CERTIFICATE OF MEDICAL NECESSITY

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	OXY	GEN							
SECTION A Cer	tification Type/Date: INITIAL//_	/ REVISED//_ RECERTIFICATION _//_							
PATIENT NAME, ADDRESS	S, TELEPHONE and HIC NUMBER SU	SUPPLIER NAME, ADDRESS, TELEPHONE and NSC NUMBER							
()	_	() NSC #							
PLACE OF SERVICE NAME and ADDRESS of FA	110. 00 0022	PT DOB//; Sex (M/F); HT(in.); WT(lbs.)							
Reverse)		YSICIAN NAME, ADDRESS, TELEPHONE and UPIN NUMBER							
SECTION B		Be Completed by the Supplier of the Items/Supplies.							
EST. LENGTH OF NEED (# OF MONTHS): 1-99 (99=LIFETIME) DIAGNOSIS CODES (ICD-9):									
ANSWERS	·								
ANSWERS ANSWER QUESTIONS 1-10. (Circle Y for Yes, N for No, or D for Does Not Apply, unless otherwise noted.) a) mm Hg b) % c)//									
Y N	Was the test in Question 1 performed EITHE days prior to discharge from an inpatient facil	R with the patient in a chronic stable state as an outpatient OR within two ity to home?							
1 2 3		test in Question 1: (1) At Rest; (2) During Exercise; (3) During Sleep							
XXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXX	Physician/provider performing test in Question 1 (and, if applicable, Question 7). Print/type name and address below: NAME: ADDRESS:								
Y N D	5. If you are ordering portable oxygen, is the patient mobile within the home? If you are <u>not</u> ordering portable oxygen, circle D.								
LPM	6. Enter the highest oxygen flow rate ordered for	r this patient in liters per minute. If less than 1 LPM, enter a "X".							
a) mm Hg b)% c)//_	PO and the this control of the test with a fact in a description of the test. Established the state of the test to								
,	1 56–59 OR OXYGEN SATURATION = 89%, AT LE <i>i</i>	AST ONE OF THE FOLLOWING CRITERIA MUST BE MET.							
Y N D	8. Does the patient have dependent edema due to congestive heart failure?								
Y N D	9. Does the patient have cor pulmonale or pulmonary hypertension documented by P pulmonale on an EKG or by an echocardiogram, gated blood pool scan or direct pulmonary artery pressure measurement?								
Y N D	10. Does the patient have a hematocrit greater th	an 56%?							
NAME OF PERSON ANS NAME:	SWERING SECTION B QUESTIONS, IF OTHER TH	HAN PHYSICIAN (Please Print): EMPLOYER:							
SECTION C		cription of Equipment and Cost							
(1) Narrative description		Supplier's charge and (3) Medicare Fee Schedule Allowance for <u>each</u> item,							
SECTION D	Physician A	ttestation and Signature/Date							
I certify that I am the treating physician identified in Section A of this form. I have received Sections A, B and C of the Certificate of Medical Necessity (including charges for items ordered). Any statement on my letterhead attached hereto, has been reviewed and signed by me. I certify that the medical necessity information in Section B is true, accurate and complete, to the best of my knowledge, and I understand that any falsification, omission, or concealment of material fact in that section may subject me to civil or criminal liability. PHYSICIAN'S SIGNATURE									

SECTION A: (May be completed by the supplier)

CERTIFICATION TYPE/DATE:

If this is an initial certification for this patient, indicate this by placing date (MM/DD/YY) needed initially in the space marked "INITIAL." If this is a revised certification (to be completed when the physician changes the order, based on the patient's changing clinical needs), indicate the initial date needed in the space marked "INITIAL," <u>and also</u> indicate the recertification date in the space marked "REVISED." If this is a recertification, indicate the initial date needed in the space marked "INITIAL," <u>and also</u> indicate the recertification date in the space marked "RECERTIFICATION." Whether submitting a REVISED or a RECERTIFIED CMN, be sure to always furnish the INITIAL date as well as the REVISED <u>or</u> RECERTIFICATION date.

PATIENT INFORMATION:

Indicate the patient's name, permanent legal address, telephone number and his/her health insurance claim number (HICN) as it appears on his/her Medicare card and on the claim form.

SUPPLIER INFORMATION:

Indicate the name of your company (supplier name), address and telephone number along with the Medicare Supplier Number assigned to you by the National Supplier Clearinghouse (NSC).

PLACE OF SERVICE:

Indicate the place in which the item is being used, i.e., patient's home is 12, skilled nursing facility (SNF) is 31, End Stage Renal Disease (ESRD) facility is 65, etc. Refer to the DMERC supplier manual for a complete list.

FACILITY NAME:

If the place of service is a facility, indicate the name and complete address of the facility.

HCPCS CODES:

List all HCPCS procedure codes for items ordered that require a CMN. Procedure codes that do not require certification should not be listed on the CMN.

PATIENT DOB, HEIGHT, WEIGHT AND SEX:

Indicate patient's date of birth (MM/DD/YY) and sex (male or female); height in inches and weight in pounds, if requested.

PHYSICIAN NAME, ADDRESS:

Indicate the physician's name and complete mailing address.

UPIN:

Accurately indicate the treating physician's Unique Physician Identification Number (UPIN).

PHYSICIAN'S TELEPHONE NO:

SECTION B:

Indicate the telephone number where the physician can be contacted (preferably where records would be accessible pertaining to this patient) if more information is needed.

(May not be completed by the supplier. While this section may be completed by a non-physician clinician, or a physician employee, it must be reviewed, and the CMN signed (in Section D) by the treating physician.)

EST. LENGTH OF NEED:

Indicate the estimated length of need (the length of time the physician expects the patient to require use of the ordered item) by filling in the appropriate number of months. If the physician expects that the patient will require the item for the duration of his/her life, then enter 99.

DIAGNOSIS CODES:

In the first space, list the ICD9 code that represents the primary reason for ordering this item. List any additional ICD9 codes that would further describe the medical need for the item (up to 3 codes).

QUESTION SECTION:

This section is used to gather clinical information to determine medical necessity. Answer each question which applies to the items ordered, circling "Y" for yes, "N" for no, "D" for does not apply, a number if this is offered as an answer option, or fill in the blank if other information is requested.

NAME OF PERSON ANSWERING SECTION B QUESTIONS: If a clinical professional other than the treating physician (e.g., home health nurse, physical therapist, dietician)

or a physician employee answers the questions of Section B, he/she must <u>print</u> his/her name, give his/her professional title and the name of his/her employer where indicated. If the physician is answering the questions, this space may be left blank.

SECTION C: (To be completed by the supplier)

NARRATIVE DESCRIPTION OF EQUIPMENT & COST: Supplier gives (1) a narrative description of the item(s) ordered, as well as all options, accessories, supplies and drugs; (2) the supplier's charge for each item, option, accessory, supply and drug; and (3) the Medicare fee schedule allowance for each item/option/accessory/supply/drug, if applicable.

SECTION D: (To be completed by the physician)

PHYSICIAN ATTESTATION:

The physician's signature certifies (1) the CMN which he/she is reviewing includes Sections A, B, C and D; (2) the answers in Section B are correct; and (3) the self-identifying information in Section A is correct.

PHYSICIAN SIGNATURE AND DATE:

After completion and/or review by the physician of Sections A, B and C, the physician must sign and date the CMN in Section D, verifying the Attestation appearing in this Section. The physician's signature also certifies the items ordered are medically necessary for this patient. Signature and date stamps are not acceptable.

According to the Paperwork Reduction Act of 1995, no persons are required to respond to a collection of information unless it displays a valid OMB control number. The valid OMB control number for this information collection is 9338-0534. The time required to complete this information collection is estimated to average 10 minutes per response, including the time to review instructions, search existing resources, gather the data needed, and complete and review the information collection. If you have any comments concerning the accuracy of the time estimate or suggestions for improving this form, please write to: HCFA, 7500 Security Blvd. N2-14-26, Baltimore, Maryland 21244-1850 and to the Office of Information and Regulatory Affairs, Office of Management and Budget, Washington, D.C. 20503.