First-Ever Redosable Gene Therapy, VYJUVEK™ (beremagene geperpavec) for the Treatment of Dystrophic Epidermolysis Bullosa

Vyjuvek[™], also known as B-VEC, a new treatment for dystrophic epidermolysis bullosa (DEB), is a step closer to being made available for DEB patients in the UK with the recent news that the US Food and Drug Administration (FDA) has approved it for treatment of DEB in patients aged 6 months and older in the US. Once available, Vyjuvek[™] will be the first-ever redosable gene therapy for treatment of DEB.

On May 19, 2023, the US FDA approved Vyjuvek™ (beremagene geperpavec), the first gene therapy for the treatment of wounds in patients with dystrophic epidermolysis bullosa (DEB). DEB is a rare genetic disorder that is hallmarked by extremely fragile skin that rips and blisters with even the slightest friction, leading to open wounds that are prone to skin infections and fibrosis. The disease is caused by mutations in the *COL7A1* gene which leads to a deficiency in collagen 7 (COL7), an essential structural protein that helps strengthen and stabilize the outer and middle layers of the skin. When COL7A1 is deficient, skin layers can separate, causing painful and debilitating blisters and wounds. Symptoms are usually present at birth and can vary widely. In mild cases, blistering is primarily found on the hands, feet, knees, and elbows. Patients with more severe disease may experience a variety of serious medical conditions including painful and debilitating widespread blistering that can lead to vision loss and fusion of the fingers and toes. Blisters in the lining of the mouth and digestive tract often result in poor growth and nutrition as well as anemia. With the severe form of DEB, the lifetime risk of developing aggressive skin cancer is higher than 90%. According to the United States National Epidermolysis Bullosa Registry, the incidence and prevalence of dystrophic epidermolysis bullosa is between 1.35 and 3.05 cases per 1 million live births.

Vyjuvek is a non-invasive, topical, redosable gene therapy designed to address the genetic cause of DEB. It uses a modified and harmless version of the herpes simplex virus to deliver two healthy copies of the *COL7A1* gene directly to the skin cells in the wounds. The new gene allows skin cells to produce functional COL7 protein, which increases the skin's structural integrity, helping to reduce blistering and repair wounds. Vyjuvek is available in a biological suspension, which is mixed with a non-active ingredient gel before being applied to the skin. The dosage can be measured in plaque-forming units (PFU, the number of individual virus particles) or in the volume applied (in milliliters). The gel is applied once per week as droplets over the wound, with each droplet spaced about a centimeter apart from other droplets. The therapy may be applied in a healthcare setting or at home, but the application should always be performed by a healthcare professional. The recommended dose depends on the size of the wound. For patients younger than age 3, the maximum weekly dose should not exceed 1.6 billion PFU (0.8 mL). For patients ages 3 and older, the maximum weekly dose is 3.2 billion PFU (1.6 mL). It may not be possible to treat all wounds in a single patient's visit. It is recommended that individual wounds be selected and treated until they are closed before treatment begins on other wounds. If a previously healed wound re-opens, it should be prioritized for treatment.

The FDA approval of Vyjuvek was based on two clinical studies, GEM-1 and GEM-3. The GEM-1 trial tested the safety and efficacy of Vyjuvek in six adults and three children who had a more severe form of DEB. In all participants, one wound was treated with Vyjuvek, while a different wound with a corresponding size was administered a placebo. The wounds that were given placebo showed varying amounts of healing or worsening. In comparison, all but one of the wounds treated with Vyjuvek closed completely and remained healed for at least three months. Complete wound closure was defined as ≥90% reduction from baseline wound surface area. The GEM-3 study enrolled 31 people, most with the more severe form of DEB. As

with GEM-1, each participant had one wound treated with Vyjuvek, while another wound was administered the placebo. The results showed that, after three months of treatment, significantly more wounds treated with Vyjuvek than placebo had healed (68% vs. 23%). A similar difference was seen after six months of treatment (65% vs. 26%), meeting the study's main objective. In both studies, Vyjuvek was generally safe and well tolerated with no reported serious treatment-related adverse events.

Current treatment for DEB has been largely supportive. It includes wound care, control of infection, along with prevention strategies, and treatment of any complications. As a result, many patients suffer from constant pain, discomfort, poor nutrition, and disfigurement. Vyjuvek appears to offer people living with DEB the opportunity for an improved quality of life. Because Vyjuvek works to correct the underlying skin defect of DEB, the therapy appears to not only close existing wounds but also prevent skin from reblistering and reopening. The durability of the wound closure remains a concern, but to date, data has initially been positive.

Outside of the US, the European Medicines Agency (EMA) has granted Vyjuvek orphan drug designation and PRIME (PRIority Medicines) eligibility for DEB. Commencement of starting the Marketing Authorization Application procedure is anticipated in the second half of 2023 with a potential approval in 2024.

References

- https://www.fda.gov/news-events/press-announcements/fda-approves-first-topical-genetherapy-treatment-wounds-patients-dystrophic-epidermolysisbullosa?utm_campaign=Weekly%20Drug%20Update&utm_medium=email&_hsmi=259899895&_ hsenc=p2ANqtz-_l3l-IdmYh6H2AdLuQLlWo8ZtN8VPtO0UBTizBYL_catptMRajguzWpF9nvDkJ4XViFE31vun1lvY6v5x7e8 ZEmZqP48h01fJETFPK2NXVlsyt6k0&utm_content=259877426&utm_source=hs_email
- 2. https://ir.krystalbio.com/news-releases/news-release-details/krystal-biotech-receives-fda-approval-first-ever-redosable-gene
- 3. https://www.pharmaceutical-technology.com/news/fda-krystal-biotech-vyjuvek/
- 4. https://www.fiercepharma.com/pharma/first-redoseable-gene-therapy-approved-treat-rare-fragile-skin-disease-krystal-biotechs
- 5. https://www.biopharmadive.com/news/krystal-fda-approval-gene-therapy-epidermolysis-bullosa/650844/
- 6. https://epidermolysisbullosanews.com/kb103/
- 7. https://ir.krystalbio.com/news-releases/news-release-details/krystal-biotech-receives-fda-approval-first-ever-redosable-gene
- 8. https://www.ncbi.nlm.nih.gov/books/NBK1304/
- 9. https://med.stanford.edu/dermatology/resources/gsdc/eb_clinic/eb-faqs.html
- 10. https://emedicine.medscape.com/article/1062939-overview#a6
- 11. https://www.europeanpharmaceuticalreview.com/news/182795/first-redosable-gene-therapy-approved/