The regulatory approval of anamorelin for treatment of cachexia in patients with non-small cell lung cancer, gastric cancer, pancreatic cancer, and colorectal cancer in Japan: facts and numbers.

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Abstract

Anamorelin is a ghrelin receptor agonist that can be administered orally and thought to improve cancer cachexia by improving appetite and increasing serum insulin-like growth factor-1. Anamorelin was not approved for use in Europe. In contrast, the use of anamorelin for cancer cachexia in four types of cancer (non-small cell lung cancer, gastric cancer, pancreatic cancer, and colorectal cancer) was approved in Japan on 11 December 2020. Phase 2 trial (ONO-7643-04) for the treatment of patients with non-small cell lung cancer and cachexia resulted in 1.56 kg lean body mass increase assessed by dual-energy X-ray absorptiometry (DXA). Another study for advanced and unresectable gastrointestinal (colorectal, gastric, or pancreatic) cancer showed 1.89 ± 0.36 kg improvement in lean body mass. Skeletal lean body mass assessed by DXA is important for diagnosing sarcopenia and cachexia in Asia. The approval of anamorelin is expected to change clinical practice of cancer cachexia in Japan and hopefully in other countries. In the past, cachexia was rarely diagnosed in Japan, because it was often thought that cachexia meant terminal stage. The dissemination of clinical findings on anamorelin from Japan, as well as the creation of consensus papers and clinical practice guidelines for cachexia in Japan and Asia, will be required to promote international expansion in the future.

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Pharmacotherapy for cachexia is limited. The Food and Drug Administration approved human growth hormone for people with acquired immunodeficiency syndrome-related cachexia more than 20 years ago in the USA.1 Megestrol acetate can be used for anorexia and cachexia in patients with cancer or acquired immunodeficiency syndrome in Korea. However, growth hormone and megestrol acetate were not approved in Japan for cachexia. The American Society of Clinical Oncology guidelines for cancer cachexia published in 2020 recommend drug intervention only by progesterone analogues and corticosteroids for appetite improvement and weight gain.2 Corticosteroids are recommended only in a short term at a dose of 3–4 mg of dexamethasone per day. However, other drugs including anamorelin are not included as a treatment option in the current recommendations.2

Anamorelin is a ghrelin receptor agonist that can be administered orally.3 Ghrelin is a peptide secreted mainly by the stomach. It is an endogenous agonist of the growth hormone receptor type 1a, which regulates energy metabolism in vivo by promoting growth hormone secretion, increasing appetite, and promoting lipogenesis.4 Anamorelin is a selective oral ghrelin-like agonist that acts on ghrelin receptors in the stomach. Anamorelin is thought to improve cancer
Anamorelin was not approved for use in Europe. The Committee for Medicinal Products for Human Use concluded that the ROMANA studies7,8 showed a marginal effect of Adlumiz (anamorelin) on lean body mass and no proven effect on hand grip strength or patients’ quality of life in 2017.9 Moreover, the safety data of the drug had not been recorded adequately.9

In Japan, the Pharmaceuticals and Medical Devices Agency is a regulatory agency, working together with the Ministry of Health, Labor and Welfare for drug approval. They conduct scientific reviews of application of medical devices as well as pharmaceuticals and are responsible for monitoring their post-marketing safety. They set a target for the total review time for new drugs at a median of 12 months for standard review products.

The use of anamorelin for cancer cachexia was approved in Japan for the first time in the world on 11 December 2020. An application was filed in Japan for manufacturing and marketing approval of anamorelin with the planned indication of ‘improvement of weight loss and anorexia in cancer cachexia’ in November 2018. The two clinical studies on which the application was based are as follows. Phase 2 trial (ONO-7643-04) for the treatment of patients with non-small cell lung cancer and cachexia resulted in 1.56 kg lean body mass increase assessed by dual-energy X-ray absorptiometry (DXA).5 Changes in lean body mass, body weight, and anorexia symptoms showed significant differences in the treatment arm compared with the placebo arm; however, handgrip strength or 6-min walk test did not show significant differences.5 Another study for advanced and unresectable gastrointestinal (colorectal, gastric, or pancreatic) cancer showed 1.89 ± 0.36 kg improvement in lean body mass assessed by DXA.6 However, functional endpoints such as muscle strength and physical function were not evaluated in this study.5 Most studies showed a significant increase in lean body mass assessed by DXA (Table 1).10

It was deliberated by the Ministry of Health, Labor and Welfare, Pharmaceutical Affairs and Food Sanitation Council, First Committee on Drugs, and decided to continue deliberation on August 2019. The reason was that there were doubts about the efficacy, safety, and the way information should be provided to medical institutions and patients.

In December 2020, the second round of deliberation was held, and the approval was granted for the indication of cancer cachexia in four types of cancer: non-small cell lung cancer, gastric cancer, pancreatic cancer, and colorectal cancer.11,12 Anamorelin treatment increased lean body mass assessed by DXA and body weight.5,6 Assessment of lean body mass by DXA is not recommended for sarcopenia diagnosis in the Sarcopenia Definition and Outcomes Consortium position statements13 but is included in the Asian Working Group for Sarcopenia 2019 consensus.14 Skeletal lean body mass assessed by DXA is important for diagnosing sarcopenia and cachexia in Asia. It is expected to be officially approved by the end of 2020 at the earliest and by the end of January 2021 at the latest.

Specific dosage and administration, precautions, warnings, contraindications, and post-marketing surveillance are planned as follows.14 The usual adult dosage of anamorelin hydrochloride is 100 mg orally once daily on an empty stomach. However, there are several precautions. If no effect is observed after 3 weeks from the start of administration, administration should be discontinued. There is no experience of administration for longer than 12 weeks. Moreover, there are several warnings. Anamorelin should be used under the supervision of a physician with sufficient knowledge and experience in cancer cachexia. The risks and benefits of administering the drug should be fully explained and made sure that the patient or family understands before administering the drug. As an example of the risks associated with the drug, cardiac risks such as QT interval prolongation may occur. In addition, the drug is contraindicated in patients with a history of hypersensitivity to any of its ingredients or in patients taking concomitant medications. Post-marketing surveillance is mandatory. The re-examination period is 8 years.

The approval of the use of anamorelin is expected to change clinical practice of cancer cachexia in Japan and hopefully in other countries. In the past, cachexia was rarely diagnosed in Japan, because it was often thought that cachexia meant terminal stage. Only 17.4% of health care professionals answered that they were evaluated for cachexia.15 However, for cachexia due to non-small cell

### Table 1 Effects of anamorelin on lean body mass

|                | Anamorelin Mean (kg) | Anamorelin SD (kg) | Placebo Mean (kg) | Placebo SD (kg) | P value |
|----------------|----------------------|--------------------|-------------------|-----------------|---------|
| Katakami et al.4 | 1.38                 | 0.18               | –0.17             | 0.17            | <0.001  |
| Hamauchi et al.6 | 1.89                 | 0.36               | 0.55              | 0.29            | 0.0516  |
| Takayama et al.10| 1.15                 | 0.31               | –0.47             | –1.00 to 0.21   | <0.001  |
| ROMANA 17       | 0.99                 | 0.61 to 1.36       | –0.98             | –1.49 to 0.41   | <0.001  |
| ROMANA 27       | 0.65                 | 0.38 to 0.91       |                   |                 |         |

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15. Katakami et al, Takayama et al.
lungenkrebs, gastrointestinal Krebs, kolorektale Krebs, und kolorektale Krebs, Cachexie ist mehr als nur diagnostiziert und Anamorelin als Gewichtsverlust bei 5% oder mehr ist beobachtet in 6 Monaten. Ferner, es ist notwendig, die Erhöhung der Bewusstheit von Krebs Cachexia und zu studieren, ob die Kombination von Therapie mit Ernährungstherapie, Übungs- und Rehabilitation Aufmerksamkeit auf Cachexia zu vergrößern und Krebs, Cachexia wird möglicherweise diagnostiziert und Lungenkrebs, Magenkrebs, Pankreaskrebs, und kolorektale Krebs. Weiterhin, in Europa und in Asien, wie Korea und Taiwan, Anamorelin wird möglicherweise als Wohlbefinden und Appetit und Gewicht durch Behandlungen von multimodalen Interventionen. In Asien, wie Korea und Taiwan, Anamorelin wird möglicherweise als Cachexie behandelt. In der anderen Hand, in Europa und den USA, approbierung ist schwierig, um keine Muskel starke, physische Funktion und Prognose zu verbessern. Die Verbreitung von klinischen Ergebnissen auf Anamorelin Japan und Asien, als die Erstellung von Konsensuspapieren und klinische Praxisrichtlinien für Cachexia in Japan und Asien, wird benötigt, um internationale Promotion zu erweitern.

**References**

1. Gilden D. Human growth hormone available for AIDS wasting. *Ghmc Treat Issues* 1995;9:9–11.
2. Roeland EJ, Bohlke K, Baracos VE, Bruera E, Del Fabbro E, Dixon S, et al. Management of cancer cachexia: ASCO guideline. *J Clin Oncol* 2020;38:2438–2453.
3. Anker SD, Coats AJ, Morley JE. Evidence for partial pharmacological reversal of the cancer anorexia-cachexia syndrome: the case of anamorelin. *J Cachexia Sarcopenia Muscle* 2015;6:275–277.
4. Kojima M, Hosoda H, Date Y, Nakazato M, Matsuo H, Kangawa K. Ghrelin is a growth hormone releasing acylated peptide from stomach. *Nature* 1999;402:656–660.
5. Katakami N, Uchino J, Yokoyama T, Naito T, Kondo M, Yamada K, et al. Anamorelin (ONO-7643) for the treatment of patients with non-small cell lung cancer and cachexia: results from a randomized, double-blind, placebo-controlled, multicenter study of Japanese patients (ONO-7643-04). *Cancer* 2018;124:606–616.
6. Hamauchi S, Furuse J, Takano T, Munemoto Y, Furuya K, Baba H, et al. A multicenter, open-label, single-arm study of anamorelin (ONO-7643) in advanced gastrointestinal cancer patients with cancer cachexia. *Cancer* 2019;125:4294–4302.
7. Temel JS, Abernethy AP, Currow DC, Friend J, Duus EM, Yan Y, et al. Anamorelin in patients with non-small-cell lung cancer and cachexia (ROMANA 1 and ROMANA 2): results from two randomised, double-blind, phase 3 trials. *Lancet Oncol* 2016;17:519–531.
8. Currow D, Temel JS, Abernethy A, Milanowski J, Friend J, Fearon KC. ROMANA 3: a phase 3 safety extension study of anamorelin in advanced non-small-cell lung cancer (NSCLC) patients with cachexia. *Ann Oncol* 2017;28:1949–1956.
9. Refusal of the marketing authorisation for adiumiz (anamorelin hydrochloride). EMA/305706/2017 rev1 (2017) EMEA/H/C/003847.
10. Takayama K, Katakami N, Yokoyama T, Atagi S, Yoshimori K, Kagamu H, et al. Anamorelin (ONO-7643) in Japanese patients with non-small cell lung cancer and cachexia: results of a randomized phase 2 trial. *Support Care Cancer* 2016;24:3495–3505.
11. Pharmaceutical Affairs and Food Sanitation Council, First Committee on Drugs, Japan’s first cancer cachexia drug Edlumiz approved after second round of deliberations. Mix Online. https://www.mixonline.jp/tabid55.5html?artid=70308 Accessed 17 Dec 2020. [In Japanese]
12. Pharmaceutical Affairs and Food Sanitation Council, First Committee on Drugs. The Ministry of Health, Labor and Welfare. https://www.mhlw.go.jp/stf/shingi/shingi-yakuji_127851.html Accessed 17 Dec 2020. [In Japanese]
13. Bhasin S, Travison TG, Manini TM, Patel S, Pencina KM, Fielding RA, et al. Sarcopenia definition: the position statements of the Sarcopenia Definition and Outcomes Consortium. *J Am Geriatr Soc* 2020;68:1410–1418.
14. Chen LK, Woo J, Assantachai P, Auyeung TW, Chou MY, Iijima K, et al. Asian Working Group for Sarcopenia: 2019 consensus update on sarcopenia diagnosis and treatment. *J Am Med Dir Assoc* 2020;21:300–307, e2.
15. Nakahara S, Wakabayashi H, Maeda K, Nishioka S, Kokura Y. Sarcopenia and cachexia evaluation in different healthcare settings: a questionnaire survey of health professionals. *Asia Pac J Clin Nutr* 2018;27:167–175.
16. Prado CM, Purcell SA, Laviano A. Nutrition interventions to treat low muscle mass in cancer. *J Cachexia Sarcopenia Muscle* 2020;11:366–380.
17. Wakabayashi H, Sakuma K. Rehabilitation nutrition for sarcopenia with disability: A combination of both rehabilitation and nutrition care management. *J Cachexia Sarcopenia Muscle* 2014;5:269–277.
18. McKeaveney C, Maxwell P, Noble H, Reid J. A critical review of multimodal interventions for cachexia. *Adv Nutr* 2020;https://doi.org/10.1093/advances/nmaa111.
19. Fearon K, Argiles JM, Baracos VE, Bernabei R, Coats A, Crawford J, et al. Request for regulatory guidance for cancer cachexia intervention trials. *J Cachexia Sarcopenia Muscle* 2015;6:272–274.
20. von Haehling S, Morley JE, Coats AJS, Anker SD. Ethical guidelines for publishing in the Journal of Cachexia, Sarcopenia and Muscle. *J Cachexia Sarcopenia Muscle* 2019;10:1143–1145.