

Referral Status:			MRN:			
	New referral		Order change			Order Renewal
Patient preferred clinic:						

<u> </u>	iair" (omaiizumab) Standard P	ian of Treatm	ent	for Orticaria	1						
PAT	TIENT DEMOGRAPHICS:										
Date of Referral:				Patient's Phone:							
Pati	ient Name:		Address:								
Date	e of Birth:		City,	City, State, Zip:							
Heiç	ght in inches: Weight: LB	or KG	Gend	ler:	Allergies:	See list	NKDA				
DIA	AGNOSIS: (PLEASE COMPLETE 2 <sup>ND</sup> AND 3	3 <sup>RD</sup> DIGITS TO CON	MPLE	TE ICD 10 FOR I	BILLING)						
	L50.0 - Allergic Urticaria			L50.1 - Chronic Idiopathic Urticaria (CIU)							
	L50.8 - Other Specified Urticaria			Other:							
REC	QUESTED DOCUMENTATION:	PREVIOUS ADMINI	STRA	TION: HAS THIS F	PATIENT TAKEN THIS M	MEDICATION BEF	ORE?				
1	Insurance information	IF NO:	IF YE	IF YES:							
2	Most recent History & Physical	PLEASE STATE	LAST INFUSION DATE:								
3	Full medication list	REQUIRED WASHOUT FROM PREVIOUS	NEXT INFUSION DATE:								
4	Tried and failed therapies	I failed therapies THERAPY:		IF ORDER CHANGE:							
5	Pre-treatment serum IgE level as required for pre treatment dosing			Continue current order until insurance approved							
	Pro	vider Attestation	for I	HCP administra	ation:						
	Provider attests that the patient or caregiver is not compe unable to administer the Xolair labeled self-administration		The location and circumstances for self-administration are not adequate for the potential treatment of anaphylaxis should that arise.								
	Patient has experienced severe hypersensitivity reactions other agents, such as foods, drugs, biologics, within the administration and direct monitoring by a healthcare prof		Patient has a history of uncontrolled disease and ordering physician attests that in their clinical opinion, it is not advisable to try the self-administration formulation of requested drug								
Patient has not received at least 3 doses of Xolair under the guidance of a healthcare provider with no hypersensitivity reactions*				Due to patient's weight, ordering provider attests that in their clinical opinion, it is not advisable to try the self-administered formulation of requested drug.							
*Spe	ecific reactions:										
_	DICATION ORDERS:										
		umab) if patient has sign	ıs/svm	ptoms of parasitic in	fection, is currently being to	reated for a parasition	c infection, or				
NOTE: Patient may be ineligible to receive Xolair® (omalizumab) if patient has signs/symptoms of parasitic infection, is currently being treated for a parasitic infection, or is having acute bronchospasm and/or asthma attack.											
MF	DICATION/FREQUENCY:										
	Xolair® (omalizumab) subcutaneously e	everv 4 weeks:		Other:							
DΩ	SE:	overy i weeke.		] •							
טט	150 mg/dose 300mg/dose										
		aubautanaaua inia	otion	to upper orm th	niah arahdaman						
CDI		subcutaneous inje	Cuon	to upper arm, th	ngn, or abdomen.						
SPE	ECIAL ORDERS:										
	ST WAIT: Extended post treatment monitoring										
	ndard Palmetto Infusion Post wait per package						njection, for				
30 H	ninutes after third injection, then monitor for 19		•	•		below.					
	Monitor patient for two (2) hours after first 3 injections, and for 30-minutes after all subsequent injections.										
	Provider specific post wait:			Refills x 12 months unless noted otherwise here:							
	RE ORDERS:			ADVERSE REACTION & ANAPHYLAXIS ORDERS:							
<b>\</b>	Provide nursing care per Palmetto Infusion			Administer acute infusion and anaphylaxis medications per Palmetto Infusion standing adverse reaction orders, which can							
	Procedures and post procedure observation	n II Indicated		be found at our website or scan here.							
PRE	ESCRIBER INFORMATION:										
PRO	OVIDER NAME:			PHONE:							
ADI	DRESS:			FAX:							
	Y, STATE, ZIP:			NPI:							
		turos)		1.4		DATE					
PKE	ESCRIBER SIGNATURE: (No stamp signa	tures				DATE					
	Dispense as written/Brand medically	necessary			Substitution permit	ted					