	PATIENT LABEL
ANZAAG	Surname:
ANZAAG Australian & New Zealand	Given Names:
Anaesthetic Allergy Group	Date of birth: Gender:
DEEEDDAL EODM	Address:
REFERRAL FORM	Record Number:
Hand print patient name Please check patient name, address and pho	one number on label are correct
Patient's Email:	
Home phone:	Mobile:
Referring Doctor (name):	
Provider Number:	
Position: Anaes consultant Anaes Regis	strar GP Anaesthetist Other:
Phone:	Mobile:
Email:	
Postal address:	
Please tick relevant conditions: Pregnant Drug Allergy (specify) Food Allergy (specify) Other Allergy (specify) Other Medical History:	Asthma Eczema Hay fever
Current Medication Tick where patient taking: Oral steroids And ACE Inhibitors/AII Receptor antagonist List medications:	ntihistamines βblockers Antidepressants NSAID

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		Given Names:	
		Date of birth: Gender:	
Anaesthetic Allergy Gr	000	Address:	
REFERRAL FOR	RM	Record Number:	
Hospital where reaction occurred: Procedure:			
Date of reaction:		Date of referral:	
Time of induction (24 hour clock):		Time reaction first noted:	
Type of Anaesthesia: General R	egional	Local IV sedation	
The patient was exposed to the following	medication	ns PRIOR to the reaction(indicate time of ex	cposure):
Agent Administered	Time	Agent Administered	Time
Please tick if the patient was exposed to t	he gaante l	istad halow (indianta tima of arnosura).	Time
	-		Time
Chlorhexidine wipes		prep Other (specify):	
Skin preparation Type	e:		
Latex Gloves	Other	r (specify):	
Contrast Agent Typ	e:		
Methylene Blue Patent B	lue		
Colloid Typ	be:		
Blood products Ty	pe:		
Antibiotics Ty	pe:		
Central venous line Chlorhex	idine coate	d Antibiotic coated Other	
Uaginal packing Ty	pe:		
	1		1
Urinary catheter Ty	pe:		



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	PATIENT LABEL
ANZAAG	Surname:
	Given Names:
acouncile Anergy Group	Date of birth: Gender:
	Address:
L FORM	Record Number:
_	
Reaction	
	No
Yes I	No
Yes I	No Type:
Yes I	No
Yes I	No Time with systolic < 60mmHgmins
Yes N	lo
Yes N	lo
Mild wh	neeze Dyspnoea reported by patient
Moderat	te wheeze Difficult to ventilate
Severe v	wheeze Very difficult to ventilate
Yes I	No SpO2 80-90 SpO2 <80
Yes I	No Localised or Generalised
Yes I	No Localised or Generalised
Yes I	No
Yes I	No
Yes I	No Site
	Duration
Yes I	No Specify:
Yes I	No Nausea Vomiting
	Abdominal cramps/pain
	Other
n you noticed?	
symptom?	
	tered) Yes I Yes I Yes I Yes I Yes I Yes I N Yes I N N Ves I N Ves I Yes I I I Yes I I I Yes I I I Yes I I I I I Yes I I I I I I I I I I I I I I I I I I I

ANZAAG Australian & New Zealand Anaesthetic Allergy Group
Anaesthetic Allergy Group

REFERRAL FOR

Assisted/Mechanical Ventilation

Vasopressors other than adrenaline given?

Specify steroid used & dose:

Specify antihistamine used & dose:

Did you use the ANZAAG Anaphylaxis Management Resource?

IV Fluids given for resuscitation?

Endotracheal intubation

Bronchospasm treatment?

Adrenaline given?

Cardiac compressions?

Steroids given?

Antihistamines used?

Cardioversion/Defibrillation

Details of Treatment

Airway Management

	Surname:		
Australian & New Zealand Anaesthetic Allergy Group	UIVEII INAIIIES.		
		G	ender:
	Address:		
EFERRAL FOR	Record Number		
_			
of Treatment			
Management			
sisted/Mechanical Ventilation 🗌 Y	es 🗌 No [Planned	Unplanned
dotracheal intubation Y	es 🗌 No [Before onset	After onset
ospasm treatment?	es 🗌 No		
Specify agent/s used & dose:			
line given?	es 🗌 No [C ETT
Total dose administered:	mcg		
ds given for resuscitation?	es 🗌 No		
Specify type/s of fluid & total volume	me:		
c compressions?	es 🗌 No How	long was CPR perf	formed?:mins
version/Defibrillation	es 🗌 No I	Number of shocks:	
essors other than adrenaline given?	${ Yes No}$		
Ephedrine Dose		l Dose	mø
			-
Vasopressin Dose			-
Noradrenaline Dose	mg Methylene I	Blue Dose	_mg
Other (specify):			
s given?	Yes No		

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No

PATIENT LABEL

Other treatments/Comments:

Yes

Please comment on any ways in which you think the resource was helpful or could be improved:

l No

Yes

	PATIENT LABEL		
ANZAAG	Surname:		
Australian & New Zealand	Given Names:		
Anaesthetic Allergy Group	Date of birth: Gender:		
	Address:		
REFERRAL FORM	Record Number:		
Investigations			
Serum tryptase taken? Yes No			
Recommended to take 10ml samples 1-2 hours, 4 hours	urs and more than 24 hours after reaction:		
Please record time samples taken and attach results	to this referral (where available)		
Sample 1: TimeResult:mcg/L	Sample 3: TimeResult:mcg/L		
Sample 2: TimeResult:mcg/L	Sample 4: TimeResult:mcg/L		
Which pathology laboratory were the specimens sent	to?		
Is there a differential diagnosis other than anaphylaxi	s that you think may have caused the reaction?		
Comments:			
Outcome/Sequelae			
Operation/procedure completed or	Operation/procedure abandoned		
Patient transferred to PACU/recovery? Yes	No		
Was the patient admitted to hospital?	No Tick if admission unplanned		
Postoperative care in ICU/HDU? Yes	No		
If yes: Was the patient still intubated/ventilated on tr	ansfer? Yes No Duration		
Was an inotrope infusion continued?	Yes No Duration		
How long was the patient in ICU?			
Were there any further complications?			
ECG Changes Coagulopathy Trop	onin rise Pneumothorax Anxiety/PTSD		
Other			
Severity of Allergic Reaction			
Please specify the Grade of Allergic Reaction from the	e categories below:		
Grade I – cutaneous-mucous signs: erythema, u	rticaria with or without angioedema		
Grade II – Moderate multivisceral signs: cutand	eous-mucous signs +/- hypotension +/- tachycardia		
+/- dyspnoea +/- gastrointestinal disturbance			
Grade III – Life-threatening mono- or multivis	ceral signs: cardiovascular collapse, tachycardia or		
bradycardia +/- cardiac dysrythmia +/- bronchosp	asm +/- cutaneous-mucous signs +/- gastrointestinal		
disturbance			
🗌 Grade IV – cardiac arrest			



REFERRAL FORM

Comments:

 PATIENT LABEL

 Surname:

 Given Names:

 Date of birth:

 Date of birth:

 Address:

 Record Number:

Please tick to acknowledge that you are aware of the following:

You are responsible for forwarding this referral and supporting documents listed to your nearest or preferred ANZAAG member. A contact list of testing specialists can be found at www.anzaag.com

A copy of the resuscitation/anaesthetic/PACU charts and tryptase results (where available) must accompany this referral.

The correct patient details have been supplied to allow follow up with the patient.

The patient is aware of the events and this referral. The patient information brochure available at <u>www.anzaag.com</u> may assist with this discussion.

The patient has a letter listing all substances administered perioperatively to show to those providing care until testing can be conducted. A form letter is available at <u>www.anzaag.com</u> to assist in this process.

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