Operating Room Management Committee

New Product/Procedure Request

VENDOR TO COMPLETE SHADED SECTION ONLY

New Product Name: The Reltok Clear-Flo™ Nasal Airway	Manufacturer: Reltok Nasal Products, LLC 9735 Wilshire Blvd. Suite 220, Beverly Hills, CA 90212			
Catalog #(s):	Cost of Equipment: N/A Cost of Disposables:			
Sales Rep:	Contact Information: Phone: Email:			
Is this product / equipment FDA approved for the application being requested? X YES NO If yes, please provide a copy of the FDA Approval Letter. Please list the approved FDA applications for this product / equipment: Nasopharyngeal Airway. Is there an approved CSMC IRB Study related to this product / equipment? YES X NO				
If yes, please provide IRB number. The requesting MD must also provide us with a copy of the Patient Consent Form approved by the IRB Office. If no, the requesting MD must provide a copy of the Patient Consent Form clearly indicating the risks associated with the off-label use. The following should also be attached: 1. Reltok Nasal Airway Research Paper, 2014 2. JAMA Facial Plastic Surgery Journal Article, September 6, 2018 3. Claim Appeal Letter				
What training / certification do you recommend for use of this new product / equipment? Not necessary. All nasal surgeons are familiar with insertion process.				
ls it generally recommended that surgeons be proctored to use	e this product / equipment? YES X NO			
Will you supply product / equipment at no charge for the purpose of evaluation? X YES NO				

SURGEON / CN IV / AUTHORIZED HOSPITAL PERSONNEL TO COMPLETE SECTIONS 1 THROUGH 7

1. New product / equipment will be used for the following type(s) of procedures / applications:			
a) Estimated # of procedures perfo	rmed in a year:		
b) Please provide a brief explanation of how this new product / equipment will add value:			
c) Will the new product / equipment of yes, please identify.	require additional e	quipment or supplies	not currently available?
Please note: You are required to submit po	eer-reviewed studies t	o support the efficac	y of the new product / equipment.
2. Please explain the clinical advantage(s) of the new product ,	/ equipment over cur	rrent method.
a) Will the new product / equipmen	nt improve patient out	come? If yes, please	explain.
Nasal Airway is a safety device. Red	uces possibility of airw	av obstruction and as	sphyxiation.
Endorsed by anesthesiologists. See attached Anesthesiologist Endorsement for Intranasal Airway.			
b) Will the new product / equipmen	nt enable us to treat m	ore or a different clas	ss of patients? If yes, please explain.
No. For all nasal and sinus surgery co	IS O S		
c) Standard of Care. Please list lead		at use the new produ	uct / equipment.
			or oquipment.
UCI Irvine; NYU Langione Medical Center; Ohio State University Health System; Wilkes Regional Medical Center, North Carolina; Marion General Hospital, Indianapolis, IN			
3. Please describe how you are currently p			
4. Please identify the product / equipmen U.S. Patent and Post-Op Nasal Airway System		_	v and unique. See attached Nasal Airway
5. In your opinion, what would be a fair ar your evaluation?	mount of time, or quai	ntity of product, / equ	uipment, in order for you to complete
6. Is vendor presence required in O.R.? YES X NO For extended use? YES X NO			
7. Do you have financial interest or investment with this vendor or distributor? YES X NO			
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EVALUATION REQUESTED BY:			
Print Name	Signature		Date
Phone	Email		
Service Line Director		Date	
O.R. CNIV / CNIII		Date	