Subject	Rel Day	Aderse					Action			
ID	of Onset	Event	Duration	Intensity	Serious	Related	Taken	Outcome		
Placebo										
Trial Number =	Trial Number = CDISCPILOT01, Site Number = 701, Subject ID = 01-701-1015, Gender = F, Race = WHITE, AGE = 63 Years									
01-701-1015	2	APPLICATION SITE	NA	MILD	N	PROBABLE		NOT RECOVERED/NOT		
		ERYTHEMA	27.4	MI D	N	DD OD A DI E		RESOLVED		
		APPLICATION SITE PRURITUS	NA	MILD	N	PROBABLE		NOT RECOVERED/NOT RESOLVED		
	8	DIARRHOEA	3 DAY	MILD	N	REMOTE		RECOVERED/RESOLVED		
Trial Number =	CDISCPILOT	701, Site Number = 701, Sub	ject ID = 01-7	701-1023, Gender = N	I, Race = WH	ITE, AGE = 64 Years	S			
01-701-1023	3	ERYTHEMA	24 DAY	MILD	N	POSSIBLE		NOT RECOVERED/NOT		
								RESOLVED		
		ERYTHEMA	NA	MODERATE	N	PROBABLE		NOT RECOVERED/NOT RESOLVED		
		ERYTHEMA	24 DAY	MILD	N	POSSIBLE		RECOVERED/RESOLVED		

Trial Number: xxxx

Subject	Rel Day	Aderse					Action			
ID	of Onset	Event	Duration	Intensity	Serious	Related	Taken	Outcome		
Placebo										
Trial Number = CDISCPILOT01, Site Number = 701, Subject ID = 01-701-1023, Gender = M, Race = WHITE, AGE = 64 Years										
01-701-1023	22	ATRIOVENTRICULA R BLOCK SECOND DEGREE	NA	MILD	N	POSSIBLE		NOT RECOVERED/NOT RESOLVED		
Trial Number =	CDISCPILOT	701, Site Number = 701, Sub	iect ID = 01-7	701-1047, Gender = F	, Race = WHI	TE, $AGE = 85 \text{ Years}$				
01-701-1047	1	HIATUS HERNIA	1 DAY	MODERATE	N	NONE		NOT RECOVERED/NOT RESOLVED		
		HIATUS HERNIA	1 DAY	MODERATE	N	NONE		RECOVERED/RESOLVED		
	23	UPPER RESPIRATORY TRACT INFECTION	NA	MILD	N	NONE		NOT RECOVERED/NOT RESOLVED		
	27	BUNDLE BRANCH BLOCK LEFT	NA	MILD	N	NONE		NOT RECOVERED/NOT RESOLVED		
Trial Number =	Trial Number = CDISCPILOT01, Site Number = 701, Subject ID = 01-701-1118, Gender = M, Race = WHITE, AGE = 52 Years									
01-701-1118		COUGH	NA	MILD	N	NONE		NOT RECOVERED/NOT RESOLVED		

Trial Number: xxxx

Subject	Rel Day	Aderse					Action			
ID	of Onset	Event	Duration	Intensity	Serious	Related	Taken	Outcome		
Placebo	Placebo									
Trial Number =	Trial Number = CDISCPILOT01, Site Number = 701, Subject ID = 01-701-1130, Gender = M, Race = WHITE, AGE = 84 Years									
01-701-1130	23	URINARY TRACT INFECTION	8 DAY	MILD	N	NONE		NOT RECOVERED/NOT RESOLVED		
		PYREXIA	1 DAY	MILD	N	NONE		RECOVERED/RESOLVED		
		URINARY TRACT INFECTION	8 DAY	MILD	N	NONE		RECOVERED/RESOLVED		
	84	EYE SWELLING	1 DAY	MILD	N	NONE		RECOVERED/RESOLVED		
		EYE ALLERGY	1 DAY	MILD	N	NONE		RECOVERED/RESOLVED		
	97	EYE PRURITUS	NA	MILD	N	REMOTE		NOT RECOVERED/NOT RESOLVED		
		PRURITUS	NA	MILD	N	REMOTE		NOT RECOVERED/NOT RESOLVED		
		NASAL CONGESTION	NA	MILD	N	REMOTE		NOT RECOVERED/NOT RESOLVED		

Trial Number: xxxx

Subject	Rel Day	Aderse					Action				
ID	of Onset	Event	Duration	Intensity	Serious	Related	Taken	Outcome			
Placebo											
Trial Number =	Trial Number = CDISCPILOT01, Site Number = 701, Subject ID = 01-701-1153, Gender = F, Race = WHITE, AGE = 79 Years										
01-701-1153	21	INCREASED APPETITE	25 DAY	MILD	N	POSSIBLE		NOT RECOVERED/NOT RESOLVED			
		INCREASED APPETITE	25 DAY	MILD	N	POSSIBLE		RECOVERED/RESOLVED			
Xanomeline Hi	Xanomeline High Dose										
Trial Number =	CDISCPILOT	701, Site Number = 701, Sub	piect ID = 01-7	701-1028, Gender = N	I, Race = WH	ITE, AGE = 71 Years	S				
01-701-1028	3	APPLICATION SITE ERYTHEMA	NA	MILD	N	POSSIBLE		NOT RECOVERED/NOT RESOLVED			
	21	APPLICATION SITE PRURITUS	NA	MILD	N	PROBABLE		NOT RECOVERED/NOT RESOLVED			
Trial Number =	CDISCPILOT	701, Site Number = 701, Sub	piect ID = 01-7	701-1034, Gender = F	, Race = WHI	TE, $AGE = 77 \text{ Years}$					
01-701-1034	58	APPLICATION SITE PRURITUS	NA	MILD	N	PROBABLE		NOT RECOVERED/NOT RESOLVED			
	125	FATIGUE	NA	MILD	N	POSSIBLE		NOT RECOVERED/NOT RESOLVED			

Subject	Rel Day	Aderse					Action			
ID	of Onset	Event	Duration	Intensity	Serious	Related	Taken	Outcome		
Xanomeline Hi	Xanomeline High Dose									
Trial Number =	Trial Number = CDISCPILOT01, Site Number = 701, Subject ID = 01-701-1133, Gender = F, Race = WHITE, AGE = 81 Years									
01-701-1133	61	APPLICATION SITE ERYTHEMA	NA	MILD	N	PROBABLE		NOT RECOVERED/NOT RESOLVED		
		APPLICATION SITE PRURITUS	NA	MILD	N	PROBABLE		NOT RECOVERED/NOT RESOLVED		
		APPLICATION SITE ERYTHEMA	NA	MODERATE	N	PROBABLE		NOT RECOVERED/NOT RESOLVED		
		APPLICATION SITE PRURITUS	NA	MODERATE	N	PROBABLE		NOT RECOVERED/NOT RESOLVED		
Trial Number =	Trial Number = CDISCPILOT01, Site Number = 701, Subject ID = 01-701-1146, Gender = F, Race = WHITE, AGE = 75 Years									
01-701-1146	-4	RASH	18 DAY	MILD	N	NONE		NOT RECOVERED/NOT RESOLVED		
		RASH	18 DAY	MILD	N	NONE		RECOVERED/RESOLVED		

Subject	Rel Day	Aderse					Action				
ID	of Onset	Event	Duration	Intensity	Serious	Related	Taken	Outcome			
Xanomeline Hi	Xanomeline High Dose										
Trial Number = CDISCPILOT01, Site Number = 701, Subject ID = 01-701-1146, Gender = F, Race = WHITE, AGE = 75 Years											
01-701-1146	13	APPLICATION SITE IRRITATION	2 DAY	MILD	N	PROBABLE		RECOVERED/RESOLVED			
	15	FATIGUE	NA	MILD	N	POSSIBLE		NOT RECOVERED/NOT RESOLVED			
	22	APPLICATION SITE ERYTHEMA	NA	MILD	N	PROBABLE		NOT RECOVERED/NOT RESOLVED			
		APPLICATION SITE PRURITUS	NA	MILD	N	PROBABLE		NOT RECOVERED/NOT RESOLVED			
		APPLICATION SITE ERYTHEMA	NA	MODERATE	N	PROBABLE		NOT RECOVERED/NOT RESOLVED			
		APPLICATION SITE PRURITUS	NA	MODERATE	N	PROBABLE		NOT RECOVERED/NOT RESOLVED			
	34	APPLICATION SITE PAIN	NA	MODERATE	N	PROBABLE		NOT RECOVERED/NOT RESOLVED			

Trial Number: xxxx

Subject	Rel Day	Aderse					Action				
ID	of Onset	Event	Duration	Intensity	Serious	Related	Taken	Outcome			
Xanomeline Hi	gh Dose										
Trial Number =	Trial Number = CDISCPILOT01, Site Number = 701, Subject ID = 01-701-1146, Gender = F, Race = WHITE, AGE = 75 Years										
01-701-1146	38	APPLICATION SITE VESICLES	NA	MODERATE	N	PROBABLE		NOT RECOVERED/NOT RESOLVED			
		APPLICATION SITE IRRITATION	NA	MODERATE	N	PROBABLE		NOT RECOVERED/NOT RESOLVED			
Trial Number =	CDISCPILOT	(01, Site Number = 701, Sub	ightarrow ject ID = 01-7	701-1148, Gender = N	A, Race = WH	ITE, AGE = 57 Years	S				
01-701-1148	-569	DYSPEPSIA	NA	MODERATE	N	NONE		NOT RECOVERED/NOT RESOLVED			
	-25	DEPRESSED MOOD	NA	MODERATE	N	NONE		NOT RECOVERED/NOT RESOLVED			
	3	APPLICATION SITE ERYTHEMA	NA	MILD	N	PROBABLE		NOT RECOVERED/NOT RESOLVED			
		APPLICATION SITE PRURITUS	NA	MILD	N	PROBABLE		NOT RECOVERED/NOT RESOLVED			
	51	LOWER RESPIRATORY TRACT INFECTION	30 DAY	MODERATE	N	NONE		NOT RECOVERED/NOT RESOLVED			

Trial Number: xxxx

Subject	Rel Day	Aderse					Action		
ID	of Onset	Event	Duration	Intensity	Serious	Related	Taken	Outcome	
Xanomeline High Dose									
Trial Number = CDISCPILOT01, Site Number = 701, Subject ID = 01-701-1148, Gender = M, Race = WHITE, AGE = 57 Years									
01-701-1148	51	LOWER RESPIRATORY TRACT INFECTION	30 DAY	MODERATE	N	NONE		RECOVERED/RESOLVED	
	115	FLANK PAIN	3 DAY	MODERATE	N	NONE		RECOVERED/RESOLVED	
	117	CALCULUS URETHRAL	1 DAY	MODERATE	N	NONE		RECOVERED/RESOLVED	
	134	EPISTAXIS	1 DAY	MILD	N	NONE		RECOVERED/RESOLVED	
	174	ACTINIC KERATOSIS	NA	MILD	N	NONE		NOT RECOVERED/NOT RESOLVED	
Trial Number =	Trial Number = CDISCPILOT01, Site Number = 701, Subject ID = 01-701-1180, Gender = M, Race = WHITE, AGE = 56 Years								
01-701-1180	1	VOMITING	1 DAY	MILD	N	NONE		NOT RECOVERED/NOT RESOLVED	
		VOMITING	1 DAY	MILD	N	NONE		RECOVERED/RESOLVED	

Subject	Rel Day	Aderse					Action				
ID	of Onset	Event	Duration	Intensity	Serious	Related	Taken	Outcome			
Xanomeline Hi	Xanomeline High Dose										
Trial Number =	Trial Number = CDISCPILOT01, Site Number = 701, Subject ID = 01-701-1180, Gender = M, Race = WHITE, AGE = 56 Years										
01-701-1180	4	MICTURITION URGENCY	33 DAY	MILD	N	POSSIBLE		NOT RECOVERED/NOT RESOLVED			
		MICTURITION URGENCY	33 DAY	MILD	N	POSSIBLE		RECOVERED/RESOLVED			
	23	NASOPHARYNGITIS	2 DAY	MILD	N	NONE		RECOVERED/RESOLVED			
	36	APPLICATION SITE VESICLES	NA	MODERATE	N	POSSIBLE		NOT RECOVERED/NOT RESOLVED			
		APPLICATION SITE PRURITUS	NA	MODERATE	N	POSSIBLE		NOT RECOVERED/NOT RESOLVED			
		APPLICATION SITE ERYTHEMA	NA	MODERATE	N	POSSIBLE		NOT RECOVERED/NOT RESOLVED			
		HEADACHE	NA	MODERATE	N	NONE		NOT RECOVERED/NOT RESOLVED			

Trial Number: xxxx

Subject	Rel Day	Aderse					Action				
ID	of Onset	Event	Duration	Intensity	Serious	Related	Taken	Outcome			
Xanomeline Hi	gh Dose										
Trial Number =	Trial Number = CDISCPILOT01, Site Number = 701, Subject ID = 01-701-1181, Gender = F, Race = WHITE, AGE = 79 Years										
01-701-1181	5	AGITATION	1 DAY	MODERATE	N	NONE		RECOVERED/RESOLVED			
Xanomeline Lo	ow Dose										
Trial Number =	CDISCPILOT	701, Site Number = 701, Sub	iect ID = 01-7	701-1097, Gender = N	I, Race = WH	ITE, AGE = 68 Years	S				
01-701-1097	3	ERYTHEMA	NA	MILD	N	POSSIBLE		NOT RECOVERED/NOT			
								RESOLVED			
	51	PRURITUS	1 DAY	MODERATE	N	POSSIBLE		RECOVERED/RESOLVED			
		GENERALISED	2 D 4 W	MIL D	3.7	DD OD A DI E		DECOMEDED/DECOMED			
		APPLICATION SITE VESICLES	3 DAY	MILD	N	PROBABLE		RECOVERED/RESOLVED			
	52	APPLICATION SITE	NA	MILD	N	POSSIBLE		NOT RECOVERED/NOT			
		PRURITUS						RESOLVED			
		APPLICATION SITE	NA	MODERATE	N	POSSIBLE		NOT RECOVERED/NOT			
		PRURITUS						RESOLVED			
	80	PRURITUS	1 DAY	MODERATE	N	POSSIBLE		RECOVERED/RESOLVED			
		GENERALISED									

Subject	Rel Day	Aderse					Action				
ID	of Onset	Event	Duration	Intensity	Serious	Related	Taken	Outcome			
Xanomeline Lo	Xanomeline Low Dose										
Trial Number = CDISCPILOT01, Site Number = 701, Subject ID = 01-701-1097, Gender = M, Race = WHITE, AGE = 68 Years											
01-701-1097	90	PRURITUS GENERALISED	1 DAY	MODERATE	N	POSSIBLE		RECOVERED/RESOLVED			
	109	PRURITUS GENERALISED	2 DAY	MODERATE	N	POSSIBLE		RECOVERED/RESOLVED			
		PHARYNGOLARYNG EAL PAIN	4 DAY	MILD	N	NONE		RECOVERED/RESOLVED			
		NASAL CONGESTION	4 DAY	MILD	N	NONE		RECOVERED/RESOLVED			
Trial Number =	CDISCPILOT	Γ01, Site Number = 701, Sub	ject ID = 01-7	701-1111, Gender = F	, Race = WHI	TE, AGE = 81 Years					
01-701-1111	-61	LOCALISED INFECTION	NA	MODERATE	N	NONE		NOT RECOVERED/NOT RESOLVED			
	-5	ERYTHEMA	6 DAY	MILD	N	NONE		NOT RECOVERED/NOT RESOLVED			
		PRURITUS	6 DAY	MILD	N	NONE		NOT RECOVERED/NOT RESOLVED			

Subject	Rel Day	Aderse					Action				
ID	of Onset	Event	Duration	Intensity	Serious	Related	Taken	Outcome			
Xanomeline Lo	Xanomeline Low Dose										
Trial Number = CDISCPILOT01, Site Number = 701, Subject ID = 01-701-1111, Gender = F, Race = WHITE, AGE = 81 Years											
01-701-1111	-5	ERYTHEMA	6 DAY	MILD	N	NONE		RECOVERED/RESOLVED			
		PRURITUS	6 DAY	MILD	N	NONE		RECOVERED/RESOLVED			
	1	MICTURITION URGENCY	NA	MILD	N	NONE		NOT RECOVERED/NOT RESOLVED			
	7	ARTHRALGIA	NA	MODERATE	N	NONE		NOT RECOVERED/NOT RESOLVED			
		CELLULITIS	NA	MODERATE	N	NONE		NOT RECOVERED/NOT RESOLVED			
Trial Number =	CDISCPILOT	Γ 01, Site Number = 701, Sub	piect ID = 01-7	701-1115, Gender = N	A, Race = WH	ITE, AGE = 84 Years	S				
01-701-1115	3	APPLICATION SITE ERYTHEMA	NA	MILD	N	PROBABLE		NOT RECOVERED/NOT RESOLVED			
		APPLICATION SITE PRURITUS	NA	MILD	N	PROBABLE		NOT RECOVERED/NOT RESOLVED			

Subject	Rel Day	Aderse					Action		
ID	of Onset	Event	Duration	Intensity	Serious	Related	Taken	Outcome	
Xanomeline Lo	Xanomeline Low Dose								
Trial Number = CDISCPILOT01, Site Number = 701, Subject ID = 01-701-1115, Gender = M, Race = WHITE, AGE = 84 Years									
01-701-1115	3	APPLICATION SITE ERYTHEMA	NA	MODERATE	N	PROBABLE		NOT RECOVERED/NOT RESOLVED	
		APPLICATION SITE PRURITUS	NA	MODERATE	N	PROBABLE		NOT RECOVERED/NOT RESOLVED	
	27	APPLICATION SITE IRRITATION	1 DAY	MILD	N	POSSIBLE		NOT RECOVERED/NOT RESOLVED	
		APPLICATION SITE IRRITATION	1 DAY	MILD	N	POSSIBLE		RECOVERED/RESOLVED	
	46	FATIGUE	NA	MODERATE	N	REMOTE		NOT RECOVERED/NOT RESOLVED	
	51	APPLICATION SITE IRRITATION	NA	MODERATE	N	PROBABLE		NOT RECOVERED/NOT RESOLVED	
		APPLICATION SITE DERMATITIS	NA	MODERATE	N	PROBABLE		NOT RECOVERED/NOT RESOLVED	

Subject	Rel Day	Aderse					Action		
ID	of Onset	Event	Duration	Intensity	Serious	Related	Taken	Outcome	
Xanomeline Lo	Xanomeline Low Dose								
Trial Number = CDISCPILOT01, Site Number = 701, Subject ID = 01-701-1188, Gender = M, Race = WHITE, AGE = 71 Years									
01-701-1188	2	ERYTHEMA	NA	MILD	N	PROBABLE		NOT RECOVERED/NOT	
								RESOLVED	
		PRURITUS	NA	MILD	N	PROBABLE		NOT RECOVERED/NOT	
								RESOLVED	
		ERYTHEMA	NA	MODERATE	N	PROBABLE		NOT RECOVERED/NOT	
								RESOLVED	
		PRURITUS	NA	MODERATE	N	PROBABLE		NOT RECOVERED/NOT	
								RESOLVED	
	18	URTICARIA	NA	MILD	N	PROBABLE		NOT RECOVERED/NOT	
								RESOLVED	
		URTICARIA	NA	MODERATE	N	PROBABLE		NOT RECOVERED/NOT	
								RESOLVED	
	35	APPLICATION SITE	NA	MILD	N	PROBABLE		NOT RECOVERED/NOT	
		URTICARIA						RESOLVED	

Trial Number: xxxx

Subject	Rel Day	Aderse					Action			
ID	of Onset	Event	Duration	Intensity	Serious	Related	Taken	Outcome		
Xanomeline Lo	Xanomeline Low Dose									
Trial Number =	Trial Number = CDISCPILOT01, Site Number = 701, Subject ID = 01-701-1188, Gender = M, Race = WHITE, AGE = 71 Years									
01-701-1188	35	URTICARIA	NA	MILD	N	PROBABLE		NOT RECOVERED/NOT RESOLVED		
Trial Number =	Trial Number = CDISCPILOT01, Site Number = 701, Subject ID = 01-701-1192, Gender = F, Race = WHITE, AGE = 80 Years									
01-701-1192	-782	COUGH	NA	MODERATE	N	NONE		NOT RECOVERED/NOT RESOLVED		
	13	NASAL MUCOSA BIOPSY	1 DAY	MILD	N	NONE		RECOVERED/RESOLVED		
	17	SECRETION DISCHARGE	39 DAY	MILD	N	NONE		NOT RECOVERED/NOT RESOLVED		
		ERYTHEMA	39 DAY	MILD	N	NONE		NOT RECOVERED/NOT RESOLVED		
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Source: [Study MK9999P001: adam-adae]