

Health Canada Santé Canada

Canada Vigilance Summary of Reported Adverse Reactions

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A 173 Report(s)

Brand Name/Active Ingredient: 'PANZYGA'

Search Date Criteria: 1965-01-01 to 2024-07-31

Reaction Term(s): All/Tous

Serious report?: Both

Type of Report: All

Source of Report: All

Gender: All

Report Outcome: All

Age: All

CAVEAT: This summary is based on information from adverse reaction reports submitted by health professionals and laypersons either directly to Health Canada or via market authorization holders. Each report represents the suspicion, opinion or observation of the individual reporter. The Canada Vigilance Program is a spontaneous reporting system that is suitable to detect signals of potential health product safety issues during the post-market period. The data has been collected primarily by a spontaneous surveillance system in which adverse reactions to health products are reported on a voluntary basis. Under reporting of adverse reactions is seen with both voluntary and mandatory spontaneous surveillance systems. Accumulated case reports should not be used as a basis for determining the incidence of a reaction or estimating risk for a particular product as neither the total number of reactions occurring, nor the number of patients exposed to the health product is known. Because of the multiple factors that influence reporting, quantitative comparisons of health product safety cannot be made from the data. Some of these factors include the length of time a drug is marketed, the market share, size and sophistication of the sales force, publicity about an adverse reaction and regulatory actions. In some cases, the reported clinical data is incomplete and there is not certainty that these health products caused the reported reactions. A given reaction may be due to an underlying disease process or to another coincidental factor. This information is provided with the understanding that the data will be appropriately referenced and used in conjunction with this caveat statement.



Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000694440	1	2017-03-24	2017-04-04	Hospital		Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Not Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions:

Patient Informa	tion			
Age	Gender	Height	Weight	Report Outcome
43 Years	Female	164 Centimeter	59 Kilogram	Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report	

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intracavernous	60.0 Gram	1 every 8 Months		Immune thrombocytopen ia

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Throat irritation	v.27.1	
Urticaria	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date	Source of Report	Market Authorization Holder AER Number	Type of Report	Reporter Type
000695154	0	2017-04-06	2017-04-06	MAH	NGAM-003-17-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient	Informa	tion	
_			

Age		Gender	Height	Weight	Report Outcome
43 Ye	ars	Female	164 Centimeter	59 Kilogram	Recovered/resolved

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	60.0 Gram	Total		Immune thrombocytopen ia

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Throat irritation	v.27.1	
Urticaria	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000697271	1	2017-05-18	2017-05-30	MAH	NGAM-008-17-CA	Spontaneous	Other health professional

Serious report?	Death: Yes	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Informa	ition			
Age Gender		Height	Weight Report Outcome	
43 Years	Female			Fatal

Link / Duplicate Report Information	1
Record Type	Link AER** Number
No duplicate or linked report	

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	35.0 Gram	1 every 1 Days		Secondary immunodeficien cy

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Condition aggravated	v.27.1	
Dyspnoea	v.27.1	
Tachycardia	v.27.1	
Tachypnoea	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000697502	2	2017-05-24	2017-06-22	MAH	NGAM-009-17-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization: Yes	Other Medically Important Conditions: Yes

Patient	Informa	tion

Age	Gender	Height	Weight	Report Outcome
63 Years	Female			Recovered/resolved

Link / Duplicate Report Information

Record Type	Link AER** Number

No duplicate or linked report.

Product Information

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Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	50.0 Gram	1 every 1 Days	2.0 Days	Eosinophilic fasciitis
ZOPICLONE	Concomitant	Tablets					

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Meningitis aseptic	v.27.1	
Nausea	v.27.1	
Pain	v.27.1	
Vomiting	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Informati	on	**AE	R = Adverse F	Reaction Re	port
Adverse Reaction Report Number	Latest AEI Num		Initial Rece	ived Date	Latest
000697506	1		2017-0	5-24	2
Serious i	eport?				Death:
Serio	us		Life Threa		ening:
Patient Informati	on				
Age	Gender		Height	V	Veight
67 Years	Female				
Link / Duplicate	Report Infor	mation			
	Record				
No duplicate or lin	ked report.				
Product Information	ion				

ed Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
-24	2017-05-31	MAH	NGAM-010-17-CA	Spontaneous	Other health professional

Death:	Disability:	Congenital Anomaly:	
Life Threatening:	Hospitalization:	Other Medically Important Conditions:	Yes

/eight	Report Outcome
	Unknown

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report	

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
ACETYLSALICYLIC ACID	Concomitant	NOT SPECIFIED					
AMLODIPINE	Concomitant	Tablets					
ATORVASTATIN CALCIUM	Concomitant	Tablets					
DIPHENHYDRAMINE	Concomitant	NOT SPECIFIED					
HYDROCHLOROTHIAZIDE	Concomitant	Tablets					
JANUVIA	Concomitant	Tablets					
LOSARTAN	Concomitant	Tablets					
METHYLPREDNISOLONE NOS	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	40.0 Gram	1 every 1 Days		Secondary immunodeficien cy
RANITIDINE	Concomitant	NOT SPECIFIED					

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Back pain	v.27.1	
Chest discomfort	v.27.1	
Chest pain	v.27.1	
Dysarthria	v.27.1	
Dyspnoea	v.27.1	
Headache	v.27.1	·
Lip swelling	v.27.1	
Musculoskeletal stiffness	v.27.1	
Nasal congestion	v.27.1	
Paraesthesia oral	v.27.1	
Rash	v.27.1	
Sensation of foreign body	v.27.1	
Swollen tongue	v.27.1	
Vision blurred	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000698047	0	2017-06-05	2017-06-05	Hospital		Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Not Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions:

Patient	Informati	tion

Age	Gender	Height	Weight	Report Outcome
65 Years	Female	158 Centimeter	56 Kilogram	Unknown

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	1000.0 mg/kg	1 every 1 Days		Immune thrombocytopen ia
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	1000.0 mg/kg	1 every 1 Days		Immune thrombocytopen ia
RED BLOOD CELLS	Concomitant	NOT SPECIFIED					

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Anti A antibody positive	v.27.1	
Anxiety	v.27.1	
Blood bilirubin increased	v.27.1	
Blood lactate dehydrogenase increased	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Chills	v.27.1	
Coombs direct test positive	v.27.1	
Heart rate increased	v.27.1	
Hyperhidrosis	v.27.1	
Nausea	v.27.1	
Pyrexia	v.27.1	
Vomiting	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

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**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000698066	0	2017-06-06	2017-06-06	Hospital		Spontaneous	Physician

Serious report?	Death:	Disability:	Congenital Anomaly:
Not Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions:

Patient Information

Age Gender		Height	Weight	Report Outcome	
31 Years	Female	165 Centimeter	69 Kilogram	Recovered/resolved	

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

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Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	30.0 Gram	1 every 4 Weeks		Nasopharyngitis

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Abdominal pain	v.27.1	
Back pain	v.27.1	
Chills	v.27.1	
Headache	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000698480	0	2017-06-16	2017-06-16	Hospital		Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization: Yes	Other Medically Important Conditions:

Patient Informatio	n
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Age	Age Gender Height		Weight	Report Outcome	
46 Years	Female	171 Centimeter	65 Kilogram	Recovered/resolved	

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
ACTONEL	Concomitant	Tablets					
CALCIUM	Concomitant	NOT SPECIFIED					
LORAZEPAM	Concomitant	NOT SPECIFIED					
METHOTREXATE	XATE Concomitant NC						
MULTIVITAMINE(S)	Concomitant	NOT SPECIFIED					
PANIZYGA Suspect SOLUTIO		SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	70.0 Gram	2 every 1 Months		Dermatomyositi s
PREDNISONE	Concomitant	NOT SPECIFIED					
SULFATRIM	Concomitant	NOT SPECIFIED					
VITAMIN D	Concomitant	NOT SPECIFIED					

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Pruritus	v.27.1	
Urticaria	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000698501	2	2017-06-14	2017-12-13	MAH	NGAM-016-17-CA	Spontaneous	Physician

Serious report?	Death:	Disability:	Congenital Anomaly:	
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions:	Yes

Patient Information

Age	Age Gender Height		Weight	Report Outcome	
31 Years	Female	165 Centimeter	69 Kilogram	Recovered/resolved	

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	14.4 Gram			Immunodeficien cy common variable

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Abdominal pain	v.27.1	
Back pain	v.27.1	
Chills	v.27.1	
Headache	v.27.1	
Maternal exposure during pregnancy	v.27.1	
Pyrexia	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Informatio	n	**AE	R = Adverse F	Reaction Re	eport								
Adverse Reaction Report Number		R Version	Initial Recei	ved Date	Latest Rece	eived Date	Sourc	e of Report	Autho	arket orization ER Number	Type of Report	Reporter	Туре
000698524		0	2017-0	6-14	2017-0	6-14		MAH	NGAM-	014-17-CA	Spontaneous	Physici	an
Serious re	port?				Death:			Disability	y:		Congenital	Anomaly:	
Seriou	ıs		L	ife Threat	ening:		Н	ospitalizatio	n:	Other Med	dically Important Co	onditions:	Yes
Patient Information	on												
Age	Gender		Height	\	Neight		Report	Outcome					
74 Years			O Centimeter	46	46 Kilogram		Recovered/resolved						
Link / Duplicate R	eport Info	rmation											
	Recor	d Type			Link AER** Number								
No duplicate or link	ed report.												
Product Informati	on												
Product Descr	ription	Health Pr	oduct Role	Dosag	je Form	Route Administr		Dose	Fre	equency	Therapy Duration	Indication	on(s)
PANZYGA		Sus	spect		JTION /ENOUS	Intravenou otherwi specifie	se	20.0 Gram	1	Total		Hypogam bulinae	
Adverse Reaction Information	Term												
	Adv	verse Reac	tion Term(s)			Me	dDRA	Version			Reaction Duration		

v.27.1

v.27.1

Flank pain

Oxygen saturation decreased

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date	Source of Report	Market Authorization Holder AER Number	Type of Report	Reporter Type
000698573	1	2017-06-16	2018-01-19	MAH	NGAM-015-17-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Informa	ition			
Age	Gender	Height	Weight	Report Outcome
65 Years	Female		56 Kilogram	Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
No displicate or linked report	

No duplicate or linked report.

Product	Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	1000.0 mg/kg	Total		Immune thrombocytopen ia
RED BLOOD CELLS	Suspect	NOT SPECIFIED	Unknown	2.0 Units	Total		Product used for unknown indication

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Antibody test positive	v.27.1	
Anxiety	v.27.1	
Blood bilirubin increased	v.27.1	
Blood lactate dehydrogenase increased	v.27.1	
Chills	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Coombs direct test	v.27.1	
Delayed haemolytic transfusion reaction	v.27.1	
Heart rate increased	v.27.1	
Hyperhidrosis	v.27.1	
Nausea	v.27.1	
Pyrexia	v.27.1	
Vomiting	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000699797	0	2017-07-07	2017-07-07	MAH	NGAM-019-17-CA	Spontaneous	Consumer/other non health professional

Serious report?	Death:	Disability:		Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Yes	Other Medically Important Conditions:

Patient Information	1
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Age	Gender	Height	Weight	Report Outcome
72 Years	Female			Recovered/resolved

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	13.0 Gram			Kidney transplant rejection

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Dyspnoea	v.27.1	
Foaming at mouth	v.27.1	
Loss of consciousness	v.27.1	
Nausea	v.27.1	
Vomiting	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000699799	1	2017-07-07	2017-07-24	MAH	GAM-150-17-CA	Spontaneous	Consumer/other non health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

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Age	Gender	Height	Weight	Report Outcome
79 Years	Female			Not recovered/not resolved

Link / Duplicate Report Information

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No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
BUPROPION	Concomitant	TABLET (EXTENDED- RELEASE)					
OCTAGAM 10% FOR I.V. INFUSION	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	25.0 Gram	1 every 1 Months		Secondary immunodeficien cy
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	25.0 Gram	1 every 1 Months		Secondary immunodeficien cy

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Hepatitis B core antibody positive	v.27.1	
Hepatitis B surface antibody positive	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Suspected transmission of an infectious agent via product	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adver Reaction Number	Report	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000699	9972	1	2017-07-12	2017-08-01	MAH	NGAM-020-17-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:		Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Yes	Other Medically Important Conditions:

Patient Information				
Age Gender		Height	Weight	Report Outcome
50 Years	Male			Recovered/resolved

Link / Duplicate Report Information	1
Record Type	Link AER** Number
No duplicate or linked report	

Product Information

Froduct information							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	95.0 Gram	Total		Multiple sclerosis

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Dizziness	v.27.1	
Headache	v.27.1	
Nausea	v.27.1	
Vomiting	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information **AER = A

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000700984	0	2017-08-01	2017-08-01	MAH	NGAM-029-17-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Information				
Age Gender		Height	Weight	Report Outcome
73 Years	Female			Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report.	

Product Information

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Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
CELECOXIB	Concomitant	Capsules					
DOMPERIDONE	Concomitant	Tablets					
HYDROCHLOROTHIAZIDE	Concomitant	Tablets					
LORAZEPAM	Concomitant	NOT SPECIFIED					
PANTOPRAZOLE	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	30.0 Milligram	1 every 1 Days		Chronic inflammatory demyelinating polyradiculoneu ropathy
PREGABALIN	Concomitant	Capsules					
TRIAMCINOLONE	Concomitant	NOT SPECIFIED					

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Abdominal pain upper	v.27.1	
Asthenia	v.27.1	
Chills	v.27.1	
Dizziness	v.27.1	
Head discomfort	v.27.1	
Headache	v.27.1	
Hypertension	v.27.1	
Malaise	v.27.1	
Nausea	v.27.1	
Palpitations	v.27.1	
Vomiting	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Adverse Reaction Repor Number	Latest AER Version Number	Initial Received Date	Latest Rece	eived Date	Source of Report	Autho	arket orization ER Number	Type of Report	Reporter '	Гуре
000701655	2	2017-08-18	2017-0	9-08	MAH	NGAM-	036-17-CA	Spontaneous	Other he profession	
Serious	report?		Death:		Disabilit	y:	Congenital A		Anomaly:	
Serie	ous	Life Threat	Life Threatening:		Hospitalization: Other Med		lically Important C	onditions:	Yes	
Patient Informat	ion									
Age	Gender	Height V	Veight		Report Outcome					
78 Years	Female				Recovered/resolved					

Product Information

No duplicate or linked report.

Product information							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
ACETAMINOPHEN	Concomitant	NOT SPECIFIED					
AMLODIPINE	Concomitant	Tablets					
IPRATROPIUM	Concomitant	NOT SPECIFIED					
MAGNESIUM	Concomitant	NOT SPECIFIED					
METOPROLOL	Concomitant	NOT SPECIFIED					
METRONIDAZOLE	Concomitant	NOT SPECIFIED					
METRONIDAZOLE	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	10.0 Gram		75.0 Minutes	Clostridium difficile infection
POTASSIUM CHLORIDE	Concomitant	NOT SPECIFIED					
VANCOMYCIN	Concomitant	NOT SPECIFIED		·			

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Hypersensitivity	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000702366	0	2017-09-06	2017-09-06	Hospital		Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization: Yes	Other Medically Important Conditions:

Patient Information

Age	Gender	Height	Weight	Report Outcome
46 Years	Female	171 Centimeter	65 Kilogram	Recovered/resolved

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

1 Todact information							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
ACTONEL	Concomitant	Tablets					
APO SULFATRIM TAB	Concomitant	Tablets					
CALCIUM	Concomitant	NOT SPECIFIED					
DIPHENHYDRAMINE HYDROCHLORIDE INJECTION USP	Concomitant	LIQUID INTRAMUSCULAR					
LORAZEPAM	Concomitant	NOT SPECIFIED					
MULTIVITAMINE(S)	Concomitant	NOT SPECIFIED					
OCTAGAM 10% FOR I.V. INFUSION	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	10.0 Gram			Dermatomyositi s
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	60.0 Gram			Dermatomyositi s

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PREDNISONE	Concomitant	NOT SPECIFIED					1
VITAMIN D	Concomitant	NOT SPECIFIED					

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Blood pressure increased	v.27.1	
Heart rate increased	v.27.1	
Rash erythematous	v.27.1	
Rash pruritic	v.27.1	
Urticaria	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31

Therapy Duration Indication(s)

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000702762	0	2017-09-14	2017-09-14	Community		Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Not Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions:

Patient Informa	tion			
Age	Gender	Height	Weight	Report Outcome
57 Years	Male	67 Inch	214 Pound	Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
Duplicate	000713845

Product Information					
Product Description	Health Product Role	Dosage Form	Route of	Dose	Fr

1 Toddot Besoription	Ticulti i Toddot itole	Dosage Form	Administration	200	Trequency	Therapy Baration	maioation(s)
BENADRYL	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	100.0 Gram	Cyclical		Myasthenia gravis

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Urticaria	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000702969	0	2017-09-19	2017-09-19	MAH	NGAM-043-17-A	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization: Yes	Other Medically Important Conditions: Yes

Patient Information	ì
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Age	Gender	Height	Weight	Report Outcome
46 Years	Female	171 Centimeter	65 Kilogram	Recovered/resolved

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

i ioaaot iiiioiiiiatioii							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
CALCIUM	Concomitant	NOT SPECIFIED					
DIPHENHYDRAMINE	Concomitant	NOT SPECIFIED					
LORAZEPAM	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	70.0 Gram	1 every 1 Months		Dermatomyositi s
PREDNISONE	Concomitant	NOT SPECIFIED					
RISEDRONATE (MANUFACTURER UNKNOWN)	Concomitant	NOT SPECIFIED					
SULFAMETHOXAZOLE AND TRIMETHOPRIM	Concomitant	NOT SPECIFIED					
VITAMIN D	Concomitant	NOT SPECIFIED					

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Blood pressure increased	v.27.1	
Heart rate increased	v.27.1	
Pruritus	v.27.1	
Rash	v.27.1	
Urticaria	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information **AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000703006	0	2017-09-19	2017-09-19	MAH	NGAM-039-17-CA	Spontaneous	Physician

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Information	
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Ì	Age	Gender	Height	Weight	Report Outcome
	69 Years	Male			Unknown

Link / Duplicate Report Information

		Record Type	Link	AER** Number	er
No duplicate or linked report.					

Product Information

Froduct information							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
DIPHENHYDRAMINE	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	30.0 Gram		55.0 Minutes	Secondary immunodeficien cy
PIPERACILLIN AND TAZOBACTAM FOR INJECTION	Concomitant	POWDER FOR SOLUTION INTRAVENOUS					

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Chills	v.27.1	
Confusional state	v.27.1	
Pyrexia	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31

173 Report(s)

Report Information

Adverse Reaction Term

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000703346	0	2017-09-27	2017-09-27	Hospital		Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Not Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions:

Patient Informa	tion			
Age	Gender	Height	Weight	Report Outcome
55 Years	Female	157 Centimeter	79 Kilogram	Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
Duplicate	000713844

Product Information							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	60.0 Gram	1 every 1 Days	2.0 Days	Immune thrombocytopen ia

Info	ormation			
	Adverse	e Reaction Term(s)	MedDRA Version	Reaction Duration
Urti	icaria		v 27 1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000703391	0	2017-09-28	2017-09-28	Hospital		Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Not Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions:

Patient Informa	tion			
Age	Gender	Height	Weight	Report Outcome
52 Years	Female	164 Centimeter	87 Kilogram	Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report.	

Product Information

Froduct information							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	25.0 Gram	1 every 4 Weeks		Secondary immunodeficien cy

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Back pain	v.27.1	
Blood pressure increased	v.27.1	
Chest pain	v.27.1	
Chills	v.27.1	
Dizziness	v.27.1	
Nausea	v.27.1	
Palpitations	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Vomiting	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information **AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000703406	0	2017-09-27	2017-09-27	MAH	NGAM-046-17-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Informa	tion			
Age	Gender	Height	Weight	Report Outcome
70 Years	Male			Recovering/resolving

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report	

Product Information

i roduct illiorillation							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
HUMALOG	Concomitant	Injection					
INSULIN	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	50.0 Gram	1 every 1 Weeks		Myasthenia gravis

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Blood pressure increased	v.27.1	
Chest discomfort	v.27.1	
Headache	v.27.1	
Heart rate increased	v.27.1	
Malaise	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000703596	0	2017-10-03	2017-10-03	Hospital		Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Information

Age	Gender	Height	Weight	Report Outcome
	Female			Recovered/resolved

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	20.0 ml	Once		Primary immunodeficien cy syndrome

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Anaphylactic reaction	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000703841	0	2017-10-10	2017-10-10	Hospital		Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Not Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions:

Patient Information

Age	Gender	Height	Weight	Report Outcome
	Female			Recovered/resolved

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	200.0 ml			Abscess drainage
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	127.0 ml			Abscess drainage

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Chills	v.27.1	
Febrile nonhaemolytic transfusion reaction	v.27.1	
Heart rate increased	v.27.1	
Нурохіа	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000703969	0	2017-10-11	2017-10-11	MAH	NGAM-047-17-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient	Informati	tion

Age	Gender	Height	Weight	Report Outcome
65 Years	Female			Recovered/resolved

Link / Duplicate Report Information

Record Type Link AEF	R** Number
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No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
ACETAMINOPHEN	Concomitant	NOT SPECIFIED					
DIPHENHYDRAMINE	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)			148.0 Minutes	Neutropenia
PRIVIGEN	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	30.0 Gram	Once		Neutropenia

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Febrile nonhaemolytic transfusion reaction	v.27.1	
Headache	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information **AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000704117	0	2017-10-13	2017-10-13	MAH	NGAM-049-17-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Information				
Age	Gender	Height	Weight	Report Outcome
52 Years	Female			Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report	

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	2.0 Gram	Total	30.0 Minutes	Primary immunodeficien cy syndrome

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Anaphylactic reaction	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000704133	0	2017-10-13	2017-10-13	MAH	NGAM-051-17-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Informa	tion			
Age	Gender	Height	Weight	Report Outcome
61 Years	Male	163 Centimeter	82 Kilogram	Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report.	

Product Information

Froduct information							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
DEXTROSE	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	30.0 Gram	1 every 1 Months	90.0 Minutes	Secondary immunodeficien cy

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Febrile nonhaemolytic transfusion reaction	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000704164	0	2017-10-16	2017-10-16	Hospital		Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening: Yes	Hospitalization:	Other Medically Important Conditions:

Patient Informa	tion			
Age	Gender	Height	Weight	Report Outcome
55 Years	Female		81 Kilogram	Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report.	

Product Information

Froduct information							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	40.0 Gram	1 every 1 Months		Immunodeficien cy

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Anaphylactic reaction	v.27.1	
Chest pain	v.27.1	
Headache	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Informatio	n	**AE	R = Adverse F	Reaction Re	port							
Adverse Reaction Report Number		ER Version mber	Initial Recei	Received Date Latest Received		ived Date	Sourc	e of Report	Market Authorization Holder AER Number		Type of Report	Reporter Type
000704354		0	2017-10-19 2017-10-1		0-19		MAH	NGAM-053-17-CA		Spontaneous	Other health professional	
Serious report? Death:			Disability:			Congenital Anomaly:						
Seriou	ıs		Life Threatening:				Hospitalization: Other Mo			Other Med	dically Important C	onditions: Yes
Patient Information	n											
Age	Gender		Height	V	Veight		Report	Outcome				
54 Years	Female				Recovered/resolved							
Link / Duplicate R	eport Info	rmation										
	Recor	d Type				Link AER*	* Numb	er				
No duplicate or link	ed report.											
Product Informati	on											
Product Descr	ription	Health Pr	oduct Role	Dosag	e Form	Route Administr		Dose	Fre	equency	Therapy Duration	Indication(s)
PANZYGA		Sus	spect		JTION /ENOUS	Intravenou otherwi specifie	se	32.7 Gran	n	Once		Incisional drainage
Adverse Reaction Information	Term											
	Ad	verse Reac	tion Term(s)			Me	dDRA	Version			Reaction Duration	

v.27.1

Febrile nonhaemolytic transfusion reaction

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000704805	1	2017-10-27	2018-02-02	MAH	NGAM-055-17-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening: Yes	Hospitalization:	Other Medically Important Conditions:

	Patient	Informa	tion
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Age	Gender	Height	Weight	Report Outcome
55 Years	Female			Unknown

Link / Duplicate Report Information

Record Type	Link AER** Number
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No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	7.0 Gram			Immunodeficien cy

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Anaphylactic reaction	v.27.1	
Chest discomfort	v.27.1	
Depressed level of consciousness	v.27.1	
Swollen tongue	v.27.1	
Throat tightness	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information	

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000704894	0	2017-11-02	2017-11-02	Community		Spontaneous	Physician

Serious report?	Death:	Disability:	Congenital Anomaly:	
Serious	Life Threatening:	Hospitalization: Yes	Other Medically Important Conditions: Y	'es

Patient Information

Age	Gender	Height	Weight	Report Outcome	
56 Years	Female	150 Centimeter	48 Kilogram	Recovered/resolved	

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	15.0 Gram	1 every 28 Days		Immunoglobulin therapy

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Bronchospasm	v.27.1	
Chest discomfort	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000704895	0	2017-11-02	2017-11-02	Community		Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization: Yes	Other Medically Important Conditions: Yes

Patient Informa	tion			
Age	Gender	Height	Weight	Report Outcome
56 Years	Female	150 Centimeter	48 Kilogram	Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
Duplicate	000713840

Product Information							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise	15.0 Gram			Immunoglobulin therapy

Adverse Reaction Term Information			
Advers	e Reaction Term(s)	MedDRA Version	Reaction Duration
Headache		v.27.1	
Migraine		v.27.1	
Visual impairment		v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000704901	0	2017-10-31	2017-10-31	Hospital		Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Not Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions:

Patient Information	tion			
Age Gender		Height	Weight	Report Outcome
30 Years	Female	147 Centimeter	46 Kilogram	Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
Duplicate	000705574

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	30.0 Gram	1 every 4 Weeks		Immune system disorder

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Anxiety	v.27.1	
Back pain	v.27.1	
Chest pain	v.27.1	
Infusion related reaction	v.27.1	
Nausea	v.27.1	
Throat tightness	v.27.1	
Vomiting	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000704947	0	2017-11-01	2017-11-01	MAH	NGAM-050-17-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Informa	ition			
Age	Gender	Height	Weight	Report Outcome
30 Years	Male			Recovered/resolved

Link / Duplicate Report Information					
Record Type	Link AER** Number				
No duplicate or linked report.					

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)	
ACETAMINOPHEN	Concomitant	NOT SPECIFIED						
FOLIC ACID	Concomitant	NOT SPECIFIED						
PANTOPRAZOLE	Concomitant	NOT SPECIFIED						
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	90.0 Gram	Total	178.0 Minutes	Immune thrombocytopen ia	
PREDNISONE	Concomitant	NOT SPECIFIED		·				

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Pruritus	v.27.1	
Rash	v.27.1	
Urticaria	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000704990	1	2017-11-02	2018-01-05	MAH	NGAM-035-17-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Informa	tion			
Age	Gender	Height	Weight	Report Outcome
17 Years	Female			Unknown

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report	

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	17.7 Gram	Total	110.0 Minutes	Encephalomyeli tis

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Anaphylactoid reaction	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date	0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	Market Authorization Holder AER Number	Type of Report	Reporter Type
000705242	0	2017-11-08	2017-11-08	MAH	NGAM-058-17-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Information

Age	Gender	Height	Weight	Report Outcome
78 Years	Male			Recovered/resolved

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
AZITHROMYCIN	Concomitant	NOT SPECIFIED					
LOVENOX	Concomitant	SOLUTION SUBCUTANEOUS					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	16.0 Gram		80.0 Minutes	Secondary immunodeficien cy
TAZOCIN	Concomitant	POWDER FOR SOLUTION INTRAVENOUS					

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Blood pressure increased	v.27.1	
Chills	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Heart rate increased	v.27.1	
Respiratory distress	v.27.1	
Skin disorder	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Re	Adverse eaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
	000705251	0	2017-11-08	2017-11-08	MAH	NGAM-060-17-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Informa	tion					
Age Gender		Height	Weight	Report Outcome		
52 Years	Female	164 Centimeter	87 Kilogram	Recovered/resolved		

Link / Duplicate Report Information

Record Type

Link AER** Number

No duplicate or linked report.

Product Information

Product information							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	25.0 Gram	1 every 4 Weeks		Secondary immunodeficien cy

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Back pain	v.27.1	
Blood pressure increased	v.27.1	
Chest pain	v.27.1	
Chills	v.27.1	
Dizziness	v.27.1	
Heart rate increased	v.27.1	
Nausea	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Vomiting	v.27.1	

Canada Vigilanaa

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

Reaction Duration

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Canada vigilance	
Summary of Reported Adverse Reactions	•

Report Information	Report Information **AER = Adverse Reaction Report												
Adverse Reaction Report Number	Reaction Report Latest AER Version Initial Received D		ved Date	Latest Received Date		Source	e of Report	Market Authorization Holder AER Number		Type of Report	Reporter	Туре	
000705490	0		2017-11	-15	2017-1	1-15		MAH	NGAM	-068-17-CA	Spontaneous	Physic	ian
Serious re	eport?				Death:			Disabilit	y:		Congenital	Anomaly:	
Seriou	us		L	ife Threat	ening:		Н	ospitalizatio	n: Yes	Other Me	edically Important Co	onditions:	Yes
Patient Information	on												
Age	Gender		Height	V	Neight		Report	Outcome					
56 Years	Female	15	0 Centimeter	48	Kilogram	gram Recovered/resolved							
Link / Duplicate R	Report Inform	mation											
	Record	Туре				Link AER**	Numb	er					
No duplicate or link	ked report.												
Product Informati	ion											_	
Product Description Hea		Health Pr	oduct Role	Dosag	je Form	Route of Administra		Dose	Fr	equency	Therapy Duration	Indicati	on(s)
PANZYGA Su		Su	spect		UTION VENOUS	Intravenous otherwis specifie	se`	15.0 Gram	n 1 ev	ery 28 Days		Prima immunod cy synd	leficien
Adverse Reaction Information	n Term												

MedDRA Version

v.27.1

v.27.1

Adverse Reaction Term(s)

Bronchospasm

Chest discomfort

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000705574	0	2017-11-15	2017-11-15	MAH	NGAM-059-17-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Informa	tion			
Age Gender		Height	Weight	Report Outcome
30 Years	Female	147 Centimeter	46 Kilogram	Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
Duplicate	000704901

Product Information							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
DANIZVOA	0	SOLUTION	Intravenous (not	00.00	1 every 4		Immunodeficien

•			Aummstration			
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	30.0 Gram	1 every 4 Weeks	Immunodeficien cy common variable
Adverse Reaction Term						

mormation		
Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Anxiety	v.27.1	
Back pain	v.27.1	
Chest pain	v.27.1	
Nausea	v.27.1	
Throat tightness	v.27.1	
Vomiting	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information **AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000706202	0	2017-11-29	2017-11-29	MAH	NGAM-074-17-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization: Ye	Other Medically Important Conditions: Yes

Patient Informa	tion			
Age	Gender	Height	Weight	Report Outcome
19 Years	Male		60 Kilogram	Unknown

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report.	

Product Information

1 Todact Illiorillation							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
IBUPROFEN	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	60.0 Gram	Total	170.0 Minutes	Thrombocytope nia
VYVANSE	Concomitant	Capsules					

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Blood pressure diastolic decreased	v.27.1	
CSF protein increased	v.27.1	
Headache	v.27.1	
Meningitis aseptic	v.27.1	
Nausea	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Vomiting	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

6	Report Informa
	Adverse Reaction Repo Number
	000706386
	Serious
	Ser
F	Patient Informa
	Age
	62 Years
L	ink / Duplicate

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000706386	1	2017-12-05	2017-12-20	MAH	NGAM-075-17-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Information					
	Age	Gender	Height	Weight	Report Outcome
	62 Years	Male			Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report	

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	8.7 Gram		95.0 Minutes	Product used for unknown indication

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Alpha haemolytic streptococcal infection	v.27.1	
Pyrexia	v.27.1	
Suspected transmission of an infectious agent via product	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Repo Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000706546	1	2017-12-08	2018-05-28	Hospital		Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Not Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions:

Patient Information				
Age	Gender	Height	Weight	Report Outcome
50 Years	Male			Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report.	

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	90.0 Gram	1 every 3 Weeks	75.0 Minutes	Immunodeficien cy common variable

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Blood culture negative	v.27.1	
Chills	v.27.1	
Hypertension	v.27.1	
Infusion related reaction	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Re	port	Infori	matio	n

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000706563	0	2017-12-11	2017-12-11	Hospital		Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Not Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions:

Patient	Information

Age	Gender	Height	Weight	Report Outcome
6 Years	Male	120 Centimeter	20 Kilogram	Recovered/resolved

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
ACETAMINOPHEN	Concomitant	NOT SPECIFIED					
ACETYLSALICYLIC ACID	Concomitant	NOT SPECIFIED					
AMOXICILLIN	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	40.0 Gram			Kawasaki's disease

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Bradycardia	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000707175	0	2017-12-22	2017-12-22	MAH	NGAM-077-17-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Informa	ation			
Age	Gender	Height	Weight	Report Outcome
6 Vears	Male	120 Centimeter	20 Kilogram	Unknown

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report.	

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
ACETAMINOPHEN	Concomitant	NOT SPECIFIED					
ACETYLSALICYLIC ACID	Concomitant	NOT SPECIFIED					
AMOXICILLIN	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	30.0 Gram			Kawasaki's disease

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Blood lactate dehydrogenase increased	v.27.1	
Bradycardia	v.27.1	
Heart rate irregular	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Informat
Adverse Reaction Repo Number
000709078
Serious
Seri
Patient Informa
Age
9 Years
Link / Duplicate
No duplicate or li
Product Informa
Product Des
PANZYGA

	Report Information	**A	ER = Adverse Reaction Re	eport				
Reaction Report		Latest AER Versio Number	n Initial Received Date	Latest Received Date	Source of Report	Market Authorization Holder AER Number	Type of Report	Reporter Type
	000709078	4	2018-02-05	2018-09-27	MAH	NGAM-002-18-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability		Congenital Anomaly:	
Serious	Life Threatening:	Hospitalization	Yes	Other Medically Important Conditions:	Yes

Patient informa	tion			
Age Gender		Height	Weight	Report Outcome
9 Years	Female			Unknown

Link / Duplicate Report Information	1
Record Type	Link AER** Number
No duplicate or linked report	

nation

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	57.0 Gram	Total		Kawasaki's disease

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Abdominal pain	v.27.1	
Anti A antibody positive	v.27.1	
Blood urine present	v.27.1	
Haemoglobin decreased	v.27.1	
Haemolysis	v.27.1	
Headache	v.27.1	
Oedema	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Pyrexia	v.27.1	
Reticulocyte count increased	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Summary of Reported Adverse Reaction

Anaphylactic transfusion reaction

Blood pressure increased

Respiratory rate increased

Report Informatio	n	**AE	R = Adverse F	Reaction Re	eport							
Adverse Reaction Report Number	action Report Latest AER Version Initial Received Date Latest Received Date Sou		Sourc	e of Report	Market Authorization Holder AER Number		Type of Report	Reporter Type				
000709440		0	2018-0	2-13	2018-0	2-13	MAH NGAM-		011-18-CA	Spontaneous	Other health professional	
Serious re	eport?				Death:			Disabilit	y:		Congenital	Anomaly:
Seriou	JS		l	ife Threat	ening:		Н	ospitalizatio	n:	Other Med	dically Important C	onditions: Yes
Patient Information	on											
Age	Gender		Height	V	Veight		Report	Outcome				
66 Years	Male					F	Recover	ed/resolved				
Link / Duplicate R	eport Info	rmation										
•		d Type				Link AER*	* Numb	er				
No duplicate or link	ked report.											
Product Informati	ion											
Product Descr	ription	Health Pr	oduct Role	Dosag	je Form	Route Administr		Dose	Fr	equency	Therapy Duration	Indication(s)
PANZYGA		Sus	spect		JTION /ENOUS	Intravenou otherwi specifie	se	70.0 Gran	า	Total	4.0 Hours	Chronic inflammatory demyelinating polyradiculoneu ropathy
Adverse Reaction Information	Term											
	Adv	verse Reac	tion Term(s)			Me	dDRA	Version			Reaction Duration	

v.27.1

v.27.1

v.27.1

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

ia

Report Information **AER = Adverse Reaction Report

Adverse Reaction Term

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000709443	0	2018-02-14	2018-02-14	MAH	NGAM-015-18-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Information				
Age Gender		Height Weight		Report Outcome
6 Years	Female			Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report.	

Product Information Route of **Product Description Health Product Role Dosage Form Therapy Duration** Indication(s) Dose **Frequency** Administration Intravenous (not Immune SOLUTION PANZYGA Suspect otherwise 6.0 Hours thrombocytopen 20.0 Gram Total INTRAVENOUS specified)

Information			
Adverse	e Reaction Term(s)	MedDRA Version	Reaction Duration
Meningitis asentic		v 27 1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

|--|

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000709474	0	2018-02-16	2018-02-16	MAH	NGAM-019-18-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization: Yes	Other Medically Important Conditions:

Patient	Information

Age	Gender	Height	Weight	Report Outcome
77 Years	Male	178 Centimeter	75 Kilogram	Recovered/resolved

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

i roduct imormation							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
ALBUTEROL	Concomitant	NOT SPECIFIED					
CALCIUM	Concomitant	NOT SPECIFIED					
LORAZEPAM	Concomitant	NOT SPECIFIED					
METOCLOPRAMIDE	Concomitant	NOT SPECIFIED					
PANTOPRAZOLE	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	175.0 Gram	Total	135.0 Minutes	Immune thrombocytopen ia
POLYETHYLENE GLYCOL	Concomitant	NOT SPECIFIED					
PREDNISONE	Concomitant	NOT SPECIFIED					
RISEDRONATE (MANUFACTURER UNKNOWN)	Concomitant	NOT SPECIFIED					
SALBUTAMOL	Concomitant	NOT SPECIFIED					

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
SEEBRI BREEZHALER	Concomitant	Capsules					
TACROLIMUS	Concomitant	Capsules					

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Abdominal pain	v.27.1	
Aortic arteriosclerosis	v.27.1	
Blood pressure increased	v.27.1	
Dyspnoea	v.27.1	
Erythema	v.27.1	
Lung hyperinflation	v.27.1	
Pain	v.27.1	
Rash	v.27.1	
Tachycardia	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Repor Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000709512	0	2018-02-20	2018-02-20	MAH	NGAM-020-18-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

F	Patient Informa	tion			
	Age	Gender	Height	Weight	Report Outcome
	15 Years	Male			Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report	

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	45.0 Gram	1 every 1 Months	4.0 Hours	Immunodeficien cy

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Headache	v.27.1	
Neck pain	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000710295	0	2018-03-07	2018-03-07	MAH	NGAM-027-18-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient	Informa	tion

Age	Gender	Height	Weight	Report Outcome
73 Years	Male			Unknown

Link / Duplicate Report Information

Record Type	Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
LORAZEPAM	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	25.0 Gram	1 every 3 Weeks		Plasma cell myeloma

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Atrial flutter	v.27.1	
Chills	v.27.1	
Dyspnoea	v.27.1	
Hyperhidrosis	v.27.1	
Hypertension	v.27.1	
Oxygen saturation decreased	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Tachycardia	v.27.1	
Tachypnoea	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000711152	0	2018-03-28	2018-03-28	MAH	NGAM-039-18-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient	Informa	tion

Age	Gender	Height	Weight	Report Outcome
78 Years	Female		44 Kilogram	Recovered/resolved

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	20.0 Gram	1 every 4 Weeks		Hypogammaglo bulinaemia

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Asthenia	v.27.1	
Discomfort	v.27.1	
Feeling abnormal	v.27.1	
Headache	v.27.1	
Heart rate increased	v.27.1	
Hypertension	v.27.1	
Oxygen saturation decreased	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000711158	0	2018-03-28	2018-03-28	MAH	NGAM-037-18-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Informa	ition			
Age Gender		Height	Weight	Report Outcome
78 Years	Male	179 Centimeter	80 Kilogram	Unknown

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report.	

Product Information

Froduct information							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)				Immune thrombocytopen ia

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Anti A antibody positive	v.27.1	
Blood lactate dehydrogenase increased	v.27.1	
Coombs direct test positive	v.27.1	
Delayed haemolytic transfusion reaction	v.27.1	
Haemoglobin decreased	v.27.1	
Reticulocyte count increased	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000711296	0	2018-04-03	2018-04-03	MAH	NGAM-042-18-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

|--|

Age	Gender	Height	Weight	Report Outcome
82 Years	Male		73 Kilogram	Recovered/resolved

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	30.0 Gram	Total	75.0 Minutes	Rheumatoid arthritis

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Blood pressure increased	v.27.1	
Chest discomfort	v.27.1	
Chest pain	v.27.1	
Chills	v.27.1	
Febrile nonhaemolytic transfusion reaction	v.27.1	
Injection site erythema	v.27.1	
Pyrexia	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Tremor	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000711504	0	2018-04-06	2018-04-06	Hospital		Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Not Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions:

Patient Information					
	Age Gender		er Height Weight		Report Outcome
	66 Years	Female			Unknown

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report.	

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)				
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	80.0 ml	Once		

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Febrile nonhaemolytic transfusion reaction	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000711630	0	2018-04-10	2018-04-10	Hospital		Spontaneous	Physician

Serious report?	Death:	Disability:	Congenital Anomaly:	
Not Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions:	

Patient Information

- L	- australia				
	Age Gender		Height	Weight	Report Outcome
	66 Years	Male	190 Centimeter	126 Kilogram	Recovered/resolved

Link / Duplicate Report Information

Record Type	Link AER** Number

No duplicate or linked report.

Product Information

1 Todaot Illiorillation							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
ACETAMINOPHEN	Concomitant	NOT SPECIFIED					
BENADRYL	Concomitant	Capsules					
GRAVOL	Concomitant	Tablets					
NAPROXEN	Concomitant	Tablets					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	300.0 ml	Once		
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	300.0 ml	Once		
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	300.0 ml	Once		

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	100.0 ml	Once		
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	300.0 ml	Once		

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Hypersensitivity	v.27.1	
Injection site rash	v.27.1	
Urticaria	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000712310	0	2018-04-17	2018-04-17	MAH	NGAM-049-18-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization: Yes	Other Medically Important Conditions: Yes

Patient	Informa	tion	

Age	Gender	Height	Weight	Report Outcome
83 Years	Male		91 Kilogram	Unknown

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	35.0 Gram	Total	90.0 Minutes	Immune thrombocytopen ia

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Blood pressure increased	v.27.1	
Chills	v.27.1	
Dyspnoea	v.27.1	
Hypothermia	v.27.1	
Oxygen saturation decreased	v.27.1	
Tremor	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information	on	**AE	R = Adverse R	eaction Re	eport							
Adverse Reaction Report Number	Latest AE Nun	R Version nber	Initial Recei	ved Date	Latest Rece	ived Date	Sourc	e of Report	Market Authorization Holder AER Number		Type of Report	Reporter Type
000712582	1	1	2018-04	l-30	2018-06-04			МАН	NGAM-	014-18-CA	Spontaneous	Other health professional
Serious re	eport?				Death:			Disabilit	y:		Congenital	Anomaly:
Serio	us		L	ife Threat	ening:		Н	ospitalizatio	n: Yes	Other Med	dically Important Co	onditions: Yes
Patient Information	on .											
Age	Gender		Height	V	Veight		Report	Outcome				
13 Months	Male		<u> </u>		Kilogram			ed/resolved				
Link / Duplicate R	Report Infor	mation										
	Record					Link AER*	* Numb	er				
No duplicate or linl	ked report.											
Product Informat	ion											
Product Desc	ription	Health Pr	oduct Role	Dosag	je Form	Route Administr		Dose	Fre	equency	Therapy Duration	Indication(s)
PANZYGA		Sus	spect		JTION /ENOUS	otherwi	Intravenous (not otherwise 17.5 Gram specified)		n	Total		Kawasaki's disease
Adverse Reaction Information	n Term											
Adverse Reaction Term(s)				Me	dDRA	Version			Reaction Duration			

v.27.1

Haemoglobin decreased

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information						
Adverse Reaction Report Number	Latest AER Num					
222712221						

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000712901	1	2018-05-07	2018-05-23	MAH	NGAM-054-18-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:	
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions:	Yes

Patient	Informati	tion

Age	Gender	Height	Weight	Report Outcome
79 Years	Male	173 Centimeter	63 Kilogram	Recovered/resolved

Link / Duplicate Report Information

Record Type Link AER Number**

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	30.0 Gram	1 every 5 Weeks	94.0 Minutes	Primary immunodeficien cy syndrome

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Blood pressure increased	v.27.1	
Chills	v.27.1	
Dyspnoea	v.27.1	
Febrile nonhaemolytic transfusion reaction	v.27.1	
Oxygen saturation decreased	v.27.1	
Productive cough	v.27.1	
Respiratory rate increased	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Tachycardia	v.27.1	
Tremor	v.27.1	
Wheezing	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31

173 Report(s)

Report Information **AER = Adverse Reaction Report								
	Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
	000713126	1	2018-05-08	2018-09-27	MAH	NGAM-053-18-CA	Spontaneous	Physician

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Informa	tion			
Age	Gender	Height	Weight	Report Outcome
64 Years	Female	165 Centimeter	54 Kilogram	Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report.	

Product Information					
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Freq

Product Description	Health Product Role	Dosage Form	Administration	Dose	Frequency	Therapy Duration	Indication(s)
ALENDRONATE	Concomitant	Tablets		_			
APO-FERROUS SULFATE	Concomitant	TABLET (ENTERIC- COATED)					
CALCIUM	Concomitant	NOT SPECIFIED					
COUMADIN	Concomitant	Tablets					
DIPHENHYDRAMINE	Concomitant	NOT SPECIFIED					
DOCUSATE SODIUM	Concomitant	NOT SPECIFIED					
HYDROXYCHLOROQUINE	Concomitant	NOT SPECIFIED					
METHYLPREDNISOLONE NOS	Concomitant	NOT SPECIFIED					
OXAZEPAM	Concomitant	Tablets					
PANTOPRAZOLE	Concomitant	NOT SPECIFIED					

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	20.0 Gram	Total	230.0 Minutes	Hodgkin's disease
PREDNISONE	Concomitant	NOT SPECIFIED					
SENNOSIDES	Concomitant	NOT SPECIFIED		_			
VITAMIN D	Concomitant	NOT SPECIFIED					

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Abdominal pain	v.27.1	
Back pain	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000713150	0	2018-05-09	2018-05-09	MAH	NGAM-058-18-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization: Yes	Other Medically Important Conditions:

Patient	Informa	tion

Age	Gender	Height	Weight	Report Outcome
52 Years	Male	173 Centimeter	88 Kilogram	Recovered/resolved

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
ACETAMINOPHEN	Concomitant	NOT SPECIFIED					
DIPHENHYDRAMINE	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)				Chronic inflammatory demyelinating polyradiculoneu ropathy

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Chills	v.27.1	
Heart rate increased	v.27.1	
Hypotension	v.27.1	
Loss of consciousness	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Oxygen saturation decreased	v.27.1	
Pyrexia	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Informatio	n	**AEI	R = Adverse	Reaction Re	port
Adverse					

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000713303	0	2018-05-14	2018-05-14	MAH	NGAM-064-18-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient	Informa	tion

Age	· G	Sender	Height	Weight	Report Outcome
52 Ye	ars	Male		89 Kilogram	Recovered/resolved

|--|

Record Type	Link AER** Number
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No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
METHYLPREDNISOLONE NOS	Concomitant	NOT SPECIFIED					
PANTOPRAZOLE	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	180.0 Gram	Total		Immune thrombocytopen ia
SULFAMETHOXAZOLE AND TRIMETHOPRIM	Concomitant	NOT SPECIFIED					

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Anaemia	v.27.1	
Coombs test positive	v.27.1	
Haemoglobin decreased	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Haemolysis	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information	**AER = Adverse Reaction Report											
Adverse Reaction Report Number	Latest AEI Num		ion Initial Received Date La		Latest Rece	Latest Received Date		e of Report	Market Authorization Holder AER Number		Type of Report	Reporter Type
000713357	2		2018-05	-15	2018-08-03			MAH	NGAM-066-18-CA		Spontaneous	Other health professional
Serious report?					Death:			Disabilit	y:		Congenital	Anomaly:
Serio	JS		Li	fe Threat	ening:		H	ospitalizatio	n:	Other Med	dically Important Co	onditions: Yes
Patient Information												
Age Gender Height		l v	Weight		Report	Outcome						
74 Years				Kilogram	Recovered/resolved							
Link / Duplicate R	eport Infor	mation										
	Record					Link AER*	* Numb	er				
No duplicate or link	ked report.			•								
Product Informati	ion											
Product Desc	ription	Health Pr	oduct Role	Dosag	ge Form	Route d Administr		Dose	Fre	equency	Therapy Duration	Indication(s)
PANZYGA S		Sus	spect		UTION VENOUS	Intravenous otherwis specifie	se	80.0 Gran		every 4 Neeks		Immune thrombocytopen ia
Adverse Reaction Information	Term											
Adverse Reaction Term(s)			Μe	dDRA '	Version			Reaction Duration				

Blood pressure increased

Haemoglobinuria

v.27.1

v.27.1

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Informa
Adverse Reaction Repo Number
000713544
Serious
Not S
Patient Informa
Age
40 Years
Link / Duplicate
No duplicate or I
Product Inform
Product Des
DIPHENHYDRA
HYDROCORTIS
PANZYGA

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000713544	1	2018-05-22	2018-05-30	MAH	NGAM-070-18-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Not Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions:

Patient Informa	tion					
Age	Gender	Height	Weight	Report Outcome		
40 Years	Female		72 Kilogram	Recovered/resolved		

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report.	

ation

i roduct imormation							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
DIPHENHYDRAMINE	Concomitant	NOT SPECIFIED					
HYDROCORTISONE	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	40.0 Gram	Total		Clostridium difficile colitis

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Body temperature increased	v.27.1	
Chills	v.27.1	
Condition aggravated	v.27.1	
Face oedema	v.27.1	
Oropharyngeal pain	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Tachycardia	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000716091	0	2018-07-25	2018-07-25	MAH	NGAM-093-18-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Informa	tion			
Age Gender		Height	Weight	Report Outcome
17 Years	Female	167 Centimeter	80 Kilogram	Recovering/resolving

Link / Duplicate Report Information	
Record Type	Link AER** Number
Nie de Perte en Pele de sent	

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
ADCETRIS	Concomitant	POWDER FOR SOLUTION INTRAVENOUS					
FENTANYL	Concomitant	NOT SPECIFIED					
LORAZEPAM	Concomitant	NOT SPECIFIED					
MIDAZOLAM (UNKNOWN)	Concomitant						
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	55.0 Gram	1 every 1 Days	160.0 Minutes	Guillain-Barre syndrome

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Blood pressure diastolic decreased	v.27.1	
Chills	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Headache	v.27.1	
Heart rate increased	v.27.1	
Meningitis aseptic	v.27.1	
Nausea	v.27.1	
Pyrexia	v.27.1	
Vomiting	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Reaction	verse on Report imber	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
0007	717005	0	2018-08-22	2018-08-22	MAH	NGAM-105-18-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Informa	tion			
Age	Gender	Height	Weight	Report Outcome
48 Years	Female			Unknown

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report.	

Product Information

Froduct information							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	50.0 Gram	Total	85.0 Minutes	Urticaria

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Back pain	v.27.1	
Chills	v.27.1	
Condition aggravated	v.27.1	
Febrile nonhaemolytic transfusion reaction	v.27.1	
Headache	v.27.1	
Heart rate increased	v.27.1	
Hypertension	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Oropharyngeal pain	v.27.1	
Urticaria	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000717114	0	2018-08-24	2018-08-24	MAH	NGAM-112-18-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Informa	tion			
Age	Gender	Height	Weight	Report Outcome
64 Years	Male			Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report	

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	50.0 Gram	1 every 1 Months		Secondary immunodeficien cy

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Chills	v.27.1	
Condition aggravated	v.27.1	
Febrile nonhaemolytic transfusion reaction	v.27.1	
Headache	v.27.1	
Hypertension	v.27.1	
Hyperthermia	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date	Source of Report	Market Authorization Holder AER Number	Type of Report	Reporter Type
000717119	1	2018-08-24	2018-10-04	MAH	NGAM-108-18-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Informa	tion			
Age	Gender	Height	Weight	Report Outcome
74 Years	Male			Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report.	

Product Information

Froduct information							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	80.0 Gram	Total	1.0 Hours	Hypogammaglo bulinaemia

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Back pain	v.27.1	
Chills	v.27.1	
Haemolysis	v.27.1	
Oxygen saturation decreased	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Reactio	verse on Report mber	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
0007	17121	0	2018-08-24	2018-08-24	MAH	NGAM-114-18-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Informa	tion			
Age	Gender	Height	Weight	Report Outcome
82 Years	Female			Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report.	

Product Information

Froduct information								
	Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
	PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	25.0 Gram	Total		Secondary immunodeficien cy

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Abdominal pain upper	v.27.1	
Back pain	v.27.1	
Hypertension	v.27.1	
Tremor	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000717127	0	2018-08-24	2018-08-24	MAH	NGAM-109-18-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Informa	tion			
Age	Gender	Height	Weight	Report Outcome
20 Months	Mala			Linknown

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report.	

Product Information

Product information							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
ACETAMINOPHEN	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	125.0 ml	Total	96.0 Minutes	Immune thrombocytopen ia

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Chills	v.27.1	
Febrile nonhaemolytic transfusion reaction	v.27.1	
Oxygen saturation decreased	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000717454	0	2018-09-04	2018-09-04	MAH	NGAM-115-18-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Informa	tion			
Age Gender		Height	Weight	Report Outcome
83 Years	Female	152 Centimeter	49 Kilogram	Unknown

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report.	

Product Information

Product information							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
DIPHENHYDRAMINE	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	15.0 Gram	1 every 1 Months		Hypogammaglo bulinaemia

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Back pain	v.27.1	
Body temperature increased	v.27.1	
Chills	v.27.1	
Hypertension	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000718185	2	2018-09-25	2018-10-25	MAH	NGAM-121-18-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization: Ye	Other Medically Important Conditions: Yes

Patient Information	tion			
Age	Gender	Height	Weight	Report Outcome
59 Years	Male			Unknown

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report	

Product Information

1 Todact Illioilliation							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
ACETAMINOPHEN	Concomitant	NOT SPECIFIED					
ACETYLSALICYLIC ACID	Concomitant	NOT SPECIFIED					
CITALOPRAM	Concomitant	Tablets					
CLOPIDOGREL	Concomitant	Tablets					
DEXLANSOPRAZOLE	Concomitant						
DIMENHYDRINATE	Concomitant	NOT SPECIFIED					
IBUPROFEN	Concomitant	NOT SPECIFIED					
INSULIN	Concomitant	NOT SPECIFIED					
JARDIANCE	Concomitant						
LEVOTHYROXINE	Concomitant	NOT SPECIFIED					
LINAGLIPTIN	Concomitant						
METFORMIN	Concomitant	Tablets					

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
METOCLOPRAMIDE	Concomitant	NOT SPECIFIED					
METOPROLOL	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	40.0 Gram	1 every 1 Days		Encephalitis autoimmune
PERINDOPRIL	Concomitant	NOT SPECIFIED					

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Arteriovenous fistula	v.27.1	
Cerebrovascular accident	v.27.1	
Hemiparesis	v.27.1	
Pain	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Informatio	n	**AE	R = Adverse F	Reaction Re	eport							
Adverse Reaction Report Number	Latest AEI Num		Initial Recei	ved Date	Latest Rece	ived Date	Sourc	e of Report	Auth	arket orization ER Number	Type of Report	Reporter Type
000718804	1		2018-1	0-11	2018-1	1-05		MAH	NGAM-	130-18-CA	Spontaneous	Other health professional
Serious re	eport?			l	Death:			Disabilit	y:		Congenital	Anomaly:
Seriou	ıs		L	ife Threat	ening:		Н	ospitalizatio	n:	Other Med	dically Important Co	onditions: Yes
Patient Information	on .											
Age	Gender		Height	\	Weight		Report	Outcome				
66 Years	Male					F	Recover	ed/resolved				
Link / Duplicate R	eport Infor	mation										
	Record	d Type				Link AER*	* Numb	er				
No duplicate or link	ked report.											
Product Informati	on											
Product Descr	ription	Health Pr	oduct Role	Dosag	ge Form	Route Administr		Dose	Fre	equency	Therapy Duration	Indication(s)
PANZYGA		Su	spect		UTION VENOUS	Intravenou otherwi specifie	se	65.0 Gram	1	Total	290.0 Minutes	Immune thrombocytopen ia
Adverse Reaction Information	Term											
	Adv	erse Reac	tion Term(s)			Me	dDRA	Version			Reaction Duration	
Anti A antibody pos	sitive						v.27	.1				

Haemoglobin decreased

Transfusion reaction

Seroconversion test positive

Hepatitis B core antibody positive

v.27.1

v.27.1

v.27.1

v.27.1

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000719429	0	2018-10-29	2018-10-29	MAH	NGAM-147-18-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization: Yes	Other Medically Important Conditions:

Patient Informa	tion			
Age	Gender	Height	Weight	Report Outcome
69 Years	Female			Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report.	

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)		
ATORVASTATIN CALCIUM	Concomitant	Tablets							
DOCUSATE SODIUM	Concomitant	NOT SPECIFIED							
ENOXAPARIN	Concomitant	Injection							
HYDROCORTISONE	Concomitant	NOT SPECIFIED							
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	54.0 Gram	Total	105.0 Minutes	Mononeuropath y multiplex		
POLYETHYLENE GLYCOL	Concomitant	NOT SPECIFIED							
SENNOSIDES	Concomitant	NOT SPECIFIED							

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Body temperature increased	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Chills	v.27.1	
Hypertension	v.27.1	
Tachycardia	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000720047	1	2018-11-21	2019-08-14	MAH	NGAM-168-18-CA	Spontaneous	Physician

Serious report?	Death:	Yes	Disability:	Congenital Anomaly:	
Serious	Life Threatening:		Hospitalization:	Other Medically Important Conditions: Yes	

Patient Information

Age	Gender	Height	Weight	Report Outcome
67 Years	Female	154 Centimeter	76 Kilogram	Fatal

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
CEFPROZIL	Concomitant	NOT SPECIFIED					
CEFTRIAXONE FOR INJECTION USP	Concomitant	POWDER FOR SOLUTION INTRAMUSCULAR					
DOXYCYCLINE	Concomitant	NOT SPECIFIED					
MEROPENEM	Concomitant	POWDER FOR SOLUTION INTRAVENOUS					
PANTOPRAZOLE	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)				Neuropathy peripheral
PREGABALIN	Concomitant	Capsules					

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Blood pressure increased	v.27.1	
Body temperature increased	v.27.1	
Chills	v.27.1	
Dyspnoea	v.27.1	
Nausea	v.27.1	
Pyrexia	v.27.1	
Respiratory rate increased	v.27.1	
Tachycardia	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000720672	0	2018-12-11	2018-12-11	MAH	NGAM-153-18-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Informa	tion			
Age	Gender	Height	Weight	Report Outcome
78 Years	Female			Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report.	

Product Information

1 Todaot IIII of III ation							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	65.0 Gram	Total		Thrombocytope nic purpura
PLATELETS	Suspect	NOT SPECIFIED	Unknown	250.0 ml	Total		Product used for unknown indication

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Anaemia	v.27.1	
Chills	v.27.1	
Confusional state	v.27.1	
Faeces discoloured	v.27.1	
Haemoglobinuria	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Haemoptysis	v.27.1	
Pyrexia	v.27.1	
Tachycardia	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000721026	0	2018-12-20	2018-12-20	MAH	NGAM-183-18-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Informa	tion			
Age	Gender	Height	Weight	Report Outcome
73 Years	Female	158 Centimeter	93 Kilogram	Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report.	

Product Information

Froduct information							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	5.0 Gram	Total		Hypogammaglo bulinaemia

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Blood pressure decreased	v.27.1	
Chest pain	v.27.1	
Erythema	v.27.1	
Hypertension	v.27.1	
Нурохіа	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Re Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
00072143	2 0	2019-01-08	2019-01-08	MAH	NGAM-188-18-CA	Spontaneous	Physician

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Information

Age	Gender	Height	Weight	Report Outcome	
83 Years	Female		49 Kilogram	Recovered/resolved	

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Froduct information							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
BISOPROLOL	Concomitant	Tablets					
CALCIUM	Concomitant	NOT SPECIFIED					
PANTOPRAZOLE	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	20.0 Gram	1 every 4 Weeks	80.0 Minutes	Hypogammaglo bulinaemia
VITAMIN D	Concomitant	NOT SPECIFIED					

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Back pain	v.27.1	
Chills	v.27.1	
Cyanosis	v.27.1	
Hypersensitivity	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Hypertension	v.27.1	
Нурохіа	v.27.1	
Tachycardia	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Re Number	Dort Latest AER Version	Initial Received Date	Latest Received Date	•	Market Authorization Holder AER Number	Type of Report	Reporter Type
00072243	0	2019-02-05	2019-02-05	MAH	NGAM-012-19-CA	Spontaneous	Physician

Serious report?Death:Disability:Congenital Anomaly:SeriousLife Threatening:Hospitalization:Other Medically Important Conditions:Yes

Patient Information

Age Gender Height		Weight	Report Outcome	
65 Years	Female	172 Centimeter	75 Kilogram	Recovered/resolved

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
ACETAMINOPHEN	Concomitant	NOT SPECIFIED					
DEXAMETHASONE	Concomitant	NOT SPECIFIED					
DICYCLOMINE HYDROCHLORIDE USP	Concomitant	NOT SPECIFIED					
DIPHENHYDRAMINE	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	140.0 Gram	Total		Muscular weakness
PYRIDOSTIGMINE BROMIDE	Concomitant	NOT SPECIFIED					
RIVAROXABAN (BAY59- 7939) VS. ASA	Concomitant						
SYMBICORT	Concomitant	Powder					

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Abdominal pain upper	v.27.1	
Asthenia	v.27.1	
Chest pain	v.27.1	
Chills	v.27.1	
Drug intolerance	v.27.1	
Electrocardiogram T wave abnormal	v.27.1	
Fatigue	v.27.1	
Headache	v.27.1	
Hypertension	v.27.1	
Pallor	v.27.1	
Syncope	v.27.1	
Tremor	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000722754	1	2019-02-14	2019-04-10	MAH	NGAM-016-19-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:		Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Yes	Other Medically Important Conditions:

Patient Information

Age	Gender	Height	Weight	Report Outcome
49 Years	Female		74 Kilogram	Unknown

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	74.0 Gram	Total		Toxic shock syndrome

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Anxiety	v.27.1	
Chest pain	v.27.1	
Chills	v.27.1	
Cough	v.27.1	
Dysphagia	v.27.1	
Dyspnoea	v.27.1	
Hypersensitivity	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Hypotension	v.27.1	
Oxygen saturation decreased	v.27.1	
Pulmonary oedema	v.27.1	
Tachycardia	v.27.1	
Tachypnoea	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000722903	0	2019-02-20	2019-02-20	MAH	NGAM-023-19-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Information						
Age Gender		Height Weight		t Report Outcome		
61 Years	Female		48 Kilogram	Recovered/resolved		

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report.	

Product Information

Froduct information							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	50.0 Gram			Immune thrombocytopen ia

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Blood pressure increased	v.27.1	
Feeling cold	v.27.1	
Headache	v.27.1	
Meningitis aseptic	v.27.1	
Neck pain	v.27.1	
Pyrexia	v.27.1	
Tremor	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000722907	0	2019-02-21	2019-02-21	MAH	NGAM-017-19-CA	Spontaneous	Physician

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Information

Age Gender		Height Weight		Report Outcome	
15 Years	15 Years Female 161 Centimeter		87 Kilogram	Recovered/resolved	

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	30.0 Milligram	Total		Immune thrombocytopen ia

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Chills	v.27.1	
Dyspnoea	v.27.1	
Febrile nonhaemolytic transfusion reaction	v.27.1	
Headache	v.27.1	
Hypertension	v.27.1	
Loss of consciousness	v.27.1	
Nausea	v.27.1	
Oxygen saturation decreased	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Vomiting	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000723306	0	2019-03-01	2019-03-01	MAH	NGAM-030-19-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Information

Age	Gender	Height	Weight	Report Outcome
56 Years	Male		100 Kilogram	Recovered/resolved

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

1 Todact Illioilliation							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
ALTEPLASE	Concomitant	LIQUID INTRAVENOUS					
CEFTRIAXONE SODIUM FOR INJECTION BP	Concomitant						
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	50.0 Gram	Total		Immunodeficien cy
PIPERACILLIN/TAZOBACT AM	Concomitant	NOT SPECIFIED					
PREDNISONE	Concomitant	NOT SPECIFIED					
VANCOMYCIN	Concomitant	NOT SPECIFIED					

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Anaphylactic reaction	v.27.1	
Blood pressure increased	v.27.1	
Chills	v.27.1	
Dyspnoea	v.27.1	
Hyperhidrosis	v.27.1	
Oxygen saturation decreased	v.27.1	
Pyrexia	v.27.1	
Swelling of eyelid	v.27.1	
Tachycardia	v.27.1	
Tremor	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000723342	0	2019-03-06	2019-03-06	MAH	NGAM-036-19-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient	Informati	tion

Age	Gender	Height	Weight	Report Outcome
56 Years	Female	147 Centimeter	84 Kilogram	Recovered/resolved

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

i roduct iiiioiiiiatioii							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	30.0 Gram	1 every 4 Weeks		Hypogammaglo bulinaemia

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Chills	v.27.1	
Erythema	v.27.1	
Febrile nonhaemolytic transfusion reaction	v.27.1	
Hypertension	v.27.1	
Hyperthermia	v.27.1	
Hypopnoea	v.27.1	
Tachycardia	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000723524	2	2019-03-11	2019-04-29	MAH	NGAM-035-19-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability: Yes	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization: Yes	Other Medically Important Conditions: Yes

Patient	Informa	tion

Age	Gender	Height	Weight	Report Outcome
29 Years	Female	170 Centimeter	87 Kilogram	Unknown

Link / Duplicate Report Information

Record Type	Link AER** Number
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No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
DEXAMETHASONE	Concomitant	NOT SPECIFIED					
ECHINACEA	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	85.0 Gram	Total		Immune thrombocytopen ia
VITAMIN C	Concomitant	NOT SPECIFIED					

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Anaemia	v.27.1	
Anti A antibody positive	v.27.1	
Body temperature increased	v.27.1	
Delayed haemolytic transfusion reaction	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Drug intolerance	v.27.1	
Jaundice	v.27.1	
Malaise	v.27.1	
Pain in extremity	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000723799	2	2019-03-19	2019-08-18	MAH	NGAM-040-19-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Informa		
Δαο	Gender	Height

Age	Gender	Height	Weight	Report Outcome
84 Years	Female	148 Centimeter	53 Kilogram	Recovered/resolved

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	20.0 Gram	1 every 6 Weeks		Secondary immunodeficien cy

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Chills	v.27.1	
Hypertension	v.27.1	
Нурохіа	v.27.1	
Tremor	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Canada Vigilance
Summary of Reported Adverse Reactions

Report Informatio	n	**AEI	R = Adverse R	eaction Re	eport							
Adverse Reaction Report Number	Latest AE Num		Initial Receiv	ed Date	Latest Rece	ived Date	Sourc	e of Report	Autho	arket orization ER Number	Type of Report	Reporter Type
000725335	()	2019-04	-25	2019-0	4-25	Н	lospital			Spontaneous	Physician
Serious re	eport?			ı	Death:			Disabilit	y:	Congenital Anomaly:		Anomaly:
Seriou	ıs		L	ife Threat	ening:		Н	ospitalizatio	n:	Other Med	dically Important Co	onditions: Yes
Patient Information	on .											
Age	Gender		Height	V	Veight		Report	Outcome				
60 Years	Female					F	Recover	ed/resolved				
Link / Duplicate R	eport Infor	mation										
	Record	І Туре				Link AER*	* Numb	er				
Duplicate						00090	9260					
Product Informati	on											
Product Descr	ription	Health Pro	oduct Role	Dosag	e Form	Route Administr		Dose	Fre	equency	Therapy Duration	Indication(s)
PANZYGA		Sus	spect		JTION /ENOUS	Intravenous otherwis specifie	se [`]	80.0 ml		Once		Pelvic inflammatory disease
Adverse Reaction Information	Term											
Adverse Reaction Term(s)			tion Term(s)		Me	dDRA	Version			Reaction Duration		

v.27.1

v.27.1 v.27.1

Anaphylactic reaction

Blood potassium decreased

Blood pressure increased

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000725404	1	2019-04-26	2019-05-21	MAH	NGAM-053-19-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability: Yes	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Information

Age	Gender	Height	Weight	Report Outcome
50 Years	Male		81 Kilogram	Unknown

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

1 Todaot Illioi Illation							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
BUPROPION	Concomitant	TABLET (EXTENDED- RELEASE)					
CEFTRIAXONE FOR INJECTION USP	Concomitant	POWDER FOR SOLUTION INTRAMUSCULAR					
CLINDAMYCIN	Concomitant	NOT SPECIFIED					
FLUOXETINE	Concomitant	NOT SPECIFIED					
OXYCODONE	Concomitant	NOT SPECIFIED					
PANTOPRAZOLE	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	20.0 Gram	Total		General physical health deterioration
PREGABALIN	Concomitant	Capsules					

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Chills	v.27.1	
Chromaturia	v.27.1	
Dyspnoea	v.27.1	
Haptoglobin increased	v.27.1	
Hypertension	v.27.1	
Нурохіа	v.27.1	
Nausea	v.27.1	
Pleuritic pain	v.27.1	
Tachycardia	v.27.1	
Tachypnoea	v.27.1	
Transfusion-related acute lung injury	v.27.1	
Vomiting	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000725886	1	2019-05-10	2019-05-29	MAH	NGAM-056-19-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient	Information	١

Age	Gender	Height	Weight	Report Outcome
54 Years	Female		91 Kilogram	Recovering/resolving

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
ACETAMINOPHEN	Concomitant	NOT SPECIFIED					
ACYCLOVIR	Concomitant	NOT SPECIFIED					
CEFTRIAXONE SODIUM FOR INJECTION BP	Concomitant						
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	30.0 Gram	Total		Hypogammaglo bulinaemia
SODIUM CHLORIDE	Concomitant	NOT SPECIFIED					
VALACYCLOVIR	Concomitant	NOT SPECIFIED					

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Acute pulmonary oedema	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Body temperature increased	v.27.1	
Chills	v.27.1	
Dyspnoea	v.27.1	
Нурохіа	v.27.1	
Tachycardia	v.27.1	
Tachypnoea	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information **AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date	Source of Report	Market Authorization Holder AER Number	Type of Report	Reporter Type
000726191	0	2019-05-17	2019-05-17	Community		Spontaneous	Pharmacist

Serious report?	Death:	Disability: Yes	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions:

Patient Information

	Age	Gender	Height	Weight	Report Outcome
	38 Years	Male	168 Centimeter	69 Kilogram	Unknown

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

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Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
ALFACALCIDOL	Concomitant						
BETAMETHASONE	Concomitant	NOT SPECIFIED					
BISOPROLOL	Concomitant	Tablets					
CHOLECALCIFEROL	Concomitant	NOT SPECIFIED					
CLINDAMYCIN	Concomitant	NOT SPECIFIED					
CLONAZEPAM	Concomitant	Tablets					
CODEINE	Concomitant	Tablets					
DARBEPOETIN ALFA	Concomitant						
DICLOFENAC	Concomitant						
FOLIC ACID	Concomitant	NOT SPECIFIED					
FOSRENOL	Concomitant	TABLET (CHEWABLE)					
HYDROCORTISONE ACETATE CREAM USP	Concomitant	Cream					

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
MAXITROL	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	27.0 Gram	8 every 1 Years		Infection prophylaxis
RANITIDINE	Concomitant	NOT SPECIFIED					
REPLAVITE	Concomitant	Tablets					
SENSIPAR	Concomitant	NOT SPECIFIED					
carbo	Concomitant	Capsules					

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Alopecia	v.27.1	
Decreased activity	v.27.1	
Product substitution issue	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000726193	2	2019-05-21	2019-11-05	MAH	NGAM-077-19-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening: Yes	Hospitalization: Yes	Other Medically Important Conditions:

Patient	Informa	tion

Age	Gender	Height	Weight	Report Outcome
67 Years	Female		67 Kilogram	Recovered/resolved

Link / Duplicate Report Information

Record Type Link AER** Number	,
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No duplicate or linked report.

Product Information

1 Todact Illioilliation							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	30.0 Gram	Total		Immune thrombocytopen ia
PREGABALIN	Concomitant	Capsules					

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Blood pressure increased	v.27.1	
Chills	v.27.1	
Dyspnoea	v.27.1	
Ecchymosis	v.27.1	
Febrile nonhaemolytic transfusion reaction	v.27.1	
Haemolysis	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Oedema peripheral	v.27.1	
Pain in extremity	v.27.1	
Pyrexia	v.27.1	
Shock	v.27.1	
Tachycardia	v.27.1	
Tremor	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000726284	0	2019-05-22	2019-05-22	MAH	NGAM-067-19-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Informa	tion			
Age Gender		Height	Weight	Report Outcome
59 Years	Male			Unknown

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report	

Product Information

Froduct information							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	30.0 Gram	Total		Leukaemia

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Dyspnoea	v.27.1	
Hypersensitivity	v.27.1	
Oxygen saturation decreased	v.27.1	
Tremor	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000726513	0	2019-05-24	2019-05-24	MAH	NGAM-079-19-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

	Patient	Informa	tion
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Age	Gender	Height	Weight	Report Outcome
82 Years	Male			Unknown

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

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Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	30.0 Gram	Total		Guillain-Barre syndrome

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Blood pressure increased	v.27.1	
Chills	v.27.1	
Oxygen saturation decreased	v.27.1	
Tachypnoea	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information **AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000727739	0	2019-07-02	2019-07-02	MAH	NGAM-099-19-CA	Spontaneous	Physician

Serious report?Death:Disability:Congenital Anomaly:SeriousLife Threatening:Hospitalization:YesOther Medically Important Conditions:

Patient Information

Age	Gender	Height	Weight	Report Outcome
54 Years	Male	177 Centimeter	99 Kilogram	Recovered/resolved

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

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Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	40.0 Gram	1 every 4 Weeks		Hypogammaglo bulinaemia

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Back pain	v.27.1	
Chills	v.27.1	
Pallor	v.27.1	
Peripheral coldness	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000727794	0	2019-07-03	2019-07-03	MAH	NGAM-105-19-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient	Informa	ation

Age	Gender	Height	Weight	Report Outcome
18 Years	Male		77 Kilogram	Recovered/resolved

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	150.0 Gram			Encephalitis autoimmune

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Anaemia	v.27.1	
Haemolysis	v.27.1	
Hepatic function abnormal	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information	n

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000727857	1	2019-07-05	2019-08-18	MAH	NGAM-111-19-CA	Spontaneous	Physician

Serious report?	Death:	Disability:	Congenital Anomaly:	
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Ye	es

Patient Information

Age	Gender	Height	Weight	Report Outcome
31 Years	Female		98 Kilogram	Unknown

Link / Duplicate Report Information

Link AER Number Record Type**

No duplicate or linked report.

Product Information

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Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
ACETAMINOPHEN	Concomitant	NOT SPECIFIED					
METOCLOPRAMIDE	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	60.0 Gram	1 every 1 Days	3.0 Days	Myasthenia gravis

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Arthralgia	v.27.1	
Dizziness	v.27.1	
Erythema	v.27.1	
Headache	v.27.1	
Hypotension	v.27.1	
Myalgia	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Received Date Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000727963	2	2019-07-02	2020-03-10	MAH	NGAM-100-19-CA	Spontaneous	Other health professional

Serious report?	Death:	No	Disability:	No	Congenital Anomaly:	No
Serious	Life Threatening: Y	Yes	Hospitalization:	Yes	Other Medically Important Conditions:	Yes

Patient Information

- and - marie						
Age Ge		Gender	Height	Weight	Report Outcome	
	28 Years	Male	165 Centimeter	54 Kilogram	Recovered/resolved	

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
ACETAMINOPHEN	Concomitant	NOT SPECIFIED					
CETIRIZINE	Concomitant	Tablets					
COVERSYL	Suspect	Tablets	Oral	2.0 Milligram	1 every 1 Days		Product used for unknown indication
DIPHENHYDRAMINE	Concomitant	NOT SPECIFIED					
HYDROCORTISONE	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	25.0 Gram	1 every 1 Months		Hypogammaglo bulinaemia

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Neurogenic shock	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Seizure	v.27.1	
Syncope	v.27.1	
Transfusion reaction	v.27.1	
Tremor	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000728494	0	2019-07-23	2019-07-23	MAH	NGAM-115-19-CA	Spontaneous	Physician

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization: Yes	Other Medically Important Conditions:

Patient Information

Age	Gender	Height	Weight	Report Outcome
64 Years	Female	150 Centimeter	68 Kilogram	Recovered/resolved

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

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Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
DEXAMETHASONE	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	35.0 Gram	1 every 1 Days	4.0 Days	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Ataxia	v.27.1	
Dizziness	v.27.1	
Hypoacusis	v.27.1	
Meningitis aseptic	v.27.1	
Salivary hypersecretion	v.27.1	
Vision blurred	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

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1245			 on
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**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000728502	0	2019-07-23	2019-07-23	MAH	NGAM-116-19-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization: Yes	Other Medically Important Conditions:

Patient	Informa	tion

Age	Gender	Height	Weight	Report Outcome
75 Years	Female	152 Centimeter	73 Kilogram	Recovered/resolved

Link / Duplicate Report Information

Record Type	Link AER** Number

No duplicate or linked report.

Product Information

1 Todact information							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
ACETYLSALICYLIC ACID	Concomitant	NOT SPECIFIED					
CLAVULIN	Concomitant	NOT SPECIFIED					
DIPHENHYDRAMINE	Concomitant	NOT SPECIFIED					
FOLIC ACID	Concomitant	NOT SPECIFIED					
HYDROCORTISONE	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	30.0 Gram	1 every 1 Months		Hypogammaglo bulinaemia

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Chills	v.27.1	
Febrile nonhaemolytic transfusion reaction	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Feeling cold	v.27.1	
Hypertension	v.27.1	
Hypoaesthesia oral	v.27.1	
Malaise	v.27.1	
Pain in jaw	v.27.1	
Pyrexia	v.27.1	
Tachypnoea	v.27.1	
Tremor	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000729039	0	2019-08-07	2019-08-07	MAH	NGAM-125-19-CA	Spontaneous	Physician

Serious report?	Death:	Disability:	Congenital Anomaly:	
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions:	Yes

Patient Information

Age Gender Height		Weight	Report Outcome	
38 Years	Male	174 Centimeter	63 Kilogram	Recovered/resolved

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	30.0 Gram	1 every 1 Days		Immune thrombocytopen ia
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	30.0 Gram	1 every 1 Days		Immune thrombocytopen ia

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Dizziness	v.27.1	
Headache	v.27.1	
Vomiting	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000729076	1	2019-08-09	2019-10-28	MAH	NGAM-139-19-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Informa	ition			
Age	Gender	Height	Weight	Report Outcome
62 Years	Male			Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report.	

Product Information

1 Todast Information							
Product Descrip	tion Health Product	Role Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	20.0 Milligram			Chronic inflammatory demyelinating polyradiculoneu ropathy

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Erythema	v.27.1	
Pruritus	v.27.1	
Skin exfoliation	v.27.1	
Urticaria	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information					
Adverse Reaction Report Number	Late				
000720077					

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000729077	0	2019-08-09	2019-08-09	MAH	NGAM-127-19-CA	Spontaneous	Physician

Seriou	us report?	Death:	Disability:	Congenital Anomaly:	
S	Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions:	Yes

Patient Information

Age	Gender	Height	Weight	Report Outcome	
15 Years	Female		119 Kilogram	Recovered/resolved	

Link / Duplicate Report Information

Link AER Number Record Type**

No duplicate or linked report.

Product Information

Product information							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
ACETAMINOPHEN	Concomitant	NOT SPECIFIED					
CLINDAMYCIN	Concomitant	NOT SPECIFIED					
MEROPENEM	Concomitant	POWDER FOR SOLUTION INTRAVENOUS					
OXYGEN	Concomitant	Gas for inhalation					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	50.0 Gram			Septic shock
VANCOMYCIN	Concomitant	NOT SPECIFIED					

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Acute pulmonary oedema	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Dyspnoea	v.27.1	
Hypotension	v.27.1	
Tachycardia	v.27.1	
Tachypnoea	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date	Source of Report	Market Authorization Holder AER Number	Type of Report	Reporter Type
000729082	2	2019-08-09	2020-02-12	MAH	NGAM-124-19-CA	Spontaneous	Physician

Serious report?	Death:	Yes	Disability:	No	Congenital Anomaly:	No
Serious	Life Threatening:	No	Hospitalization:	No	Other Medically Important Conditions:	Yes

Patient Information

Age	Gender	Height	Weight	Report Outcome
13 Years	Male		32 Kilogram	Fatal

Link / Duplicate Report Information	
Record Type	Link AER** Number
Duplicate	

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
CEFAZOLIN	Concomitant						
CEFTRIAXONE FOR INJECTION USP	Concomitant	POWDER FOR SOLUTION INTRAMUSCULAR					
OXYGEN	Concomitant	Gas for inhalation					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	30.0 Gram			Myocarditis

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Acute respiratory distress syndrome	v.27.1	
Chills	v.27.1	
Dyspnoea	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Encephalopathy	v.27.1	
Hypertension	v.27.1	
Livedo reticularis	v.27.1	
Metabolic disorder	v.27.1	
Multiple organ dysfunction syndrome	v.27.1	
Pyrexia	v.27.1	
Tachycardia	v.27.1	
Tachypnoea	v.27.1	
Troponin increased	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date	Source of Report	Market Authorization Holder AER Number	Type of Report	Reporter Type
000729208	0	2019-08-13	2019-08-13	MAH	NGAM-141-19-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient	Informa	tion

Age	Gender	Height	Weight	Report Outcome
48 Years	Male		70 Kilogram	Recovered/resolved

Link / Duplicate Report Information

Record Type	Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	70.0 Gram			Immune thrombocytopen ia

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Dizziness	v.27.1	
Headache	v.27.1	
Meningitis aseptic	v.27.1	
Neck pain	v.27.1	
Pyrexia	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000729744	0	2019-08-30	2019-08-30	MAH	NGAM-156-19-CA	Spontaneous	Physician

Serious report?	Death:	Disability:	Congenital Anomaly:	
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions:	Yes

Patient Information

Age	Gender	Height	Weight	Report Outcome
68 Years	Male		95 Kilogram	Recovered/resolved

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

i ioaaot iiiioiiiiatioii							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
CANDESARTAN	Concomitant						
LEVOTHYROXINE	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous drip	190.0 Gram	Total		Immune thrombocytopen ia
PREDNISONE	Suspect	NOT SPECIFIED					

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Anaemia	v.27.1	
Coombs direct test positive	v.27.1	
Haemolysis	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date	Source of Report	Market Authorization Holder AER Number	Type of Report	Reporter Type
000729770	0	2019-09-02	2019-09-02	MAH	NGAM-163-19-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization: Yes	Other Medically Important Conditions:

Patient Information			
Age	Gender	Height	Weight

Age	Gender	Height	Weight	Report Outcome
75 Years	Male			Unknown

Link / Duplicate Report Information	
Record Type	Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	30.0 Gram			Leukaemia

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Hypotension	v.27.1	
Tachycardia	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information	on	**AE	R = Adverse R	Reaction Re	eport							
Adverse Reaction Report Number		ER Version nber	Initial Recei	ved Date	Latest Rece	eived Date	Source	e of Report	Auth	arket orization AER Number	Type of Report	Reporter Typ
000730414		0	2019-09	9-19	2019-0	9-19		МАН	NGAM-	168-19-CA	Spontaneous	Other health professional
Serious re	eport?				Death:			Disabilit	y:		Congenital	Anomaly:
Serio	us		L	ife Threat	ening:		H	ospitalizatio	n:	Other Med	dically Important C	onditions: Ye
Patient Information	on											
Age	Gender		Height	V	Veight		Report	Outcome				
8 Years	Male					F	Recover	ed/resolved				
Link / Duplicate R	eport Info	rmation										
	Recor	d Type				Link AER*	* Numb	er				
No duplicate or link	ked report.											
Product Informati	ion											
Product Desci	ription	Health Pro	oduct Role	Dosag	je Form	Route Administr		Dose	Fr	equency	Therapy Duration	Indication(s
PANZYGA		Sus	spect		JTION /ENOUS	Intravenou otherwi specifie	se`	20.0 Gran	n			Chronic inflammatory demyelinatin polyradiculone ropathy
Adverse Reaction Information	n Term											
	Adv	verse Reac	tion Term(s)			Me	edDRA	Version			Reaction Duration	

v.27.1

Hypersensitivity

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

Reaction Duration

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Informatio	n	**AE	R = Adverse F	Reaction Re	port							
Adverse Reaction Report Number		ER Version nber	Initial Rece	ved Date	Latest Rece	eived Date	Sourc	e of Report	Autho	arket orization LER Number	Type of Report	Reporter Type
000730420		0	2019-09-19 2019-09-			9-19		MAH	NGAM-166-19-CA		Spontaneous	Other health professional
Serious report?				Death: Disability:				Congenital	Anomaly:			
Seriou	ıs		ı	ife Threat	ening:		Н	ospitalizatio	n:	Other Med	dically Important Co	onditions: Yes
Patient Information	n											
Age	Gender		Height	V	Veight		Report	Outcome				
66 Years	Male		•		Recovered/resolved							
Link / Duplicate R	eport Info	rmation										
		d Type				Link AER*	* Numb	er				
No duplicate or link	ed report.											
Product Informati	on											
Product Descr	ription	Health Pr	oduct Role	Dosag	e Form	Route Administr	-	Dose	Fre	equency	Therapy Duration	Indication(s)
DEXTROSE		Conc	omitant	NOT SP	ECIFIED							
PANZYGA		Sus	spect	INTRAVENOUS other		Intravenou otherwi specifie	ise`	30.0 Gran	n			Chronic lymphocytic leukaemia
Adverse Reaction Information	Term											

MedDRA Version

v.27.1

Adverse Reaction Term(s)

Hypersensitivity

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000730584	0	2019-09-24	2019-09-24	MAH	NGAM-174-19-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Informa	tion			
Age Gender		Height	Weight	Report Outcome
88 Years	Female			Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report.	

Product Information

1 Todaot IIII o I III ation	·						
Product Descript	tion Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous drip	30.0 Gram	Total		Chronic inflammatory demyelinating polyradiculoneu ropathy

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Chest pain	v.27.1	
Chills	v.27.1	
Headache	v.27.1	
Hypertension	v.27.1	
Nausea	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000730593	0	2019-09-24	2019-09-24	MAH	NGAM-170-19-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient	Informa	tion

Age	Gender	Height	Weight	Report Outcome
65 Years	Male		71 Kilogram	Unknown

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	60.0 Gram	Total		Immune thrombocytopen ia

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Dyspnoea	v.27.1	
Hypertension	v.27.1	
Нурохіа	v.27.1	
Tachycardia	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000730596	2	2019-09-24	2019-10-28	MAH	NGAM-175-19-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Informa	tion			
Age	Gender	Height	Weight	Report Outcome
51 Years	Female			Unknown

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report.	

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	25.0 Gram	Total	115.0 Minutes	Chronic inflammatory demyelinating polyradiculoneu ropathy

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Chills	v.27.1	
Febrile nonhaemolytic transfusion reaction	v.27.1	
Headache	v.27.1	
Product contamination microbial	v.27.1	
Pyrexia	v.27.1	
Suspected transmission of an infectious agent via product	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000730795	0	2019-09-30	2019-09-30	MAH	NGAM-183-19-CA	Spontaneous	Physician

Serious report?Death:Disability:Congenital Anomaly:SeriousLife Threatening:Hospitalization:Other Medically Important Conditions:Yes

Patient Information

Age	Age Gender Height		Weight	Report Outcome	
78 Years	Male	173 Centimeter	66 Kilogram	Unknown	

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Froduct information							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
ACETAMINOPHEN	Concomitant	NOT SPECIFIED					
FUROSEMIDE	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	20.0 Gram	Total		Autoimmune haemolytic anaemia
RED BLOOD CELLS	Suspect	NOT SPECIFIED	Unknown	1.0 Dosage forms			Product used for unknown indication

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Chills	v.27.1	
Dyspnoea	v.27.1	
Febrile nonhaemolytic transfusion reaction	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Hypertension	v.27.1	
Oxygen saturation decreased	v.27.1	
Pyrexia	v.27.1	
Tachycardia	v.27.1	
Tachypnoea	v.27.1	
Tremor	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000730989	0	2019-10-07	2019-10-07	MAH	NGAM-184-19-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Informa	tion			
Age	Gender	Height	Weight	Report Outcome
78 Years	Female	158 Centimeter	44 Kilogram	Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report.	

Product Information

Froduct information							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	25.0 Gram		105.0 Minutes	Haemolytic anaemia

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Headache	v.27.1	
Hyperhidrosis	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000731022	3	2019-10-07	2021-04-12	MAH	NGAM-181-19-CA	Spontaneous	Physician

Serious report?	Death: Ye	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Information

Age Gender Height		Weight	Report Outcome	
76 Years	Female	158 Centimeter	65 Kilogram	Fatal

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

1 Todast Information							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
DIPHENHYDRAMINE	Concomitant	NOT SPECIFIED					
FOLIC ACID	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	60.0 Gram	Total		Bicytopenia
PLATELETS	Suspect	NOT SPECIFIED	Intravenous (not otherwise specified)	210.0 ml			Bicytopenia
PREDNISONE	Concomitant	NOT SPECIFIED					
PRIVIGEN	Suspect	SOLUTION INTRAVENOUS	Unknown				Bicytopenia

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Chills	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Feeling cold	v.27.1	
Haemolysis	v.27.1	
Pyrexia	v.27.1	
Tachycardia	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Re	Adverse eaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
	000731026	1	2019-10-08	2019-10-29	MAH	NGAM-158-19-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient	Information	
		_

Age Gender Height		Weight	Report Outcome	
83 Years	Male	96 Centimeter	96 Kilogram	Recovered/resolved

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
ACETYLSALICYLIC ACID	Concomitant	NOT SPECIFIED					
CEFADROXIL	Concomitant	NOT SPECIFIED					
FAMOTIDINE	Concomitant	NOT SPECIFIED					
FENTANYL	Concomitant	NOT SPECIFIED					
FOLIC ACID	Concomitant	NOT SPECIFIED					
INSULIN	Concomitant	NOT SPECIFIED					
MAGNESIUM	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	40.0 Gram			Guillain-Barre syndrome
PROPOFOL	Concomitant						
RANITIDINE	Concomitant	NOT SPECIFIED					
ROCURONIUM	Concomitant	NOT SPECIFIED			_		

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Bradycardia	v.27.1	
Hyperhidrosis	v.27.1	
Hypotension	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000732542	1	2019-11-05	2020-06-11	MAH	NGAM-197-19-CA	Spontaneous	Other health professional

Serious report?	Death:		Disability:		Congenital Anomaly:
Serious	Life Threatening:	Yes	Hospitalization:	Yes	Other Medically Important Conditions: Yes

Patient Informa	tion			
Age	Gender	Height	Weight	Report Outcome
69 Years	Male		81 Kilogram	Unknown

Link / Duplicate Report Information	
Record Type	Link AER** Number
Duplicate	000731493

Product Information							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
AMOXICILLIN	Concomitant	NOT SPECIFIED					
AS-3 RBC LR (E7962V00)	Suspect		Intravenous (not otherwise specified)			145.0 Minutes	Neutropenia
MEROPENEM	Concomitant	POWDER FOR SOLUTION INTRAVENOUS					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	30.0 Gram		2.0 Days	Non-Hodgkin's lymphoma

Adverse Reaction Term Information Suspect INTRAVENOUS otherwise specified) 30.0 Gram 2.0 Days lymphoma

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Abdominal pain upper	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Chest pain	v.27.1	
Dyspnoea	v.27.1	
Heart rate increased	v.27.1	
Pleuritic pain	v.27.1	
Pulmonary embolism	v.27.1	
Respiratory rate increased	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000732670	0	2019-11-06	2019-11-06	MAH	NGAM-200-19-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Informa	tion			
Age	Gender	Height	ght Weight Report Outcome	
75 Years	Male		83 Kilogram	Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report.	

Product Information

Froduct information							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
NITROGLYCERIN	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	80.0 Gram	Total	95.0 Minutes	Immune thrombocytopen ia

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Angina pectoris	v.27.1	
Chills	v.27.1	
Dyspnoea	v.27.1	
Febrile nonhaemolytic transfusion reaction	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000732922	0	2019-11-11	2019-11-11	MAH	NGAM-073-19-CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Informa	ition			
Age	Gender	Height	Weight	Report Outcome
61 Vears	Male			Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report.	

Product Information

Froduct information							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	35.0 Gram			Myasthenia gravis

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Acute pulmonary oedema	v.27.1	
Cough	v.27.1	
Dyspnoea	v.27.1	
Hyperhidrosis	v.27.1	
Hypertension	v.27.1	
Muscle spasms	v.27.1	
Stridor	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Tachycardia	v.27.1	
Tachypnoea	v.27.1	
Wheezing	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000732960	0	2019-11-14	2019-11-14	MAH	NGAM20619CA	Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient	Informa	tion

Age	Gender	Height	Weight	Report Outcome
65 Years	Female			Recovered/resolved

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
DARATUMUMAB	Concomitant	SOLUTION INTRAVENOUS					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	35.0 Gram	Total		Plasma cell myeloma

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Chills	v.27.1	
Dyspnoea	v.27.1	
Hypersensitivity	v.27.1	
Tremor	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000734289	0	2019-12-09	2019-12-09	MAH		Spontaneous	Other health professional

Serious report?	Death:	No	Disability:	No	Congenital Anomaly:	No
Serious	Life Threatening:	No	Hospitalization:	No	Other Medically Important Conditions:	Yes

Patient	Informa	tion

Age	Gender	Height	Weight	Report Outcome
42 Years	Female	161 Centimeter	95 Kilogram	Recovered/resolved

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	160.0 Gram		2.0 Days	Guillain-Barre syndrome

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Anaemia	v.27.1	
Haemolysis	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000734798	0	2019-12-17	2019-12-17	Hospital		Spontaneous	Other health professional

Serious report?	Death:	No	Disability:	No	Congenital Anomaly:	No
Not Serious	Life Threatening:	No	Hospitalization:	No	Other Medically Important Conditions:	No

Patient Information

Age	Gender	Height	Weight	Report Outcome
39 Years	Male		90 Kilogram	Recovered/resolved

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

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Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	240.0 ml		35.0 Minutes	Immune thrombocytopen ia
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	1.0 Gram		160.0 Minutes	Immune thrombocytopen ia

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Chills	v.27.1	
Feeling cold	v.27.1	
Headache	v.27.1	
Myalgia	v.27.1	
Pain	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Urticaria	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000735133	1	2019-12-18	2021-10-18	MAH		Spontaneous	Other health professional

Serious report?	Death:	No	Disability:	No	Congenital Anomaly:	No
Serious	Life Threatening:	No	Hospitalization:	Yes	Other Medically Important Conditions:	No

Patient	Informa	tion

Age	Gender	Height	Weight	Report Outcome
71 Years	Female			Recovered/resolved

Link / Duplicate Report Information

Record Type Link AER** Number	
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No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	30.0 Gram	1 every 6 Weeks		Chronic inflammatory demyelinating polyradiculoneu ropathy

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Chest pain	v.27.1	
Hypertension	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000735439	0	2019-12-27	2019-12-27	Hospital		Spontaneous	Other health professional

Serious report?	Death:	No	Disability:	No	Congenital Anomaly: No
Serious	Life Threatening:	No	Hospitalization:	No	Other Medically Important Conditions: Yes

Patient Informa	tion			
Age	Gender	Height	Weight	Report Outcome
72 Years	Male		82 Kilogram	Unknown

Link / Duplicate Report Information	
Record Type	Link AER** Number
Duplicate	000919087

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	50.0 ml		45.0 Minutes	Guillain-Barre syndrome
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	300.0 ml		3.0 Hours	Guillain-Barre syndrome

Adverse Reaction Term Information			
Adverse Reaction Term(s)		MedDRA Version	Reaction Duration
Febrile nonhaemolytic transfusion reaction		v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000735562	0	2019-12-30	2019-12-30	Hospital		Spontaneous	Other health professional

Serious report?	Death:	No	Disability:	No	Congenital Anomaly:	No
Not Serious	Life Threatening:	No	Hospitalization:	No	Other Medically Important Conditions:	No

Patient Informa	tion			
Age	Gender	Height	Weight	Report Outcome
72 Years	Female		70 Kilogram	Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report.	

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	100.0 ml			Hypogammaglo bulinaemia
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	200.0 ml			Hypogammaglo bulinaemia
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	100.0 ml			Hypogammaglo bulinaemia
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	200.0 ml			Hypogammaglo bulinaemia

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Allergic transfusion reaction	v.27.1	64 Days
Pruritus	v.27.1	64 Days
Urticaria	v.27.1	64 Days

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Summary of Reported Adverse Rea

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000737353	0	2020-01-16	2020-01-16	Hospital		Spontaneous	Other health professional

Serious report?	Death:	No	Disability:	No	Congenital Anomaly:	No
Serious	Life Threatening:	No	Hospitalization:	No	Other Medically Important Conditions:	Yes

Patient	Informa	ation

Age	Gender	Height	Weight	Report Outcome
55 Years	Female	160 Centimeter	91 Kilogram	Recovered/resolved

Link / Duplicate Report Information

Record Type	Link AER** Number
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No duplicate or linked report.

Product Information

1 TOGGOT IIIIOTIIIGGOTI							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
CYTOSAR	Concomitant	POWDER FOR SOLUTION INTRATHECAL					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)			50.0 Minutes	
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)			38.0 Minutes	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Arthralgia	v.27.1	
Blood pressure decreased	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Dizziness	v.27.1	
Haemolysis	v.27.1	
Headache	v.27.1	
Nausea	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000908444	0	2020-01-31	2020-01-31	Community		Spontaneous	Other health professional

Serious report?	Death:	No	Disability:	No	Congenital Anomaly:	No
Not Serious	Life Threatening:	No	Hospitalization:	No	Other Medically Important Conditions:	No

Patient I	nforma	tion		
			_	-

Age	Gender	Height	Weight	Report Outcome
67 Years	Male			Unknown

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS		250.0 ml		112.0 Minutes	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Chills	v.27.1	
Hypertension	v.27.1	
Pyrexia	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000909637	0	2020-02-14	2020-02-14	Hospital		Spontaneous	Other health professional

Serious report?	Death:	No	Disability:	No	Congenital Anomaly:	No
Serious	Life Threatening:	No	Hospitalization:	No	Other Medically Important Conditions:	Yes

Patient Information				
Age	Gender	Height	Weight	Report Outcome
78 Years	Male	159 Centimeter	70 Kilogram	Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS		100.0 ml		54.0 Minutes	Platelet count decreased
PANZYGA	Suspect	SOLUTION INTRAVENOUS		600.0 ml		4.0 Hours	Platelet count decreased

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Chills	v.27.1	
Febrile nonhaemolytic transfusion reaction	v.27.1	
Hypertension	v.27.1	
Nausea	v.27.1	
Vomiting	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000910579	0	2020-02-21	2020-02-21	Hospital		Spontaneous	Other health professional

Serious report?	Death:	No	Disability:	No	Congenital Anomaly: No
Serious	Life Threatening:	Yes	Hospitalization:	Yes	Other Medically Important Conditions: Yes

Patient	Informati	tion

Age	Gender	Height	Weight	Report Outcome
15 Years	Male	175 Centimeter	87 Kilogram	Unknown

Link / Duplicate Report Information

Record Type	Link AER** Number
Linked	000989072

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
BIPHENTIN	Concomitant	CAPSULE, EXTENDED RELEASE					
DEXAMETHASONE	Concomitant	LIQUID INTRA- ARTICULAR					
PANZYGA	Suspect		Intravenous (not otherwise specified)	75.0 Gram			Thrombocytope nic purpura
PREDNISONE	Concomitant	NOT SPECIFIED					
RITUXAN	Concomitant						

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Cerebral venous sinus thrombosis	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000910798	1	2020-02-24	2020-03-19	MAH		Spontaneous	Physician

Serious report?	Death:	No	Disability:	No	Congenital Anomaly:	No
Serious	Life Threatening:	No	Hospitalization:	No	Other Medically Important Conditions:	Yes

Patient Information

Age	Gender	Height	Weight	Report Outcome
35 Years	Female			Unknown

Link / Duplicate Report Information

Record Type Link AER Number**

No duplicate or linked report.

Product Information

1 Todast IIII STIII ation							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS					Neonatal alloimmune thrombocytopen ia
PANZYGA	Suspect	SOLUTION INTRAVENOUS					Neonatal alloimmune thrombocytopen ia
PANZYGA	Suspect	SOLUTION INTRAVENOUS					Neonatal alloimmune thrombocytopen ia
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)				Neonatal alloimmune thrombocytopen ia

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Maternal exposure during pregnancy	v.27.1	
Treponema test positive	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000911183	0	2020-03-03	2020-03-03	Hospital		Spontaneous	Other health professional

Serious report?	Death:	No	Disability:	No	Congenital Anomaly:	No
Serious	Life Threatening:	No	Hospitalization:	No	Other Medically Important Conditions:	Yes

Patient Informa	tion			
Age	Gender	Height	Weight	Report Outcome
6 Years	Female	110 Centimeter	19 Kilogram	Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
ACETAMINOPHEN	Concomitant	NOT SPECIFIED					
BENADRYL	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	180.0 ml	Once	5.0 Hours	Epilepsy, CSWS syndrome, Attention deficit hyperactivity disorder

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Chills	v.27.1	
Headache	v.27.1	
Nausea	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Pyrexia	v.27.1	
Vomiting	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000911288	0	2020-02-21	2020-02-21	Hospital		Spontaneous	Other health professional

Serious report?	Death:	No	Disability:	No	Congenital Anomaly:	No
Serious	Life Threatening:	No	Hospitalization:	Yes	Other Medically Important Conditions:	No

Patient Information

Age	Gender	Height	Weight	Report Outcome
9 Years	Male		35 Kilogram	Recovered/resolved

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

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Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)		
ACETAMINOPHEN	Concomitant	NOT SPECIFIED							
CALCIUM CARBONATE	Concomitant	Tablets							
CHOLECALCIFEROL	Concomitant	NOT SPECIFIED							
FLUTICASONE	Concomitant	SPRAY, METERED DOSE							
HYDROCORTISONE	Concomitant	POWDER FOR SOLUTION INTRAMUSCULAR							
MONTELUKAST	Concomitant	NOT SPECIFIED							
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	20.0 Gram	1 every 1 Months		Hypogammaglo bulinaemia		

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Headache	v.27.1	1 Days

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000911638	0	2020-03-10	2020-03-10	Hospital		Spontaneous	Other health professional

Serious report?	Death:	No	Disability:	No	Congenital Anomaly:	No
Serious	Life Threatening:	No	Hospitalization:	No	Other Medically Important Conditions:	Yes

Patient Informa	tion			
Age Gender		Height	Weight	Report Outcome
70 Years	Male	120 Centimeter		Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report.	

Product Information

1 Todaot IIII of III ation							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	144.0 ml	1 every 1 Days	116.0 Minutes	Immune thrombocytopen ia, Anaemia
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	200.0 ml	1 every 1 Days	117.0 Minutes	Immune thrombocytopen ia, Anaemia

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Blood pressure increased	v.27.1	
Body temperature increased	v.27.1	
Chills	v.27.1	
Febrile nonhaemolytic transfusion reaction	v.27.1	
Heart rate increased	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Nausea	v.27.1	
Oxygen saturation increased	v.27.1	
Respiratory rate increased	v.27.1	
Vomiting	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000913092	0	2020-03-04	2020-03-04	Hospital		Spontaneous	Other health professional

Serious report?	Death:	No	Disability:	No	Congenital Anomaly:	No
Serious	Life Threatening:	No	Hospitalization:	Yes	Other Medically Important Conditions:	No

Patient Information

Age	Gender	Height	Weight	Report Outcome
7 Years	Male		22 Kilogram	Recovered/resolved

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
GABAPENTIN	Concomitant	Capsules					
IBUPROFEN	Concomitant	NOT SPECIFIED					
METHYLPREDNISOLONE NOS	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	2.5 Gram	Once		Myelitis
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	2.5 Gram	Once		Myelitis
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	10.0 Gram	Once		Myelitis

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	10.0 Gram	Once		Myelitis
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	10.0 Gram	Once		Myelitis
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	10.0 Gram	Once		Myelitis

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Chills	v.27.1	
Meningitis aseptic	v.27.1	2 Days
Nausea	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000913238	1	2020-04-01	2020-09-29	MAH		Spontaneous	Other health professional

Serious report?	Death:	No	Disability:	No	Congenital Anomaly: No
Serious	Life Threatening:	No	Hospitalization:	Yes	Other Medically Important Conditions: No

Patient Information

Age	Gender	Height	Weight	Report Outcome
79 Years	Female			Unknown

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Froduct information							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
CETIRIZINE	Concomitant	Tablets					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	40.0 Gram	1 every 4 Weeks	26.0 Days	Secondary immunodeficien cy
PSEUDOEPHEDRINE	Concomitant	NOT SPECIFIED					
RED BLOOD CELLS	Concomitant	NOT SPECIFIED					

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Anaemia	v.27.1	
Anti A antibody positive	v.27.1	
Asthenia	v.27.1	
Chills	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Crepitations	v.27.1	
Cyanosis	v.27.1	
Dyspnoea	v.27.1	
Febrile nonhaemolytic transfusion reaction	v.27.1	
Feeling cold	v.27.1	
Haemolysis	v.27.1	
Hypertension	v.27.1	
Nausea	v.27.1	
Pallor	v.27.1	
Pulmonary oedema	v.27.1	
Tachycardia	v.27.1	
Tachypnoea	v.27.1	
Tremor	v.27.1	
Wheezing	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000913443	0	2020-04-02	2020-04-02	Hospital		Spontaneous	Other health professional

Serious report?	Death:	No	Disability:	No	Congenital Anomaly: No	
Serious	Life Threatening:	No	Hospitalization:	No	Other Medically Important Conditions: Yes	s

Patient Informa	tion			
Age	Gender	Height	Weight	Report Outcome
53 Years	Female	163 Centimeter	53 Kilogram	Recovered/resolved

Link / Duplicate Report Information			
Record Type	Link AER** Number		
Duplicate	000932116		

Product Information							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	200.0 ml		130.0 Minutes	Thrombocytope nia, Immune thrombocytopen ia, Hypersplenism
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	50.0 ml		37.0 Minutes	Thrombocytope nia, Immune thrombocytopen ia, Hypersplenism

Adverse Reaction Term Information Adverse Reaction Term(s) Adverse Reaction Term(s) Abdominal pain upper Back pain Negation Duration V.27.1 V.27.1

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Chills	v.27.1	
Febrile nonhaemolytic transfusion reaction	v.27.1	3 Days
Headache	v.27.1	
Nausea	v.27.1	
Pyrexia	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000915292	1	2020-04-23	2020-04-23	Hospital		Spontaneous	Other health professional

Serious report?	Death:	No	Disability:	No	Congenital Anomaly:	No
Serious	Life Threatening:	Yes	Hospitalization:	Yes	Other Medically Important Conditions:	No

Patient	Informa	ation

Age Gender Height		Weight	Report Outcome	
56 Years	Female	180 Centimeter	118 Kilogram	Recovered/resolved

Link / Duplicate Report Information

Record Type	Link AER** Number
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No duplicate or linked report.

Product Information

1 TOGGOT IIIIOTTIIGGOTI							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
ANTIHISTAMINE ORAL PWR	Concomitant	Powder					
CORTICOSTEROID(S)	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	200.0 ml		90.0 Minutes	Immunodeficien cy common variable
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	200.0 ml		60.0 Minutes	Immunodeficien cy common variable

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Blood pressure increased	v.27.1	
Dysphonia	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Dyspnoea	v.27.1	
Erythema	v.27.1	
Heart rate increased	v.27.1	
Hyperhidrosis	v.27.1	
Hypersensitivity	v.27.1	
Hypertension	v.27.1	
Pharyngeal oedema	v.27.1	
Stridor	v.27.1	
Throat tightness	v.27.1	
Use of accessory respiratory muscles	v.27.1	
Wheezing	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000922912	1	2020-07-29	2020-10-13	Hospital		Spontaneous	Other health professional

Serious report?	Death:	No	Disability:	No	Congenital Anomaly: No
Serious	Life Threatening:	No	Hospitalization:	No	Other Medically Important Conditions: Yes

Patient Informa	tion			
Age	Gender	Height	Weight	Report Outcome
70 Years	Female	161 Centimeter	41 Kilogram	Recovered/resolved

Link / Duplicate Report Information

Record Type
Link AER** Number

No duplicate or linked report.

Product Information

i roduct imormation							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
ACETAMINOPHEN	Concomitant	NOT SPECIFIED					
CETIRIZINE	Concomitant	Tablets					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)			43.0 Minutes	Infection prophylaxis
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)			33.0 Minutes	Infection prophylaxis

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Back pain	v.27.1	
Blood pressure increased	v.27.1	
Bradycardia	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Chills	v.27.1	
Fatigue	v.27.1	
Febrile nonhaemolytic transfusion reaction	v.27.1	
Nausea	v.27.1	
Oxygen saturation decreased	v.27.1	
Oxygen therapy	v.27.1	
Product intolerance	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000924997	2	2020-09-01	2021-10-08	MAH		Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization: Yes	Other Medically Important Conditions: Yes

Patient Informa	tion			
Age	Gender	Height	Weight	Report Outcome
72 Years	Female			Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report.	

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
OXYGEN	Concomitant	Gas for inhalation					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	30.0 Gram	Total		Hypogammaglo bulinaemia
PREDNISONE	Concomitant	NOT SPECIFIED					

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Chest pain	v.27.1	
Chills	v.27.1	
Dyspnoea	v.27.1	
Headache	v.27.1	
Hypertension	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Нурохіа	v.27.1	
Pain in jaw	v.27.1	
Tachypnoea	v.27.1	
Transfusion-related acute lung injury	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000926151	0	2020-09-17	2020-09-17	Hospital		Spontaneous	Other health professional

Serious report?	Death:	No	Disability:	No	Congenital Anomaly: No
Serious	Life Threatening:	No	Hospitalization:	No	Other Medically Important Conditions: Yes

Patient Informa	tion			
Age	Gender	Height	Weight	Report Outcome
49 Years	Female		66 Kilogram	Unknown

Link / Duplicate Report Information	
Record Type	Link AER** Number
Duplicate	000943694

Product Information							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	200.0 ml		3.0 Hours	
PANZYGA	Suspect	SOLUTION INTRAVENOUS		50.0 ml		74.0 Minutes	

Adverse Reaction Term Information			
Adverse	e Reaction Term(s)	MedDRA Version	Reaction Duration
Headache		v.27.1	
Hypertension	l de la companya de	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000927019	0	2020-09-28	2020-09-28	MAH		Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization: Yes	Other Medically Important Conditions:

Patient Informa	tion

Age	Gender	Height	Weight	Report Outcome
79 Years				Recovered/resolved

Link / Duplicate Report Information

Record Type	Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
ACETAMINOPHEN	Concomitant	NOT SPECIFIED					
CETIRIZINE	Concomitant	Tablets					
INSULIN	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	40.0 Gram	1 every 4 Weeks		Secondary immunodeficien cy

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Acute pulmonary oedema	v.27.1	
Bronchospasm	v.27.1	
Chills	v.27.1	
Cyanosis	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Dyspnoea	v.27.1	
Febrile nonhaemolytic transfusion reaction	v.27.1	
Hypertension	v.27.1	
Hypoaesthesia	v.27.1	
Tachycardia	v.27.1	
Tachypnoea	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000928557	0	2020-10-16	2020-10-16	MAH		Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Informa	tion			
Age	Gender	Height	Weight	Report Outcome
7 Years	Male			Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report	

Product Information

Froduct information							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	15.0 Gram		1.0 Days	Myelitis transverse

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Chills	v.27.1	
Discomfort	v.27.1	
Meningitis aseptic	v.27.1	
Musculoskeletal stiffness	v.27.1	
Nausea	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

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Report Information	

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000931283	0	2020-11-17	2020-11-17	MAH		Spontaneous	Nurse

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization: Yes	Other Medically Important Conditions:

Patient Information

Age Gender		Height Weight		Report Outcome	
14 Years	Male	140 Centimeter	30 Kilogram	Recovered/resolved	

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	20.0 Gram			Immunodeficien cy common variable

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Chills	v.27.1	
Drug intolerance	v.27.1	
General physical health deterioration	v.27.1	
Headache	v.27.1	
Meningitis aseptic	v.27.1	
Nausea	v.27.1	
Pyrexia	v.27.1	
Vomiting	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000932116	1	2020-11-24	2021-03-05	MAH		Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Information					
	Age Gender 53 Years Female		Height	Weight	Report Outcome
			163 Centimeter	53 Kilogram	Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
Duplicate	000913443
Duplicate	000914025

Product Information							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION	Intravenous (not otherwise	55.0 Gram			Immune thrombocytopen

			Administration			
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	55.0 Gram		Immune thrombocytopen ia
Adverse Reaction Term Information						

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Abdominal pain upper	v.27.1	
Back pain	v.27.1	
Chills	v.27.1	
Febrile nonhaemolytic transfusion reaction	v.27.1	
Feeling cold	v.27.1	
Headache	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Nausea	v.27.1	
Somnolence	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000936791	0	2021-01-22	2021-01-22	MAH		Spontaneous	Physician

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Information

Age	Gender	Height	Weight	Report Outcome
71 Years	Female			Recovered/resolved

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
CLINDAMYCIN	Concomitant	NOT SPECIFIED	Unknown				
DEXAMETHASONE	Concomitant	NOT SPECIFIED	Unknown				
MEROPENEM	Concomitant	POWDER FOR SOLUTION INTRAVENOUS					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	80.0 Gram			Autoimmune haemolytic anaemia
RED BLOOD CELLS	Suspect	NOT SPECIFIED	Unknown				Product used for unknown indication

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Respiratory failure	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Tachypnoea	v.27.1	
Wheezing	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31

173 Report(s)

Report Information	
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**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000942344	1	2021-03-10	2021-03-22	MAH		Spontaneous	Other health professional

Serious report?	Death:	Disability:		Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Yes	Other Medically Important Conditions:

Patient Information

Age	Gender	Height	Weight	Report Outcome
61 Years	Female		90 Kilogram	Recovered/resolved

Link / Duplicate Report Information

Record Type Link AER Number**

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
FRAGMIN	Concomitant	SOLUTION INTRAVENOUS					
IRBESARTAN	Concomitant	Tablets	Unknown				
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	90.0 Gram	Total		Paraneoplastic syndrome
POLYETHYLENE GLYCOL	Concomitant	NOT SPECIFIED					
TAMSULOSIN	Concomitant	NOT SPECIFIED					
THIAMINE	Concomitant	Tablets					

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Anaemia	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Angioedema	v.27.1	
Chest pain	v.27.1	
Drug intolerance	v.27.1	
Dyspnoea	v.27.1	
Nausea	v.27.1	
Platelet count decreased	v.27.1	
Swelling	v.27.1	
Tachycardia	v.27.1	
Vomiting	v.27.1	
Wheezing	v.27.1	
White blood cell count decreased	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000943694	2	2021-03-25	2021-12-22	MAH		Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Informa	tion			
Age	Gender	Height	Weight	Report Outcome
49 Years	Female		66 Kilogram	Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
Duplicate	000926151
Duplicate	E2B_04443527

Product Information Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	25.0 Gram		254.0 Minutes	Myasthenic syndrome
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	25.0 Gram		254.0 Minutes	Myasthenic syndrome

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Headache	v.27.1	
Hypertension	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31

173 Report(s)

Report Informatio	n
Adverse Reaction Report Number	L
000944692	
Serious re	p
Seriou	IS
Patient Informatio	n
Age	
59 Years	
Link / Duplicate R	ep
Duplicate	
Product Informati	or

Adverse Reaction Term

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000944692	0	2021-04-06	2021-04-06	Hospital		Spontaneous	Other health professional

Serious report?	Death:	No	Disability:	No	Congenital Anomaly:	No
Serious	Life Threatening:	No	Hospitalization:	No	Other Medically Important Conditions:	Yes

Patient Informa	tion			
Age	Gender	Height	Weight	Report Outcome
59 Years	Female	166 Centimeter	84 Kilogram	Recovering/resolving

Link / Duplicate Report Information	
Record Type	Link AER** Number
Duplicate	E2B_04605968

Product Information							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise	50.0 Gram	1 every 1 Months		Immunodeficien cy

Information		
Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Back pain	v.27.1	
Chills	v.27.1	
Tremor	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

R	Adverse eaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
	000947626	0	2021-04-26	2021-04-26	MAH		Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Informa	tion			
Age	Gender	Height	Weight	Report Outcome
66 Years	Female		57 Kilogram	Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
Duplicate	E2B_04605842

Product Information							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
FLUTICASONE	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	30.0 Gram	1 every 1 Days	1.0 Days	Chronic inflammatory demyelinating polyradiculoneu ropathy

Adverse Reaction Term Information Adverse Reaction Term(s) MedDRA Version V.27.1 Haemolysis V.27.1

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000949075	0	2021-05-06	2021-05-06	MAH		Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Informa	ition			
Age	Gender	Height	Weight	Report Outcome
94 Years	Female			Unknown

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report.	

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	20.0 Gram			Product used for unknown indication

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Acute pulmonary oedema	v.27.1	
Dyspnoea	v.27.1	
Hypertension	v.27.1	
Tachypnoea	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000950358	1	2021-05-17	2021-09-30	Other		Spontaneous	Other health professional

Serious report?	Death:	No	Disability:	No	Congenital Anomaly:	No
Serious	Life Threatening:	No	Hospitalization:	No	Other Medically Important Conditions:	Yes

Patient	Informa	tion

Age	Gender	Height	Weight	Report Outcome
94 Years	Female			Unknown

Link / Duplicate Report Information

Record Type	Link AER** Number

No duplicate or linked report.

Product Information

1 Toddot IIII Officiation							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
AS-3 RBC LR (E7962V00)	Suspect		Intravenous (not otherwise specified)			3.0 Hours	
PANZYGA	Suspect	SOLUTION INTRAVENOUS		150.0 ml		90.0 Minutes	
PLTA-1 LR IRR (E3056V00)	Suspect	NOT SPECIFIED	Intravenous (not otherwise specified)			100.0 Minutes	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Acute pulmonary oedema	v.27.1	
Blood culture negative	v.27.1	
Blood pressure increased	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Dyspnoea	v.27.1	
Hypertension	v.27.1	
Hypervolaemia	v.27.1	
Нурохіа	v.27.1	
Oxygen saturation decreased	v.27.1	
Oxygen therapy	v.27.1	
Respiratory rate increased	v.27.1	
Tachycardia	v.27.1	
Wheezing	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000950469	0	2021-05-18	2021-05-18	MAH		Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Informa	tion			
Age	Gender	Height	Weight	Report Outcome
63 Years	Female		82 Kilogram	Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
Na disellanta an linka dinament	

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
ACETAMINOPHEN	Concomitant	NOT SPECIFIED					
AMOXICILLIN/CLAVULANI C ACID	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	35.0 Gram	Cyclical	2.0 Days	Chronic inflammatory demyelinating polyradiculoneu ropathy
triptan	Concomitant						

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Back pain	v.27.1	
Drug ineffective	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Headache	v.27.1	
Hot flush	v.27.1	
Nausea	v.27.1	
Oxygen saturation decreased	v.27.1	
Sleep apnoea syndrome	v.27.1	
Vomiting	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information	**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000953798	0	2021-06-09	2021-06-09	MAH		Spontaneous	Physician

Serious report?	Death:	Disability:	Congenital Anomaly:	
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Ye	es

Patient Information	
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Age	Gender	Height	Weight	Report Outcome
80 Years	Male		90 Kilogram	Recovered/resolved

Link / Duplicate Report Information

Record Type	Link AER** Number
No duplicate or linked report.	

Product Information

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Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	50.0 Gram			Polymyositis

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Chills	v.27.1	
Dyspnoea	v.27.1	
Hypersensitivity	v.27.1	
Hypertension	v.27.1	
Wheezing	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000954890	0	2021-06-17	2021-06-17	MAH		Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Informa	tion			
Age	Gender	Height	Weight	Report Outcome
61 Years	Female			Recovered/resolved

Link / Duplicate Report Information				
Record Type	Link AER** Number			
No duplicate or linked report.				

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
LIRAGLUTIDE	Concomitant						
PANZYGA	Suspect	SOLUTION INTRAVENOUS		50.0 Gram			Immunodeficien cy

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Chills	v.27.1	
Muscle spasms	v.27.1	
Nausea	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000954896	0	2021-06-17	2021-06-17	Hospital		Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization: Yes	Other Medically Important Conditions: Yes

Patient Informa	tion			
Age Gender		Height	Weight	Report Outcome
51 Years	Female			Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report.	

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	20.0 Gram	1 every 1 Months		
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	30.0 Gram	1 every 1 Months		

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Meningitis aseptic	v.27.1	368 Days

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

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**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000960103	0	2021-07-22	2021-07-22	MAH		Spontaneous	Physician

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization: Yes	Other Medically Important Conditions: Yes

Patient Informa	tion			
Age	Gender	Height	Weight	Report Outcome
51 Years	Female		77 Kilogram	Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report.	

Product	Information
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Report Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	80.0 Gram		142.0 Minutes	Chronic cutaneous lupus erythematosus

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Headache	v.27.1	
Meningitis aseptic	v.27.1	
Nausea	v.27.1	
Vomiting	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000961132	0	2021-07-29	2021-07-29	Hospital		Spontaneous	Other health professional

Serious report?	Death:	No	Disability:	No	Congenital Anomaly: No	
Serious	Life Threatening:	No	Hospitalization:	No	Other Medically Important Conditions: Yes	s

Patient Information

Age	Gender	Height	Weight	Report Outcome	
72 Years	Male		118 Kilogram	Recovered/resolved	

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	25.0 Gram	Once		Squamous cell carcinoma of skin, Pyrexia

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Blood pressure decreased	v.27.1	
Blood pressure increased	v.27.1	
Chills	v.27.1	
Cyanosis	v.27.1	
Heart rate increased	v.27.1	
Hypertension	v.27.1	
Oxygen saturation decreased	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Respiratory rate increased	v.27.1	
Tachycardia	v.27.1	1 Days
Transfusion reaction	v.27.1	
Transfusion-associated dyspnoea	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000963722	1	2021-08-18	2021-08-24	MAH		Spontaneous	Consumer/other non health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization: Yes	Other Medically Important Conditions:

Patient Information

Age	Gender	Height	Weight	Report Outcome
52 Years	Male		79 Kilogram	Unknown

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report.	

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)				Secondary immunodeficien cy

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Asthenia	v.27.1	
Circulatory collapse	v.27.1	
Sepsis	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000965250	0	2021-08-27	2021-08-27	MAH		Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Information

Age	Gender	Height	Weight	Report Outcome
	Male		118 Kilogram	Recovered/resolved

Link / Duplicate Report Information

Record Type	Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	25.0 Gram	Once		Immunodeficien cy
PANZYGA	Suspect	SOLUTION INTRAVENOUS					Immunodeficien cy

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Chills	v.27.1	
Cyanosis	v.27.1	
Dyspnoea	v.27.1	
Hypertension	v.27.1	
Peripheral coldness	v.27.1	

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Tachycardia	v.27.1	
Tachypnoea	v.27.1	
Wheezing	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000969403	3	2021-09-28	2021-12-21	MAH		Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Not Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: No

Patient Informa	tion			
Age	Gender	Height	Weight	Report Outcome
75 Years	Female			Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report.	

Product Information

Froduct information							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	35.0 Gram		56.0 Minutes	Immune thrombocytopen ia
PANZYGA	Suspect	SOLUTION INTRAVENOUS					Immune thrombocytopen ia
PANZYGA	Suspect	SOLUTION INTRAVENOUS					Immune thrombocytopen ia
PANZYGA	Suspect	SOLUTION INTRAVENOUS					Immune thrombocytopen ia
PLATELETS	Suspect	NOT SPECIFIED					
RED BLOOD CELLS	Suspect	NOT SPECIFIED					

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Chills	v.27.1	
Haemolysis	v.27.1	
Headache	v.27.1	
Hypertension	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information	
[2] [0] [0] [0] [1] [0] [1] [1] [1] [1] [1] [1]	

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000981444	0	2021-12-22	2021-12-22	MAH		Spontaneous	Physician

Serious report?	Death:	Disability:	Congenital Anomaly:	
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions:	Yes

Patient Information

	Age	Gender	Height	Weight	Report Outcome
	77 Years	Male	172 Centimeter	65 Kilogram	Recovered/resolved

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS					Immune thrombocytopen ia
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	65.0 Gram			Immune thrombocytopen ia

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Anaemia	v.27.1	
Haemolysis	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000982315	0	2022-01-03	2022-01-03	MAH		Spontaneous	Physician

Serious report?	Death:	Disability:	Congenital Anomaly:	
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Ye	es

Patient Information

Age	Age Gender Height		Weight	Report Outcome	
61 Years	Male		86 Kilogram	Recovered/resolved	

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

i ioaast iiiioiiiiatioii							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	30.0 Gram	Total		Immune thrombocytopen ia
PREDNISONE	Concomitant	NOT SPECIFIED					

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Dizziness	v.27.1	
Hyperhidrosis	v.27.1	
Hypersensitivity	v.27.1	
Hypotension	v.27.1	
Incorrect drug administration rate	v.27.1	
Syncope	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000982374	1	2022-01-04	2022-03-22	МАН		Spontaneous	Consumer/other non health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Information

Age	Gender	Height	Weight	Report Outcome
86 Years	Female			Recovered/resolved

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	70.0 Gram	1 every 1 Days	1.0 Days	Platelet count decreased

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Anaemia	v.27.1	
Anti A antibody positive	v.27.1	
Anti B antibody positive	v.27.1	
Haemolysis	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information **AE

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
000989072	0	2022-02-22	2022-02-22	МАН		Other	Consumer/other non health professional

Serious report?	Death:		Disability:		Congenital Anomaly:	
Serious	Life Threatening:	Yes	Hospitalization:	Yes	Other Medically Important Conditions:	Yes

Patient	Informa	tion

· attorit iiii oi iiia				
Age	Gender	Height	Weight	Report Outcome
15 Years	Male	175 Centimeter	87 Kilogram	Unknown

Link / Duplicate Report Information
Record Type

Record Type	Link AER** Number
Linked	000910579

Product Information

· · · · · · · · · · · · · · · · · · ·							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
DEXAMETHASONE	Concomitant	Tablets					
METHYLPHENIDATE	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	75.0 Gram			Thrombocytope nic purpura
PREDNISONE	Concomitant	NOT SPECIFIED					
RITUXIMAB GP2013	Concomitant						

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Cerebral venous sinus thrombosis	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information	on	**AE	R = Adverse F	Reaction Re	eport									
Adverse Reaction Report Number		ER Version mber	Initial Recei	ved Date	Latest Rece	ived Date	Sourc	e of Report	Market Authorization Holder AER Number				Type of Report	Reporter Type
000989412		0	2022-0	2-23	2022-02-23			MAH			AH		Spontaneous	Consumer/other non health professional
Serious r	eport?				Death:			Disabilit	y:		Congenital	Anomaly:		
Serio	us		L	ife Threat	ening:		Н	ospitalizatio	n: Yes	Other Med	dically Important Co	onditions:		
Patient Information	on .													
Age	Gender		Height	V	Veight		Report	Outcome						
9 Years	Male							ed/resolved						
Link / Duplicate F	Report Info	rmation												
		d Type				Link AER*	* Numb	er						
No duplicate or lin	ked report.													
Product Informat	ion													
Product Desc	ription	Health Pr	oduct Role	Dosag	e Form	Route d Administr		Dose	Fre	equency	Therapy Duration	Indication(s)		
PANZYGA		Sus	spect		JTION /ENOUS	Intravenous otherwis specifie	se	20.0 Gran		every 1 Months		Hypogammaglo bulinaemia		
Adverse Reaction Information	n Term													
	Ad	verse Reac	tion Term(s)			Me	dDRA	Version			Reaction Duration			

v.27.1

Headache

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

Adverse Reaction Term

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
001001084	1	2022-05-10	2022-06-15	Hospital		Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Information	tion			
Age	Gender	Height	Weight	Report Outcome
83 Years	Male			Recovered/resolved

Link / Duplicate Report Information					
Record Type	Link AER** Number				
Duplicate	001053725				

Product Information							
Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	20.0 Gram	1 every 2 Days	80.0 Minutes	Immune thrombocytopen ia
PLATELETS	Concomitant	NOT SPECIFIED				52.0 Minutes	

Information **Adverse Reaction Term(s) MedDRA Version Reaction Duration** Chills v.27.1 Electrocardiogram abnormal v.27.1 v.27.1 Hypertension v.27.1 Lung opacity Oxygen saturation decreased v.27.1 Tachycardia v.27.1

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Tachypnoea	v.27.1	
Transfusion reaction	v.27.1	
Transfusion-associated dyspnoea	v.27.1	
Use of accessory respiratory muscles	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
001012819	0	2022-08-09	2022-08-09	Hospital		Spontaneous	Other health professional

Serious report?	Death:	No	Disability:	No	Congenital Anomaly: No
Serious	Life Threatening:	No	Hospitalization:	No	Other Medically Important Conditions: Yes

Patient	Information

Age	Gender	Height	Weight	Report Outcome
48 Years	Male	181 Centimeter	81 Kilogram	Recovered/resolved

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
ACETAMINOPHEN	Concomitant	NOT SPECIFIED					
BENADRYL	Concomitant	NOT SPECIFIED					
MOXIFLOXACIN	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	35.0 Gram	1 every 4 Weeks		

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Hypersensitivity	v.27.1	
Transfusion reaction	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
001029898	0	2022-12-19	2022-12-19	Community		Spontaneous	Other health professional

Serious report?	Death:	No	Disability:	No	Congenital Anomaly:	No
Serious	Life Threatening:	No	Hospitalization:	No	Other Medically Important Conditions:	Yes

Patient	Informa	tion	
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Age	Gender	Height	Weight	Report Outcome
86 Years	Female	153 Centimeter	50 Kilogram	Recovered/resolved

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS		10.0 Gram	Once		Secondary immunodeficien cy

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
COVID-19	v.27.1	
Chills	v.27.1	
Dyspnoea	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
001030107	0	2022-12-21	2022-12-21	Hospital		Spontaneous	Other health professional

Serious report?	Death:	No	Disability:	No	Congenital Anomaly: No	
Serious	Life Threatening:	Yes	Hospitalization:	No	Other Medically Important Conditions: Ye	s

Patient	Informa	tion

Age	Gender	Height	Weight	Report Outcome
67 Years	Female	152 Centimeter	56 Kilogram	Recovered/resolved

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

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Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
CALCIUM CARBONATE	Concomitant	NOT SPECIFIED					
PANTOPRAZOLE	Concomitant	POWDER FOR SOLUTION INTRAVENOUS					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	5.0 ml		5.0 Minutes	Immune- mediated myositis
PREDNISONE	Concomitant	NOT SPECIFIED					

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Anaphylactic reaction	v.27.1	
Hypertension	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
001042372	0	2023-03-28	2023-03-28	MAH		Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions: Yes

Patient Informa	tion			
Age	Gender	Height	Weight	Report Outcome
63 Years	Female			Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
No duplicate or linked report.	

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
DIPHENHYDRAMINE	Concomitant	NOT SPECIFIED					
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	60.0 Gram		4.0 Hours	Immune thrombocytopen ia

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Chest pain	v.27.1	
Dyspnoea	v.27.1	
Hypersensitivity	v.27.1	
Pruritus	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
001048002	0	2023-05-16	2023-05-16	Hospital		Spontaneous	Other health professional

Serious report?	Death:	No	Disability:	No	Congenital Anomaly: No
Serious	Life Threatening:	Yes	Hospitalization:	No	Other Medically Important Conditions: No

Patient	Informa	tion	

Age	Gender	Height	Weight	Report Outcome
69 Years				Unknown

Link / Duplicate Report Information	
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Record Type	Link AER** Number		
Linked	001070686		

Product Information

Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	75.0 Gram	Once		Myositis

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Chest pain	v.27.1	
Troponin increased	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31

173 Report(s)

Report Information

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
001053725	0	2023-06-22	2023-06-22	MAH		Spontaneous	Other health professional

Serious report?	Death:	Disability:	Congenital Anomaly:
Serious	Life Threatening:	Hospitalization: Yes	Other Medically Important Conditions: Yes

Patient Informa	tion			
Age Gender		Height	Weight	Report Outcome
83 Years	Male	164 Centimeter	87 Kilogram	Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
Duplicate	001001084

Product Informati	tion
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Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	80.0 Gram	1 every 2 Days		Immune thrombocytopen ia
PLATELETS	Concomitant	NOT SPECIFIED					

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Chills	v.27.1	
Hypertension	v.27.1	
Tachycardia	v.27.1	
Transfusion reaction	v.27.1	
Transfusion-associated dyspnoea	v.27.1	
Use of accessory respiratory muscles	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports:

2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information	**∆∤

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
001055286	0	2023-07-05	2023-07-05	MAH		Spontaneous	Physician

Serious report?	Death:	Disability:	Congenital Anomaly:	
Serious	Life Threatening:	Hospitalization:	Other Medically Important Conditions:	Yes

Patient Information

Age	Gender	Height	Weight	Report Outcome
86 Years	Female			Recovered/resolved

Link / Duplicate Report Information

Record Type Link AER** Number

No duplicate or linked report.

Product Information

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Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	10.0 Gram	Once		Secondary immunodeficien cy

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
COVID-19	v.27.1	
Chills	v.27.1	
Dyspnoea	v.27.1	
Febrile nonhaemolytic transfusion reaction	v.27.1	
Hypertension	v.27.1	

Report Runtime: Initial Received Date: Latest Received Date: Total Number of Reports: 2024-12-05 - 07:03:57 AM 1965-01-01 to 2024-07-31 N/A

173 Report(s)

Report Information	

**AER = Adverse Reaction Report

Adverse Reaction Report Number	Latest AER Version Number	Initial Received Date	Latest Received Date		Market Authorization Holder AER Number	Type of Report	Reporter Type
001070686	2	2023-11-08	2023-11-20	MAH		Spontaneous	Consumer/other non health professional

Serious report?	Death:		Disability:		Congenital Anomaly:
Serious	Life Threatening: Ye	'es	Hospitalization:	Yes	Other Medically Important Conditions:

Patient Informa	ation
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Age	Gender	Height	Weight	Report Outcome
69 Years				Recovered/resolved

Link / Duplicate Report Information	
Record Type	Link AER** Number
Linked	001048002

Product Information

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Product Description	Health Product Role	Dosage Form	Route of Administration	Dose	Frequency	Therapy Duration	Indication(s)
PANZYGA	Suspect	SOLUTION INTRAVENOUS	Intravenous (not otherwise specified)	75.0 Gram	Total		Myositis

Adverse Reaction Term(s)	MedDRA Version	Reaction Duration
Chest pain	v.27.1	
Troponin increased	v.27.1	
Vasospasm	v.27.1	