

# <u>Long-Acting Granulocyte Colony Stimulating Factors (LA-gCSF)</u>: Neulasta®; Fulphila®; Udenyca®; Ziextenzo®; Nyvepria™; Fylnetra®; Stimufend®; Rolvedon®; Ryzneuta®

(Subcutaneous)

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#### I. Length of Authorization <sup>1-9,16-21</sup>

- Bone marrow transplantation (BMT) failure or engraftment delay: Coverage will be provided for 1 dose only and may not be renewed.
- Peripheral blood progenitor cell (PBPC) mobilization and transplant: Coverage will be provided for 1 dose only and may not be renewed.
- Acute exposure to myelosuppressive doses of radiation (Hematopoietic Acute Radiation Syndrome [H-ARS]): Coverage will be provided for 2 doses and may not be renewed.
- All other indications: Coverage will be provided for four months and may be renewed unless otherwise specified.

## II. Dosing Limits

## A. Quantity Limit (max daily dose) [NDC Unit]:

- Neulasta 6 mg single-dose prefilled syringe: 1 syringe per 14 days
- Neulasta 6 mg single-dose prefilled syringe Onpro kit: 1 kit per 14 days
- Fulphila 6 mg single-dose prefilled syringe: 1 syringe per 14 days
- Udenyca 6 mg single-dose prefilled syringe: 1 syringe per 14 days
- Udenyca 6 mg single-dose prefilled autoinjector: 1 autoinjector per 14 days
- Udenyca 6 mg single-dose prefilled syringe ONBODY kit: 1 kit per 14 days
- Ziextenzo 6 mg single-dose prefilled syringe: 1 syringe per 14 days
- Nyvepria 6 mg single-dose prefilled syringe: 1 syringe per 14 days
- Fylnetra 6 mg single-dose prefilled syringe: 1 syringe per 14 days
- Stimufend 6 mg single-dose prefilled syringe: 1 syringe per 14 days



- Rolvedon 13.2 mg single-dose prefilled syringe: 1 syringe per 14 days
- Ryzneuta 20 mg single-dose prefilled syringe: 1 syringe per 14 days

#### B. Max Units (per dose and over time) [HCPCS Unit]:

Drug Name	Indication	Billable Units
Neulasta, Fulphila, Udenyca, Ziextenzo,	Acute Radiation Exposure	12 billable units weekly x 2 doses
Nyvepria, Fylnetra, and Stimufend	BMT failure or engraftment delay/ PBPC mobilization and transplant	12 billable units x 1 dose
	All other indications	12 billable units per 14 days
Rolvedon	Acute Radiation Exposure	132 billable units weekly x 2 doses
	All other indications	132 billable units per 14 days
Ryzneuta	Acute Radiation Exposure	20 mg weekly x 2 doses
	All other indications	20 mg per 14 days

## III. Initial Approval Criteria <sup>1-9</sup>

Coverage is provided in the following conditions:

- Patient must try and have an inadequate response, contraindication, or intolerance to Neulasta AND Fulphila, **OR**
- Patient is continuing treatment with a different peg-filgrastim product
- Patient is at least 18 years of age (Rolvedon and Ryzneuta ONLY); AND

#### Prophylactic use in patients with solid tumors or non-myeloid malignancy † ‡ 1-12,22,24-30

- Patient is undergoing myelosuppressive chemotherapy with an expected incidence of febrile neutropenia ♣ of > 20% §; OR
- Patient is undergoing myelosuppressive chemotherapy with an expected incidence of febrile neutropenia� of 10% to 20% § AND <u>one</u> or more patient-related risk factors ¥; OR

**\*\***Use in this setting is based on clinical judgment.

**Note**: Dose-dense therapy, in general, requires growth factor support to maintain dose intensity and schedule. In the palliative setting, consideration should be given to dose reduction or change in regimen.

Patient who experience a neutropenic complication from a prior cycle of the same chemotherapy  $\ddagger$   $^{11,12}$ 



**Note**: Dose-dense therapy, in general, requires growth factor support to maintain dose intensity and schedule. In the palliative setting, consideration should be given to dose reduction or change in regimen.

Patients acutely exposed to myelosuppressive doses of radiation (Hematopoietic Acute Radiation Syndrome [H-ARS])  $\dagger \ddagger \Phi$  <sup>1,3,4,7,11,12,31,32</sup>

Bone marrow transplantation (BMT) failure or engraftment delay ‡ <sup>16-20</sup> (Neulasta, Fulphila, Udenyca, Ziextenzo, Nyvepria, Fylnetra, and Stimufend ONLY)

Peripheral blood progenitor cell (PBPC) mobilization and transplant ‡ <sup>11</sup> (Neulasta, Fulphila, Udenyca, Ziextenzo, Nyvepria, Fylnetra, and Stimufend ONLY)

Wilms Tumor (Nephroblastoma) ‡<sup>11</sup> (Neulasta, Fulphila, Udenyca, Ziextenzo, Nyvepria, Fylnetra, and Stimufend ONLY)

- Patient has favorable histology disease; AND
- Used in combination with a cyclophosphamide-based chemotherapy regimen (i.e., Regimen M or Regimen I only)

FDA Approved Indication(s); Compendia Recommended Indication(s); Orphan Drug

#### ¥ Patient risk factors for febrile neutropenia <sup>12</sup>

- Age >65 years receiving full dose intensity chemotherapy
- Prior exposure to chemotherapy or radiation therapy
- Persistent neutropenia (ANC  $\leq$  1000/mm3)
- Bone marrow involvement by tumor
- Patient has a condition that can potentially increase the risk of serious infection (i.e., HIV/AIDS with low CD4 counts)
- Recent surgery and/or open wounds
- Poor performance status
- Renal dysfunction (creatinine clearance <50 mL/min)
- Liver dysfunction (elevated bilirubin >2.0 mg/dL)
- $\bullet \quad {\rm Chronic\ immunosuppression\ in\ the\ post-transplant\ setting,\ including\ organ\ transplant}$

#### Febrile neutropenia is defined as: <sup>12</sup>

- <u>Temperature</u>: a single temperature ≥38.3 °C orally or ≥38.0 °C over 1 hour; **AND**
- <u>Neutropenia</u>: <500 neutrophils/mcL or <1,000 neutrophils/mcL and a predicted decline to ≤500 neutrophils/mcL over the next 48 hours

§ Expected incidence of febrile neutropenia percentages for myelosuppressive chemotherapy regimens can be found in the NCCN Hematopoietic Growth Factors Clinical Practice Guideline at NCCN.org <sup>12</sup>

## IV. Renewal Criteria <sup>1-9,16-21</sup>

Coverage for all other indications can be renewed based upon the following criteria:

 

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- Patient continues to meet indication-specific relevant criteria such as concomitant therapy requirements (not including prerequisite therapy), performance status, etc. identified in section III; **AND**
- Absence of unacceptable toxicity from the drug. Examples of unacceptable toxicity include: splenic rupture, acute respiratory distress syndrome (ARDS), serious allergic reactions/anaphylaxis, sickle cell crisis, glomerulonephritis, leukocytosis, thrombocytopenia, capillary leak syndrome, potential for tumor growth stimulation of malignant cells, aortitis, myelodysplastic syndrome and acute myeloid leukemia in patients with breast and lung cancer, etc.; **AND**

Bone marrow transplantation (BMT) failure or engraftment delay (Neulasta, Fulphila, Udenyca, Ziextenzo, Nyvepria, Fylnetra, and Stimufend ONLY)

• Coverage may not be renewed

**Peripheral blood progenitor cell (PBPC) mobilization and transplant** (Neulasta, Fulphila, Udenyca, Ziextenzo, Nyvepria, Fylnetra, and Stimufend ONLY)

• Coverage may not be renewed

Acute exposure to myelosuppressive doses of radiation (Hematopoietic Acute Radiation Syndrome [H-ARS])

• Coverage may not be renewed

# V. Dosage/Administration <sup>1-9,16-21</sup>

#### Neulasta, Fulphila, Udenyca, Ziextenzo, Nyvepria, Fylnetra, and Stimufend

Indication	Dose
Prophylactic use in patients with non-myeloid malignancy	• Administer 6 mg subcutaneously once per chemotherapy cycle and dosed no more frequently than every 14 days
Patient who experienced a	• For pediatric patients weighing <45 kg:
neutropenic complication from a prior	- <10 kg = 0.1 mg/kg
cycle of the same chemotherapy	- 10-20 kg = 1.5 mg
	- 21-30 kg = 2.5 mg
Wilms Tumor (Nephroblastoma)	- 31-44 kg = 4 mg
Acute Radiation Exposure	• Administer 6 mg subcutaneously weekly x 2 doses
(Hematopoietic Acute Radiation	• For pediatric patients weighing <45 kg:
Syndrome)	- <10 kg = 0.1 mg/kg
	- 10-20 kg = 1.5 mg
	- 21-30 kg = 2.5 mg
	- 31-44 kg = 4 mg

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BMT failure or engraftment delay	Administer 6 mg subcutaneously for 1 dose only
PBPC mobilization and transplant	
NOTE	

#### NOTE:

- Do not administer within 14 days before and 24 hours after administration of cytotoxic chemotherapy.
- Use of the pre-filled syringe products may be self-administered or administered by a caregiver or healthcare professional.
- A healthcare provider must fill the on-body injector with Neulasta or Udenyca using the prefilled • syringe and then apply the on-body injector to the patient's skin (abdomen or back of arm).
- On-body Injectors may be applied on the same day as chemotherapy as long as the Neulasta or Udenyca is administered no less than 24 hours after administration of chemotherapy. Not recommended for use in patients with acute radiation exposure or in pediatric patients.

#### Rolvedon

Dose
• Administer 13.2 mg subcutaneously once per chemotherap cycle approximately 24 hours after cytotoxic chemotherapy
• Administer 13.2 mg subcutaneously weekly x 2 doses

- Do not administer within 14 days before and 24 hours after administration of cytotoxic chemotherapy.
- Rolvedon may be self-administered or administered by a caregiver or healthcare professional.

## <u>Ryzneuta</u>

Indication	Dose
Prophylactic use in patients with non-myeloid malignancy Patient who experienced a neutropenic complication from a prior cycle of the same chemotherapy	• Administer 20 mg subcutaneously once per chemotherapy cycle at least 24 hours after cytotoxic chemotherapy.
Acute Radiation Exposure (Hematopoietic Acute Radiation Syndrome)	• Administer 20 mg subcutaneously weekly x 2 doses

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	FYLNETRA <sup>®</sup> ; STIMUFEND <sup>®</sup> ; ROLVEDON <sup>®</sup> ; RYZNEUTA <sup>®</sup> )	Magg
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#### <u>NOTE:</u>

- Do not administer within 14 days before and <24 hours after administration of cytotoxic chemotherapy.
- Ryzneuta is administered subcutaneously via a single-dose prefilled syringe by a healthcare professional.

# VI. Billing Code/Availability Information

#### HCPCS Code(s):

- J2506 Injection, pegfilgrastim, excludes biosimilar, 0.5 mg; 1 billable unit = 0.5 mg (*Neulasta only*)
- Q5108 Injection, pegfilgrastim-jmdb, biosimilar, (Fulphila), 0.5 mg; 1 billable unit = 0.5 mg
- Q5111 Injection, pegfilgrastim-cbqv, biosimilar, (Udenyca), 0.5 mg; 1 billable unit = 0.5 mg
- Q5120 Injection, pegfilgrastim-bmez, biosimilar, (Ziextenzo), 0.5 mg; 1 billable unit = 0.5 mg
- Q5122 Injection, pegfilgrastim-apgf, biosimilar, (Nyvepria), 0.5 mg; 1 billable unit = 0.5 mg
- Q5127 Injection, pegfilgrastim-fpgk, biosimilar, (Stimufend), 0.5 mg; 1 billable unit = 0.5 mg
- Q5130 Injection, pegfilgrastim-pbbk, biosimilar, (Fylnetra), 0.5 mg; 1 billable unit = 0.5 mg
- J1449 Injection, eflapegrastim-xnst, 0.1 mg; 1 billable unit = 0.1 mg (Rolvedon only)
- J3590 Unclassified biologics (Ryzneuta only)

#### NDC(s):

- Neulasta 6 mg single-dose prefilled syringe: 55513-0190-xx
- Neulasta 6 mg single-dose prefilled syringe Onpro Kit: 55513-0192-xx
- Fulphila 6 mg single-dose prefilled syringe: 83257-0005-xx
- Udenyca 6 mg single-dose prefilled syringe: 70114-0101-xx
- Udenyca 6 mg single-dose prefilled autoinjector: 70114-0120-xx
- Udenyca 6 mg single-dose prefilled syringe ONBODY kit: 70114-0130-xx
- Ziextenzo 6 mg single-dose prefilled syringe: 61314-0866-xx
- Nyvepria 6 mg single-dose prefilled syringe: 00069-0324-xx
- Fylnetra 6 mg single-dose prefilled syringe: 70121-1627-xx
- Stimufend 6 mg single-dose prefilled syringe: 65219-0371-xx
- Rolvedon 13.2 mg single-dose prefilled syringe: 76961-0101-xx
- Ryzneuta 20 mg/mL prefilled syringe: 73491-0627-xx

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#### **Appendix 1 – Covered Diagnosis Codes**

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#### Neulasta, Fulphila, Udenyca, Ziextenzo, Nyvepria, Fylnetra, & Stimufend

ICD-10	ICD-10 Description
D61.810	Antineoplastic chemotherapy induced pancytopenia
C64.1	Malignant neoplasm of right kidney, except renal pelvis
C64.2	Malignant neoplasm of left kidney, except renal pelvis
C64.9	Malignant neoplasm of unspecified kidney, except renal pelvis
D70.1	Agranulocytosis secondary to cancer chemotherapy
D70.9	Neutropenia, unspecified
T45.1X5A	Adverse effect of antineoplastic and immunosuppressive drugs initial encounter
T45.1X5D	Adverse effect of antineoplastic and immunosuppressive drugs subsequent encounter
T45.1X5S	Adverse effect of antineoplastic and immunosuppressive drugs sequela
T66.XXXA	Radiation sickness, unspecified, initial encounter
T66.XXXD	Radiation sickness, unspecified, subsequent encounter
T66.XXXS	Radiation sickness, unspecified, sequela
W88.1	Exposure to radioactive isotopes
W88.8	Exposure to other ionizing radiation
Z41.8	Encounter for other procedures for purposes other than remedying health state
Z48.290	Encounter for aftercare following bone marrow transplant
Z51.11	Encounter for antineoplastic chemotherapy
Z51.12	Encounter for antineoplastic immunotherapy

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ICD-10	ICD-10 Description
Z51.89	Encounter for other specified aftercare
Z52.011	Autologous donor, stem cells
Z52.091	Other blood donor, stem cells
Z76.89	Persons encountering health services in other specified circumstances
Z94.81	Bone marrow transplant status
Z94.84	Stem cells transplant status

#### Rolvedon & Ryzneuta

ICD-10	ICD-10 Description
D61.810	Antineoplastic chemotherapy induced pancytopenia
D61.811	Other drug-induced pancytopenia
D61.818	Other pancytopenia
D70.1	Agranulocytosis secondary to cancer chemotherapy
D70.9	Neutropenia, unspecified
T45.1X5A	Adverse effect of antineoplastic and immunosuppressive drugs initial encounter
T45.1X5D	Adverse effect of antineoplastic and immunosuppressive drugs subsequent encounter
T45.1X5S	Adverse effect of antineoplastic and immunosuppressive drugs sequela
T66.XXXA	Radiation sickness, unspecified, initial encounter
T66.XXXD	Radiation sickness, unspecified, subsequent encounter
T66.XXXS	Radiation sickness, unspecified, sequela
W88.1	Exposure to radioactive isotopes
W88.8	Exposure to other ionizing radiation
Z41.8	Encounter for other procedures for purposes other than remedying health state
Z51.11	Encounter for antineoplastic chemotherapy
Z51.12	Encounter for antineoplastic immunotherapy
Z51.89	Encounter for other specified aftercare
Z76.89	Persons encountering health services in other specified circumstances

## Appendix 2 – Centers for Medicare and Medicaid Services (CMS)

The preceding information is intended for non-Medicare coverage determinations. Medicare coverage for outpatient (Part B) drugs is outlined in the Medicare Benefit Policy Manual (Pub. 100-2), Chapter 15, §50 Drugs and Biologicals. In addition, National Coverage Determinations (NCDs) and/or Local Coverage Determinations (LCDs) may exist and compliance with these policies is required where applicable. Local Coverage Articles (LCAs) may also exist for claims payment purposes or to clarify benefit eligibility under Part B for drugs which may be self-administered. The following link may be used to search for NCD, LCD, or LCA documents: <a href="https://www.cms.gov/medicare-coverage-">https://www.cms.gov/medicare-coverage-</a>

	Long-Acting Granulocyte Colony Stimulating Factors (LA-gCSF) (NEULASTA®; FULPHILA™; UDENYCA®; ZIEXTENZO®; NYVEPRIA™; FYLNETRA®; STIMUFEND®; ROLVEDON®; RYZNEUTA®)	Ν
Page 11	Prior Auth Criteria	1
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	without approval.	
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<u>database/search.aspx</u>. Additional indications, including any preceding information, may be applied at the discretion of the health plan.

Medicare Part B Covered Diagnosis Codes			
Jurisdiction	NCD/LCA/LCD	Contractor	
	Document (s)		
J, M	A56748	Palmetto GBA	
J, M	A54682	Palmetto GBA	

Medicare Part B Administrative Contractor (MAC) Jurisdictions			
Jurisdiction	Applicable State/US Territory	Contractor	
E (1)	CA, HI, NV, AS, GU, CNMI	Noridian Healthcare Solutions, LLC	
F (2 & 3)	AK, WA, OR, ID, ND, SD, MT, WY, UT, AZ	Noridian Healthcare Solutions, LLC	
5	KS, NE, IA, MO	Wisconsin Physicians Service Insurance Corp (WPS)	
6	MN, WI, IL	National Government Services, Inc. (NGS)	
H (4 & 7)	LA, AR, MS, TX, OK, CO, NM	Novitas Solutions, Inc.	
8	MI, IN	Wisconsin Physicians Service Insurance Corp (WPS)	
N (9)	FL, PR, VI	First Coast Service Options, Inc.	
J (10)	TN, GA, AL	Palmetto GBA	
M (11)	NC, SC, WV, VA (excluding below)	Palmetto GBA	
L (12)	DE, MD, PA, NJ, DC (includes Arlington & Fairfax counties and the city of Alexandria in VA)	Novitas Solutions, Inc.	
K (13 & 14)	NY, CT, MA, RI, VT, ME, NH	National Government Services, Inc. (NGS)	
15	KY, OH	CGS Administrators, LLC	



# Nondiscrimination & Language Access Policy



Discrimination is Against the Law. Aspirus Health Plan, Inc. complies with applicable Federal civil rights laws and does not discriminate on the basis of race, color, national origin, age, disability, or sex, (including sex characteristics, including intersex traits; pregnancy or related conditions; sexual orientation, gender identity and sex stereotypes), consistent with the scope of sex discrimination described at 45 CFR § 92.101(a)(2). Aspirus Health Plan, Inc. does not exclude people or treat them less favorably because of race, color, national origin, age, disability, or sex.

Aspirus Health Plan, Inc.:

Provides people with disabilities reasonable modifications and free appropriate auxiliary aids and services to communicate effectively with us, such as:

- Qualified sign language interpreters.

- Written information in other formats (large print, audio, accessible electronic formats, other formats).

Provides free language assistance services to people whose primary language is not English, which may include:

- Qualified interpreters.
- Information written in other languages.

If *you* need reasonable modifications, appropriate auxiliary aids and services, or language assistance services, contact the Nondiscrimination Grievance Coordinator at the address, phone number, fax number, or email address below.

If you believe that Aspirus Health Plan, Inc. has failed to provide these services or discriminated in another way on the basis of race, color, national origin, age, disability, or sex, you can file a grievance with:

Nondiscrimination Grievance Coordinator Aspirus Health Plan, Inc. PO Box 1890 Southampton, PA 18966-9998 Phone: 1-866-631-5404 (TTY: 711) Fax: 763-847-4010 Email: customerservice@aspirushealthplan.com

You can file a grievance in person or by mail, fax, or email. If you need help filing a grievance, the Nondiscrimination Grievance Coordinator is available to help you.

*You* can also file a civil rights complaint with the U.S. Department of Health and Human Services, Office for Civil Rights, electronically through the Office for Civil Rights Complaint Portal, available at https://ocrportal.hhs.gov/ocr/portal/lobby.jsf, or by mail or phone at:

U.S. Department of Health and Human Services 200 Independence Avenue, SW Room 509F, HHH Building Washington, D.C. 20201 1.800.368.1019, 800.537.7697 (TDD)

Complaint forms are available at http://www.hhs.gov/ocr/office/file/index.html. This notice is available at Aspirus Health Plan, Inc.'s website: https://aspirushealthplan.com/webdocs/70021-AHP-NonDiscrim\_Lang-Assist-Notice.pdf.

#### Language Assistance Services

Albanian: KUJDES: Nëse flitni shqip, për ju ka në dispozicion shërbime të asistencës gjuhësore, pa pagesë. Telefononi në 1-800-332-6501 (TTY: 711). (711: (تق هاتف الصم والبك) 1-800-332-6501 تنبيه : إذا كنت تتحدث اللغة العربية، فإن خدمات المساعدة اللغوية متاحة لك مجاناً التصل بن اعلى رقم الهاتف ال-800-332-6501 (TTY: 711). French: ATTENTION: Si vous parlez français, des services d'aide linguistique vous sont proposés gratuitement. Appelez le 1-800-332-6501 (ATS: 711). German: ACHTUNG: Wenn Sie Deutsch sprechen, stehen Ihnen kostenlos sprachliche Hilfsdienstleistungen zurVerfügung. Rufnummer: 1-800-332-6501 (TTY: 711).

Hindi: \_यान द\_: य\_द आप िहंदी बोलते ह\_ तो आपके िलए मु\_त म\_ भाषा सहायता सेवाएं उपल\_ध ह\_11-800-332-6501 (TTY: 711) पर कॉल कर\_। Hmong: LUS CEEV: Yog tias koj hais lus Hmoob, cov kev pab txog lus, muaj kev pab dawb rau koj. Hu rau 1-800-332-6501 (TTY: 711).

Korean: 주의: 한국어를 사용하시는 경우, 언어 지원 서비스를 무료로 이용하실 수 있습니다.1-800-332-6501 (TTY: 711)번으로 전화해 주십시오.

Polish: UWAGA: Jeżeli mówisz po polsku, możesz skorzystać z bezpłatnej pomocy językowej. Zadzwoń pod numer1-800-332-6501 (TTY: 711). Russian: ВНИМАНИЕ: Если вы говорите на русском языке, то вам доступны бесплатные услуги перевода.Звоните 1-800-332-6501 (телетайп: 711).

Spanish: ATENCIÓN: si habla español, tiene a su disposición servicios gratuitos de asistencia lingüística. Llame all-800-332-6501 (TTY: 711). Tagalog: PAUNAWA: Kung nagsasalita ka ng Tagalog, maaari kang gumamit ng mga serbisyo ng tulong sa wika nangwalang bayad. Tumawag sa 1-800-332-6501 (TTY: 711).

Traditional Chinese: 注意: 如果您使用繁體中文, 您可以免費獲得語言援助服務。請 致電 1-800-332-6501 (TTY: 711)

Vietnamese: CHÚ Ý: Nếu bạn nói Tiếng Việt, có các dịch vụ hỗ trợ ngôn ngữ miễn phí dành cho bạn. Gọi số 1-800-332-6501 (TTY: 711). Pennsylvania Dutch: Wann du Deitsch (Pennsylvania German / Dutch) schwetzscht, kannscht du mitaus Koschte ebbergricke, ass dihr helft mit die englisch Schprooch. Ruf selli Nummer uff: Call 1-800-332-6501 (TTY: 711).

Lao: ໂປດຊາບ: ຖ້າວ່າ ທ່ານເວົ້າພາສາ ລາວ, ການບໍລິການຊ່ວຍເຫຼືອດ້ານພາສາ,ໂດຍບໍ່ເສັງຄ່າ, ແມ່ນມີພ້ອມໃຫ້ທ່ານ. ໂທຣ 1-800-332-6501 (TTY: 711).