 92 mg/m³ 8 hours. Issued/Revised: 4/2002 TWA: 25 ppm 8 hours. Issued/Revised: 4/2002
2-methylpropan-2-ol	Safe Work Australia (Australia). STEL: 455 mg/m³ 15 minutes. Issued/ Revised: 5/1995 STEL: 150 ppm 15 minutes. Issued/ Revised: 5/1995 TWA: 303 mg/m³ 8 hours. Issued/Revised: 5/1995 TWA: 100 ppm 8 hours. Issued/Revised: 5/1995
diisopropyl ether	Safe Work Australia (Australia). STEL: 1300 mg/m³ 15 minutes. Issued/ Revised: 5/1995 STEL: 310 ppm 15 minutes. Issued/ Revised: 5/1995 TWA: 1040 mg/m³ 8 hours. Issued/Revised: 5/1995 TWA: 250 ppm 8 hours. Issued/Revised: 5/1995
Polycyclic aromatic hydrocarbons (PAHs)	Safe Work Australia (Australia). TWA: 0.2 mg/m³ 8 hours.

Appropriate engineering controls

All activities involving chemicals should be assessed for their risks to health, to ensure exposures are adequately controlled. Personal protective equipment should only be considered after other forms of control measures (e.g. engineering controls) have been suitably evaluated. Personal protective equipment should conform to appropriate standards, be suitable for use, be kept in good condition and properly maintained.

Your supplier of personal protective equipment should be consulted for advice on selection and appropriate standards. For further information contact your national

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Section 8. Exposure controls and personal protection

organisation for standards.

Provide exhaust ventilation or other engineering controls to keep the relevant airborne concentrations below their respective occupational exposure limits. The final choice of protective equipment will depend upon a risk assessment. It is important to ensure that all items of personal protective equipment are compatible.

Environmental exposure controls

Emissions from ventilation or work process equipment should be checked to ensure they comply with the requirements of environmental protection legislation. In some cases, fume scrubbers, filters or engineering modifications to the process equipment will be necessary to reduce emissions to acceptable levels.

Individual protection measures

Hygiene measures

Wash hands, forearms and face thoroughly after handling chemical products, before eating, smoking and using the lavatory and at the end of the working period. Appropriate techniques should be used to remove potentially contaminated clothing. Wash contaminated clothing before reusing. Ensure that eyewash stations and safety showers are close to the workstation location.

Eye/face protection Skin protection Hand protection

Chemical splash goggles.

Wear chemical resistant gloves.

Do not re-use gloves. Protective gloves must give suitable protection against mechanical risks (i.e. abrasion, blade cut and puncture). Protective gloves will deteriorate over time due to physical and chemical damage. Inspect and replace gloves on a regular basis. The frequency of replacement will depend upon the circumstances of use.

Skin protection

Recommended: Gloves made from fluoroelastomer resistant to hydrocarbons and a wide range of chemicals. Nitrile gloves.

✓ se of protective clothing is good industrial practice.

Personal protective equipment for the body should be selected based on the task being performed and the risks involved and should be approved by a specialist before handling this product.

Cotton or polyester/cotton overalls will only provide protection against light superficial contamination that will not soak through to the skin. Overalls should be laundered on a regular basis. When the risk of skin exposure is high (e.g. when cleaning up spillages or if there is a risk of splashing) then chemical resistant aprons and/or impervious chemical suits and boots will be required.

Wear suitable protective clothing.

Footwear highly resistant to chemicals.

When there is a risk of ignition wear inherently fire resistant protective clothes and gloves.

When there is a risk of ignition from static electricity, wear anti-static protective clothing. For greatest effectiveness against static electricity, overalls, boots and gloves should all be anti-static.

When the risk of skin exposure is high (from experience this could apply to the following tasks: cleaning work, maintenance and service, filling and transfer, taking samples and cleaning up spillages) then a chemical protective suit and boots will be required.

Work clothing / overalls should be laundered on a regular basis. Laundering of contaminated work clothing should only be done by professional cleaners who have been told about the hazards of the contamination. Always keep contaminated work clothing away from uncontaminated work clothing and uncontaminated personal clothes.

Other skin protection

Appropriate footwear and any additional skin protection measures should be selected based on the task being performed and the risks involved and should be approved by a specialist before handling this product.

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Section 8. Exposure controls and personal protection

Respiratory protection Se with adequate ventilation.

If there is a requirement for the use of a respiratory protective device, but the use of breathing apparatus (independent of ambient atmosphere) is not required, then a

suitable filtering device must be worn.

The filter class must be suitable for the maximum contaminant concentration (gas/

vapour/aerosol/particulates) that may arise when handling the product.

Recommended: If ventilation is inadequate, use respirator that will protect against

organic vapour and dust/mist.

Refer to standards: Respiratory protection: AS/NZS 1715 and AS/NZS 1716

Gloves: AS/NZS 2161.1

Eye protection: AS/NZS 1336 and AS/NZS 1337

Section 9. Physical and chemical properties

Appearance

Physical stateLiquid. Clear and BrightColourPale Yellow.to Orange/Red.OdourHydrocarbon. [Strong]

Odour thresholdNot available.pHNot available.Melting pointNot available.

Boiling point 30 to 210°C (86 to 410°F)

Flash point Closed cup: <-40°C (<-40°F) [Pensky-Martens.]

Evaporation rate Not available.

Flammability (solid, gas) Not applicable. Based on - Physical state

Lower and upper explosive Lower: 1.4% (flammable) limits Upper: 7.6%

Vapour pressure 30.1 to 100.3 kPa (225.6 to 752 mm Hg)

Vapour density Not available.

Relative density Not available.

Density 710 to 750 kg/m³ (0.71 to 0.75 g/cm³)

Solubility Water soluble.(Moderate.)

Partition coefficient: n-

octanol/water

reactions

Not available.

Auto-ignition temperature >350°C (>662°F) **Decomposition temperature**Not available.

Viscosity

Kinematic: 0.4 to 0.55 mm²/s (0.4 to 0.55 cSt) at 40°C

Remarks

Reid vapor pressure (RVP): 55 to 100 kPa (40 °C)

Section 10. Stability and reactivity

ReactivityNo specific test data available for this product. Refer to Conditions to avoid and

Incompatible materials for additional information.

Chemical stability The product is stable.

Possibility of hazardousUnder normal conditions of storage and use, hazardous reactions will not occur.

Under normal conditions of storage and use, hazardous polymerisation will not

occur.

Conditions to avoid Avoid all possible sources of ignition (spark or flame). Avoid excessive heat.

Incompatible materials

Reactive or incompatible with the following materials: oxidising materials.

Hazardous decomposition

Under normal conditions of storage and use, hazardous decomposition products

products should not be produced.

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Information on toxicological effects

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Acute toxicity					
Product/ingredient name	Result	Species	;	Dose	Exposure
Sasoline	LC50 Inhalation Vapour	Rat		>7630 mg/m³ Nominal	4 hours
	LC50 Inhalation Vapour	Rat		>5610 mg/m³ analytical	4 hours
	LD50 Dermal	Rabbit		>2000 mg/kg	-
	LD50 Oral	Rat		>5000 mg/kg	-
Ethanol	LC50 Inhalation Vapour	Rat		124.7 mg/l	4 hours
	LC50 Inhalation Vapour	Rat		116.9 mg/l	4 hours
	LC50 Inhalation Vapour	Rat		133.8 mg/l	4 hours
	LD50 Oral	Rat		10470 mg/kg	-
tert-butyl methyl ether	LC50 Inhalation Vapour	Rat		85 mg/l	4 hours
	LD50 Dermal	Rat		>2000 mg/kg	-
	LD50 Oral	Rat		>2000 mg/kg	-
2-methylpropan-2-ol	LC50 Inhalation Vapour	Rat		>10000 ppm	4 hours
	LD50 Oral	Rabbit		3559 mg/kg	-
	LD50 Oral	Rat		2743 mg/kg	-
diisopropyl ether	LC50 Inhalation Vapour	Rat		40.5 mg/m³	1 hours
	LD50 Dermal	Rabbit		2000 mg/kg	-
	LD50 Oral	Rat		8470 mg/kg	-
Irritation/Corrosion					
Product/ingredient name	Result	Species	Score	e Exposure	Observation
Sasoline	Skin - Irritant	Rabbit	_	-	_
	Eyes - Non-irritating to the eyes.	Rabbit	-	-	-
Ethanol	Skin - Non-irritant to skin.	Rabbit	-	-	-
	Eyes - Cornea opacity	Rabbit	-	-	-

Rabbit

Rabbit

Rabbit

Eyes - Non-irritating to the Rabbit eyes.

Eyes Causes serious eye irritation.

Eyes - Iris lesion

Causes skin irritation.

Eyes - Irritant

Skin - Irritation

Mutagenicity

Skin

tert-butyl methyl ether

Product/ingredient name	Test	Experiment	Result
Sasoline	Equivalent to OECD 476	Experiment: In vitro	Negative
		Subject: Mammal - species unspecified	
	Equivalent to OECD 471	Experiment: In vitro	Negative
		Subject: Non-mammalian species	
	EPA OPPTS 870.5395	Experiment: In vivo Subject: Unspecified Cell: Germ	Negative
	Equivalent to OECD 475	Experiment: In vivo	Negative
		Subject: Unspecified Cell: Germ	
Ethanol	Equivalent to OECD 476	Experiment: In vitro	Negative
		Subject: Mammal - species unspecified	
	Equivalent to OECD	Experiment: In vitro	Negative

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Section 11. Toxico	ologicai	IIIIOIIIIa	atioi					
	473		0.1.					
	Equivalent 478	to OECD		ect: Non-ma riment: In v		ian species	Negative	
tert-butyl methyl ether	EU B 13/14	Į.	Cell:	ect: Unspec Germ riment: In v			Negative	
	0505.474					ian species	NI C	
	OECD 471			riment: In v		ian species	Negative	
	OECD 476		Expe	riment: In v	vitro	ian species	Negative	
	Equivalent 473	to OECD		riment: In v		·	Negative	
	Equivalent 486	to OECD	Expe	riment: In v	vivo	ian species	Negative	
				ect: Unspec	cified			
	Equivalent OPPTS 87			Somatic riment: In v	vivo		Negative	
	0111007	0.0000		ect: Unspec	cified			
	Equivalent OPPTS 79			Somatic riment: In v	vivo		Negative	
	0111079	0.000		ect: Unspec	cified			
Conclusion/Summary Carcinogenicity	May cau	se genetic de	efects.					
Product/ingredient name	Result			Species		Dose	Expo	NEURO
Sasoline	Negative -	Inhalation -		Rat		_		weeks
padomio	Unspecified Negative -	b		Mouse		-		weeks
	Unspecified	b						
Ethanol		Oral - Unspec Oral - Unspe		Mouse Rat		-		weeks weeks
tert-butyl methyl ether		nhalation -	omou	Rat		-	2 yea	
Conclusion/Summary	•	se cancer						
Reproductive toxicity								
Product/ingredient name	Maternal toxicity	Fertility	Deve toxii	elopmental 1	Speci	es	Dose	Exposure
Sasoline	-	Negative	-		Rat		Inhalation	2 generation
Ethanol	-	- Positive	Nega -	ntive	Rat Rat		Inhalation Oral	14 days 2 generation
	-	-	Nega	itive	Rat		Inhalation	18 days
tert-butyl methyl ether	-	Negative	-		Rat		Inhalation	2
	-	-	Nega	itive	Rat		Inhalation	generation 9 days
Specific target organ toxici	ty (single ex	posure)						
Name			C	ategory		Route of exposure	Target	organs

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Category 3Not applicable.Narcotic effectsBenzeneCategory 3Not applicable.Narcotic effectsCategory 3Not applicable.Respiratory tract

irritation

diisopropyl ether Category 3 Not applicable. Narcotic effects

Specific target organ toxicity (repeated exposure)

Name Category Route of Target organs

exposure

Senzene Category 1 Not determined blood system

Aspiration hazard

Name Result

Sasoline ASPIRATION HAZARD - Category 1

Information on likely routes

of exposure

Routes of entry anticipated: Dermal, Inhalation.

Potential acute health effects

Eye contact Causes serious eye irritation.

Inhalation Can cause central nervous system (CNS) depression. May cause drowsiness or

dizziness.

Skin contact Causes skin irritation.

Ingestion Irritating to mouth, throat and stomach. Aspiration hazard if swallowed -- harmful or

fatal if liquid is aspirated into lungs.

Symptoms related to the physical, chemical and toxicological characteristics

Eye contact Adverse symptoms may include the following:

pain or irritation watering

redness

Inhalation Adverse symptoms may include the following:

nausea or vomiting

headache

drowsiness/fatigue dizziness/vertigo unconsciousness

Skin contact Adverse symptoms may include the following:

irritation redness

reduced foetal weight increase in foetal deaths skeletal malformations

Ingestion Adverse symptoms may include the following:

nausea or vomiting reduced foetal weight increase in foetal deaths skeletal malformations

Delayed and immediate effects as well as chronic effects from short and long-term exposure

Eye contact Vapour, mist or fume may cause eye irritation. Exposure to vapour, mist or fume

may cause stinging, redness and watering of the eyes.

Inhalation Vapour, mist or fume may irritate the nose, mouth and respiratory tract.

Skin contact Prolonged or repeated contact can defat the skin and lead to irritation, cracking and/

or dermatitis.

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Ingestion If swallowed, may irritate the mouth, throat and digestive system. If swallowed, may

cause abdominal pain, stomach cramps, nausea, vomiting, diarrhoea, dizziness and

drowsiness.

General Solvent "sniffing" (abuse) or intentional overexposure to vapours can produce

serious central nervous system effects, including unconsciousness, and possibly

death.

Carcinogenicity May cause cancer. Risk of cancer depends on duration and level of exposure.

Mutagenicity May cause genetic defects.

Teratogenicity

Suspected of damaging the unborn child.

No known significant effects or critical hazards.

Fertility effects

No known significant effects or critical hazards.

Numerical measures of toxicity

Acute toxicity estimates

Route ATE value Inhalation (vapours) 1100 mg/l

Other information

Gasoline - Excess exposure to vapors may produce headaches, dizziness, nausea, drowsiness, irritation of eyes, nose and throat and central nervous system depression. Aspiration of this material into the lungs can cause chemical pneumonia and can be fatal. Aspiration into the lungs can occur while vomiting after ingestion of this product. Inhalation of unleaded gasoline vapors did not produce birth defects in laboratory animals. Ingestion of this material can cause gastrointestinal irritation and diarrhea.

In a long-term inhalation study of whole unleaded gasoline vapors, exposure-related kidney damage and kidney tumors were observed in male rats. Similar kidney effects were not seen in female rats or in mice. At the highest exposure level (2056 ppm), female mice had an increased incidence of liver tumors. Results from subsequent scientific studies have shown that a broad variety of chemicals cause these kidney effects only in the male rat. Further studies have discovered the means by which the physiology of the male rat uniquely predispose it to these effects. Consequently, the Risk Assessment Forum of the Environmental Protection Agency has recognized that these responses are not predictive of a human health hazard. The liver tumors that were increased in the high-dose female mice are likewise of questionable significance because of their high spontaneous occurrence even without chemical exposure and because the rate of their occurrence is accelerated by a broad spectrum of chemicals not commonly considered to be carcinogens (e.g., phenobarbital).

Thus, the significance of the mouse liver tumor response in terms of human health is questionable.

Gasoline is a complex mixture of hydrocarbons and contains benzene (typically no more than 2 volume%), toluene, and xylene. Chronic exposure to high levels of benzene has been shown to cause cancer (leukemia) in humans and other adverse blood effects (anemia). Benzene is considered a human carcinogen by IARC, NTP and OSHA. Over exposure to xylene and toluene can cause irritation to the upper respiratory tract, headache and narcosis. Some liver damage and lung inflammation were seen in chronic studies on xylene in guinea pigs but not in rats.

Solvent "sniffing" (abuse) or intentional overexposure to vapors can produce serious central nervous system effects, including unconsciousness, and possibly death.

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Gasoline: Additional toxicity information on the components:

Benzene: Acute toxicity of benzene results primarily from depression of the central nervous system (CNS). Inhalation of concentrations over 50 ppm can produce headache, lassitude, weariness, dizziness, drowsiness, or excitation. Exposure to very high levels can result in unconsciousness and death.

Benzene: Long-term overexposure to benzene has been associated with certain types of leukemia in humans. In addition, the International Agency for Research on Cancer (IARC), the National Toxicology Program, and OSHA consider benzene to be a human carcinogen. Chronic exposures to high levels of benzene have been reported to cause adverse blood effects including anemia. Benzene exposure can occur by inhalation and absorption through the skin.

Inhalation and forced feeding studies of benzene in laboratory animals have produced a carcinogenic response in a variety of organs, including possibly leukemia, other adverse effects on the blood, chromosomal changes and some effects on the immune system. Exposure to benzene at levels up to 300 ppm did not produce birth defects in animal studies; however, exposure to higher dosage levels resulted in a reduction of body weight of the rat pups (fetotoxicity). Changes in the testes have been observed in mice exposed to benzene at 300 ppm, but reproductive performance was not altered in rats exposed to benzene at the same level. Aspiration of this material into the lungs can cause chemical pneumonia and can be fatal. Aspiration into the lungs can occur while vomiting after ingestion of this material.

Toluene: Aspiration of this material into the lungs can cause chemical pneumonia and can be fatal. Aspiration into the lungs can occur while vomiting after ingestion of this material. Deliberate inhalation of high concentrations of toluene has been linked to damage of the brain, liver and kidney. Inhalation of very high concentrations of toluene, such as in cases of solvent abuse, has resulted in sudden death which may be a result of cardiac arrhythmia or central nervous system depression. Mental and/or growth retardation has been reported in children of women who deliberately inhale toluene during pregnancy (usually at thousands of ppm). Foetal developmental toxicity was observed when pregnant rats were exposed to toluene at levels of 1500 ppm. Maternal toxicity was also observed at this concentration. Prolonged, high level exposure to toluene in laboratory animals has resulted in hearing loss. Exposure studies in rats have resulted in adverse effects on the kidney, liver and central nervous system. Studies in occupationally exposed individuals indicate that toluene exposure has been associated with impaired colour vision and decreased performance in some neurobehavioural tests. There are occupational studies which report an association between inhalation exposure to toluene and adverse effects on reproduction including spontaneous abortion. The methodology of these studies and the reliability of the results have been questioned. In a two-generation study in rats, inhalation of toluene at levels up to 2000 ppm did not produce adverse effects on fertility or reproductive performance.

Xylenes: Xylene has been reported to cause central nervous system effects at concentrations above the recommended exposure limit. Xylene vapour becomes irritating at relatively high levels. In one study, eye irritation was reported at exposures of 460 ppm and in one person at 230 ppm after 15 minutes. In another study, no one reported eyes, nose and throat irritation at mixed xylene exposures up to 230 ppm for 30 minutes. Dermal LD50 is expected to be greater than 10g/kg in rabbits, based on test results from similar materials.

Mixed xylenes caused slight hearing loss in rats exposed to 800 ppm in the air for 14 hours/day for six weeks. There is no information available for lower concentrations; however, similar chemicals that have caused these hearing effects at similar concentrations have not caused effects at lower concentrations.

Pregnant animals exposed to xylene or its isomers have been reported to cause development toxicity in rodents when exposed by inhalation. The developmental

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effects observed consisted of delayed development and minor skeletal variations, but no malformations. Because of the high exposure levels used in these studies, we do not believe that these results imply an increased risk of reproductive toxicity to workers exposed to xylene levels at or below the exposure limits.

Xylene and its isomers are not genotoxic.

Technical grade xylene has been tested in a National Toxicology Program carcinogenicity study in rats and mice dosed orally for two years. There was no evidence of carcinogenicity.

Ethylbenzene - The National Toxicology Program (NTP) conducted a 13-week inhalation study with male and female rats and mice at exposure concentrations ranging from 100 to 1000 ppm ethylbenzene. No rats or mice died during the study. Kidney, liver, and lung weights were increased in the exposed rats, while weight increases were observed only in the livers of exposed mice. Treatment-related histopathologic changes were not observed in any tissues of rats and mice. NTP also exposed male and female rats and mice by inhalation to 0, 75, 250, or 750 ppm ethylbenzene for 2 years. There was a statistically significant increase in the number of kidney tumors in male and female rats at 750 ppm. There were also increased incidences of lung tumors in male mice and liver tumors in female mice that were statistically significant at 750 ppm. Except for the male rat kidney tumors, the incidence of the tumors were within the range observed for non-exposed animals from other studies conducted by NTP. The significance of these findings to humans is unknown. Ethylbenzene is not genotoxic. The International Agency for Research on Cancer (IARC) has evaluated ethylbenzene and found it to be possibly carcinogenic to humans (Group 2B).

Ethylbenzene is not genotoxic.

Naphthalene has been reported to cause developmental toxicity in mice after oral exposure to relatively high dose levels, but developmental toxicity was not observed in NTP (National Toxicology Program) sponsored studies in rats and rabbits. Ingestion or inhalation of naphthalene can result in hemolysis and other blood abnormalities, and individuals (and infants) deficient in glucose-6-phosphate dehydrogenase may be especially susceptible to these effects. Inhalation of naphthalene may cause headache and nausea. Airborne exposure can result in eye irritation. Naphthalene exposure has been associated with cataracts in animals and humans.

Ethanol - Human data: In humans excessive consumption of alcoholic beverages during pregnancy is associated with the induction of Fetal Alcohol Syndrome in the offspring. Reduced birth weight and physical and mental defects occur. There is no evidence that such effects might be caused by exposures other than direct ingestion of alcoholic drinks. In humans high lifetime consumption of alcoholic beverages can be associated with certain cancers and effects on the liver. There is no evidence that these can be caused by exposure other than direct ingestion of alcoholic drinks (IARC 1988).

Fish

Section 12. Ecological information

Toxicity

Product/ingredient name Result **Species Exposure G**asoline Acute EC50 15.41 mg/l Nominal Fresh Micro-organism 40 hours Acute EL50 3.1 mg/l Nominal Fresh Algae 72 hours water Acute EL50 3.7 mg/l Nominal Fresh Algae 96 hours water Acute EL50 4.5 mg/l Nominal Fresh Daphnia 48 hours water

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Acute LL50 10 mg/l Nominal Fresh

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96 hours

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	water Acute LL50 8.2 mg/l Nominal Fresh	Fish	96 hours
	water		
	Acute NOELR 0.5 mg/l Nominal Fresh water	Algae	72 hours
	Acute NOELR 0.5 mg/l Nominal Fresh water	Daphnia	48 hours
	Chronic EL50 10 mg/l Nominal Fresh water	Daphnia	21 days
	Chronic EL50 >40 mg/l Nominal Fresh water	Daphnia	21 days
	Chronic EL50 10 mg/l Nominal Fresh water	Fish	21 days
	Chronic LL50 5.2 mg/l Nominal Fresh water	Fish	14 days
	Chronic NOELR 2.6 mg/l Nominal Fresh water	Daphnia	21 days
	Chronic NOELR 16 mg/l Nominal Fresh water	Daphnia	21 days
	Chronic NOELR 2.6 mg/l Nominal Fresh water	Fish	14 days
	Chronic NOELR 2.6 mg/l Nominal Fresh water	Fish	21 days
	Chronic PNEC >0.4 mg/kg	soil, plants	_
Ethanol	EC50 675 mg/l	Algae	4 days
	EC50 4432 mg/l	Aquatic plants	7 days
	Acute LC50 5012 mg/l	Daphnia	48 hours
	Acute LC50 153 g/l	Fish	96 hours
	Acute LC50 14.2 g/l	Fish	96 hours
	Chronic LC50 2 mg/l	Daphnia	10 days
	Chronic LC50 9.6 mg/l	Daphnia	9 days
tert-butyl methyl ether	Acute EC50 472 mg/l Fresh water	Daphnia	48 hours
, ,	Acute LC50 200 mg/l Marine water	Crustaceans	96 hours
	Acute LC50 672 mg/l Fresh water	Fish	96 hours
	Acute LC50 574 mg/l Marine water	Fish	96 hours
	Chronic NOEC 26 mg/l Marine water	Crustaceans	28 days
	Chronic NOEC 51 mg/l Fresh water	Daphnia	21 days

Conclusion/Summary

Toxic to aquatic life with long lasting effects.

Persistence and degradability

The biodegradability of this material has not been determined.

Product/ingredient name	Test	Result	Dose	Inoculum
⊵ thanol	EPA	95 % - Readily - 15 days	_	-
	EPA	84 % - Readily - 20 days	-	-
	EPA	74 % - Readily - 5 days	-	-
	EPA	74 % - Readily - 10 days	-	-
tert-butyl methyl ether	not guideline	100 % - 1.25 days	-	-
	Modelled data	61 to 69 % - 151 days	-	-
	OECD 301 D	9.24 % - Not readily - 28 days	-	-
	OECD 301 D	1.8 % - Not readily - 28 days	-	-
	OECD 301 D	0 % - Not readily - 28 days	-	-
	Modelled data	0 % - 250 days	-	-
Conclusion/Summary	Partially biode	egradable.		
Product/ingredient name	Aquatic half-life	Photolysi	S	Biodegradability
⊵ thanol	-	-		Readily

Bioaccumulative potential

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This product is not expected to bioaccumulate through food chains in the environment.

Product/ingredient name	LogPow	BCF	Potential
G asoline	2 to 7	-	high
Ethanol	-0.35	-	low
Benzene	2.13	11	low
tert-butyl methyl ether	1.04	1.5	low
2-methylpropan-2-ol	0.317	-	low
diisopropyl ether	2.4	-	low

Mobility in soil

Soil/water partition coefficient (Koc)

Not available.

Mobility Spillages may penetrate the soil causing ground water contamination.

Other ecological information

Spills may form a film on water surfaces causing physical damage to organisms. Oxygen transfer could also be impaired.

Section 13. Disposal considerations

Disposal methods

The generation of waste should be avoided or minimised wherever possible. Significant quantities of waste product residues should not be disposed of via the foul sewer but processed in a suitable effluent treatment plant. Dispose of surplus and non-recyclable products via a licensed waste disposal contractor. Disposal of this product, solutions and any by-products should at all times comply with the requirements of environmental protection and waste disposal legislation and any regional local authority requirements. Waste packaging should be recycled. Incineration or landfill should only be considered when recycling is not feasible. This material and its container must be disposed of in a safe way. Care should be taken when handling emptied containers that have not been cleaned or rinsed out. Empty containers or liners may retain some product residues. Vapour from product residues may create a highly flammable or explosive atmosphere inside the container. Do not cut, weld or grind used containers unless they have been cleaned thoroughly internally. Avoid dispersal of spilt material and runoff and contact with soil, waterways, drains and sewers.

Special Precautions for Landfill or Incineration

No additional special precautions identified.

Section 14. Transport information

	ADG	IMDG	IATA
UN number	UN1203	UN1203	UN1203
UN proper shipping name	MOTOR SPIRIT or GASOLINE or PETROL	SASOLINE or MOTOR SPIRIT. Marine pollutant	MOTOR SPIRIT or GASOLINE or PETROL
Transport hazard class(es)	3	3	3
Packing group	II	II	II
Environmental hazards	No.	Yes.	Yes. The environmentally hazardous substance mark is not required.

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Section 14. Transport information Additional Hazchem code 3YE The marine pollutant mark is The environmentally information Initial emergency response not required when transported hazardous substance mark <u>quide</u> 14 in sizes of ≤5 L or ≤5 kg. may appear if required by other transportation **Emergency schedules** F-E. S-E regulations.

Special precautions for user

Not available.

Transport in bulk according to Annex II of Marpol and the IBC Code

Proper shipping name

MARPOL Annex 1 rules apply for bulk shipments

by sea.

Category: gasoline and spirits

Section 15. Regulatory information

Standard Uniform Schedule of Medicine and Poisons

Not scheduled - When packed in containers having capacity of greater than 20 litres.

S5 - When packed in containers having capacity of less than 20 litres.

Consumer products - This product is exempt per Appendix A of the SUSMP.

Industrial Products - Labelling requirements for SUSMP do not apply to a poison that is packed and sold solely for industrial, laboratory or manufacturing use. However, this product is labelled in accordance with NOSHC National Code of Practice for labelling of workplace substances.

Model Work Health and Safety Regulations - Scheduled Substances

No listed substance

Montreal Protocol (Annexes A, B, C, E)

Ingredient name Not listed.	List name	Status		
Stockholm Convention on Persistent Organic Persiste	<u>ollutants</u>			
Ingredient name Not listed.	List name	Status		
Rotterdam Convention on Prior Informed Consent (PIC)				
Ingredient name Not listed.	List name	Status		

International lists

National inventory

REACH Status For the REACH status of this product please consult your company contact, as

identified in Section 1.

Australia inventory (AICS) All components are listed or exempted.

All components are listed or exempted. Canada inventory

China inventory (IECSC) At least one component is not listed.

Not determined. **Japan inventory (ENCS) Korea inventory (KECI)** Not determined. **Philippines inventory** Not determined.

(PICCS)

Not determined.

Taiwan Chemical Substances Inventory

(TCSI)

At least one component is not listed.

United States inventory (TSCA 8b)

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Section 16. Any other relevant information

History

Date of printing 1/23/2020 Date of issue/Date of 1/23/2020

revision

Date of previous issue 9/28/2015

Version 2

Prepared by Product Stewardship

Key to abbreviations ADG = Australian Dangerous Goods

ATE = Acute Toxicity Estimate BCF = Bioconcentration Factor

GHS = Globally Harmonized System of Classification and Labelling of Chemicals

IATA = International Air Transport Association

IBC = Intermediate Bulk Container

IMDG = International Maritime Dangerous Goods

LogPow = logarithm of the octanol/water partition coefficient

MARPOL = International Convention for the Prevention of Pollution From Ships,

1973 as modified by the Protocol of 1978. ("Marpol" = marine pollution) NOHSC = National Occupational Health and Safety Commission

REACH = Registration, Evaluation, Authorisation and Restriction of Chemicals

Regulation [Regulation (EC) No. 1907/2006]

STEL = Short term exposure limit

SUSMP = Standard Uniform Schedule of Medicine and Poisons

UN = United Nations

TWA = Time weighted average VOC = Volatile Organic Compound

SADT = Self-Accelerating Decomposition Temperature

Varies = may contain one or more of the following 64741-88-4, 64741-89-5, 64741-95-3, 64741-96-4, 64742-01-4, 64742-44-5, 64742-45-6, 64742-52-5, 64742-53-6, 64742-54-7, 64742-55-8, 64742-56-9, 64742-57-0, 64742-58-1, 64742-62-7, 64742-63-8, 64742-65-0, 64742-70-7, 72623-85-9, 72623-86-0,

72623-87-1

Procedure used to derive the classification

Classification	Justification
F am. Liq. 1, H224	On basis of test data
Skin Irrit. 2, H315	Calculation method
Eye Irrit. 2A, H319	Calculation method
Muta. 1B, H340	Expert judgment
Carc. 1B, H350	Expert judgment
Repr. 2, H361 (Unborn child)	Calculation method
STOT SE 3, H336	Calculation method
Asp. Tox. 1, H304	Calculation method

▼ Indicates information that has changed from previously issued version.

Notice to reader

All reasonably practicable steps have been taken to ensure this data sheet and the health, safety and environmental information contained in it is accurate as of the date specified below. No warranty or representation, express or implied is made as to the accuracy or completeness of the data and information in this data sheet.

The data and advice given apply when the product is sold for the stated application or applications. You should not use the product other than for the stated application or applications without seeking advice from BP Group.

It is the user's obligation to evaluate and use this product safely and to comply with all applicable laws and regulations. The BP Group shall not be responsible for any damage or injury resulting from use, other than the stated product use of the material, from any failure to adhere to recommendations, or from any hazards inherent in the nature of the material. Purchasers of the product for supply to a third party for use at work, have a duty to take all necessary steps to ensure that any person handling or using the product is provided with the information in this sheet. Employers have a duty to tell employees and others who may be affected of any hazards described in this sheet and of any precautions that should be taken. You can contact the BP Group to ensure that this document is the most current available.

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Section 16. Any other relevant information Alteration of this document is strictly prohibited.

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