Molnupiravir: First Approval

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Abstract
Molnupiravir (Lagevrio®) is an orally-administered antiviral prodrug that inhibits replication of RNA viruses through viral error induction. It is being developed by Merck and Ridgeback Biotherapeutics for the prevention and treatment of Coronavirus disease 2019 (COVID-19). Molnupiravir received its first approval on 4 November 2021 in the UK for the treatment of mild to moderate COVID-19 in adults with a positive severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) diagnostic test and who have at least one risk factor for developing severe illness. Molnupiravir is filed for approval and has emergency use authorization for the treatment of COVID-19 in several countries, including the USA, Japan and those in the EU. This article summarizes the milestones in the development of molnupiravir leading to this first approval for COVID-19.

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Molnupiravir (Lagevrio®): Key Points
- A broad-spectrum RNA antiviral prodrug being developed by Merck and Ridgeback Biotherapeutics for the treatment of COVID-19
- Received its first approval on 4 November 2021 in the UK
- Approved for use in adults with mild to moderate COVID-19 at risk of progression to severe illness

1 Introduction
Molnupiravir (Lagevrio®) is an orally-administered, small-molecule, antiviral prodrug that inhibits replication of RNA viruses through viral error induction [1]. The drug was invented by Drug Innovation Ventures at Emory (DRIVE; wholly owned by Emory University) and is being developed by Merck, in collaboration with Ridgeback Biotherapeutics, for the prevention and treatment of Coronavirus disease 2019 (COVID-19) [1, 2]. Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), the causative virus of COVID-19, uses an RNA-dependent RNA polymerase (RdRp) complex for replication [3]. Therefore, RdRp is a promising target for both developing new antiviral drugs against SARS-CoV-2 as well as for repurposing existing ones [4]. A number of drug candidates with activity against SARS-CoV-2 RdRp are being evaluated for the treatment of COVID-19 [4, 5]. Remdesivir, an intravenously administered RdRp inhibitor, is approved for the treatment of adults and paediatric patients with COVID-19 [6]. However, new treatment options are required to prevent disease progression in at-risk populations; in addition to improving patient-level clinical outcomes, such treatments can reduce healthcare system burden. In treating COVID-19, oral pills are preferable to intravenous drugs in terms of patient convenience, use during self-isolation, faster treatment uptake, healthcare resource use, scalability, logistics and cost.

Molnupiravir received its first approval on 4 November 2021 in the UK for the treatment of mild-to-moderate COVID-19 in adults with a positive SARS-CoV-2 diagnostic test and who have at least one risk factor for developing severe illness [7, 8]. The recommended dosage of molnupiravir is 800 mg every 12 h for 5 days. It should be administered as soon as possible after a COVID-19 diagnosis and within 5 days of symptom onset [7]. Molnupiravir underwent an accelerated first-in-human clinical trial programme through
extraordinary collaboration between the drug developers, contract research organization and regulatory authorities [2].

Molnupiravir has been filed for approval and has emergency use authorization for the treatment of COVID-19 in several countries, including the USA, Japan and those in the EU. In addition, a multinational phase III trial is evaluating molnupiravir for post-exposure prophylaxis against COVID-19.

1.1 Company Agreements

In May 2020, Ridgeback Biotherapeutics entered into a licensing agreement with Merck for the development of molnupiravir. Under the terms of agreement, Merck has acquired exclusive worldwide rights to develop and commercialise molnupiravir and related molecules [9, 10]. Ridgeback Biotherapeutics acquired an exclusive license to DRIVE’s molnupiravir in March 2020 [11].

In June 2021, Cipla, Dr Reddys Laboratories, Emcure Pharmaceuticals Limited, Sun Pharmaceutical Industries and Torrent Pharmaceuticals entered into a collaboration agreement to conduct a clinical trial of molnupiravir in the treatment of mild COVID-19 in an outpatient setting in India. In March and April 2021, the five companies had individually entered into a non-exclusive voluntary licensing agreement with Merck to manufacture and supply molnupiravir to India and over 100 low- and middle-income countries [12]. As of January 2022, Merck has signed non-exclusive licensing agreements with additional generic drug manufacturers in India.

In October 2021, Merck granted a license to the Medicines Patent Pool to increase broad access to molnupiravir in low- and middle-income countries upon regulatory approvals. Under this agreement, Merck, Ridgeback Biotherapeutics and Emory University will not receive royalties for molnupiravir sales as long as COVID-19 remains classified as a Public Health Emergency of International Concern by the World Health Organization [13].

In January 2022, Merck and Ridgeback Biotherapeutics entered into a long-term supply agreement with UNICEF, under which Merck will allocate up to 3 million courses of molnupiravir to UNICEF during the first half of 2022 for distribution in more than 100 low- and middle-income countries upon local regulatory authorizations [14]. Merck also has a procurement agreement with the US Government, under which the US Government purchased a total of approximately 3.1 million courses of molnupiravir upon emergency use authorization by the US FDA [14].

2 Scientific Summary

2.1 Pharmacodynamics

Molnupiravir is an orally available prodrug that is rapidly converted in plasma by host’s esterases to the ribonucleoside analogue N-hydroxycytidine (NHC) [15]. After intracellular uptake, NHC is phosphorylated into its pharmacologically active ribonucleoside triphosphate
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NHC-TP is incorporated into viral RNA by the RdRp, resulting in viral error induction (i.e. accumulation of errors in the viral genome ultimately leading to inhibition of replication) [7, 16, 17]. Consistent with this mechanism of action, molnupiravir treatment was associated with numerically higher SARS-CoV-2 viral RNA error rates versus placebo in patients with COVID-19 in clinical trials [18–20].

Molnupiravir inhibited SARS-CoV-2 in cell culture assays, with 50% effective concentrations (EC₅₀) ranging between 0.32 and 2.66 µM [7]. Molnupiravir has broad-spectrum in vitro activity against SARS-CoV-2 variants of concern such as B.1.1.529 (omicron) [21–26], and B.1.1.7 (alpha), B.1351 (beta), P.1 (gamma) and B.1.617.2 (delta) [7]. The EC₅₀ values were 1.86–1.95 µM [21], and 1.59, 1.77, 1.32 and 1.68 µM [7], respectively. In immunodeficient mice implanted with human lung tissue, therapeutic and prophylactic administration of molnupiravir inhibited SARS-CoV-2 replication in vivo [27]. Treatment of SARS-CoV-2-infected ferrets with molnupiravir reduced SARS-CoV-2 levels in the upper respiratory tract and completely blocked transmission of the virus to untreated contact animals [28]. Molnupiravir inhibited SARS-CoV-2 in human airway epithelial cell cultures; in SARS-CoV-2-infected mice, molnupiravir decreased viral load and body weight loss and improved pulmonary function [29].

The potential for SARS-CoV-2 resistance to molnupiravir is low [30]. In phase II trials of molnupiravir in patients with COVID-19, no amino acid substitutions in SARS-CoV-2 associated with resistance to NHC have been detected [7]. Preclinical data suggest that molnupiravir is of low risk for genotoxicity or mutagenicity in clinical use [31–33].

### 2.2 Pharmacokinetics

The pharmacokinetics of molnupiravir have been evaluated in healthy volunteers [15] and patients with COVID-19 [19, 34]. According to a population pharmacokinetic model, plasma NHC pharmacokinetics following single and multiple twice-daily molnupiravir administrations can be adequately described by a linear two-compartment model, with sigmoid absorption and first-order elimination [35]. Molnupiravir pharmacokinetics did not differ significantly between healthy volunteers and COVID-19 patients [35]. In patients with COVID-19, NHC exhibited dose-proportional pharmacokinetics and the exposures were comparable across all evaluated demographic and clinical characteristics, including age, sex, body-mass index, body weight, ethnicity and kidney function [36].

Following oral administration of molnupiravir 800 mg every 12 h, the median time to peak plasma NHC concentrations was 1.5 h [7]. In healthy volunteers receiving a single molnupiravir 200 mg dose with a high-fat meal, NHC peak concentration was reduced by 35%, while area under the plasma concentration was not significantly affected. NHC does not bind to plasma proteins and it has an effective half-life of ≈ 3.3 h. In healthy volunteers, ≤ 3% of the administered dose is excreted as NHC in the urine [7].

### 2.3 Therapeutic Trials

#### 2.3.1 Non-hospitalized Patients

Molnupiravir was evaluated independently in outpatients with PCR-confirmed SARS-CoV-2 infection in a phase Ib/IIa trial (NCT04746183), using the AGILE platform. From the

### Features and properties of molnupiravir

| Alternative names | Lagevrio; EIDD-2801; MK-4482 |
|-------------------|-------------------------------|
| Class             | Antivirals; esters; hydroxylamines; pyrimidinones; ribonucleosides; small molecules |
| Mechanism of action | Viral replication inhibitors |
| Route of administration | Oral |
| Pharmacodynamics | Molnupiravir is hydrolysed to N-hydroxycytidine (NHC) which is phosphorylated to pharmacologically active N-hydroxycytidine triphosphate. Inhibits SARS-CoV-2 replication via viral error induction; robust in vitro and in vivo activity against SARS-CoV-2; retains activity against SARS-CoV-2 variants |
| Pharmacokinetics of NHC | tₘax 1.5 h; a high-fat meal reduces Cₘax by 35%; does not bind to plasma proteins; t₁/₂ 3.3 h; gender, race, age, and kidney and liver impairment have no clinically relevant effect |
| Most frequent adverse events | Diarrhoea, nausea, dizziness, headache |
| ATC codes | WHO ATC code: J05A-X (other antivirals) |
| EphMRA ATC code | J5B9 (antivirals, others) |
| Chemical name | [(2R,3S,4R,5R)-3,4-dihydroxy-5-[4-(hydroxyamino)-2-oxopyrimidin-1-yl]oxolan-2-yl]methyl 2-methylpropanoate |

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Molnupiravir significantly reduced nasopharyngeal SARS-CoV-2 load in outpatients with COVID-19 in a randomized, double-blind, placebo-controlled phase II trial (NCT04045570) [20]. Patients with confirmed SARS-CoV-2 infection and symptom onset within 7 days were randomized to molnupiravir 200, 400 or 800 mg twice daily or placebo (1:1 for 200 mg and 3:1 for other doses) for 5 days. The primary endpoint was time to SARS-CoV-2 RNA clearance (< 1,018 copies/mL), as measured by quantitative reverse transcriptase PCR of nasopharyngeal swabs. By day 28 (end of study), SARS-CoV-2 RNA clearance was seen in 91.3, 78.7 and 92.5% of patients in the molnupiravir 200, 400 and 800 mg groups, respectively, versus 80.3% in the placebo group. The median time to SARS-CoV-2 RNA clearance was significantly (p = 0.013) shorter for molnupiravir 800 mg versus placebo (14.0 vs 15.0 days), with no significant between-group differences seen for other doses. Molnupiravir also dose-dependently decreased infectious SARS-CoV-2. On day 5, the proportion of patients positive for infectious SARS-CoV-2 in nasopharyngeal swabs was significantly lower in the 400 mg (0% of 42; p = 0.034) and 800 mg (0% of 53; p = 0.027) groups than in the placebo group (11.1% of 54) [20].

Molnupiravir reduced the risk of hospitalization or death in patients with COVID-19 in a randomized, double-blind, placebo-controlled, global, phase II/III trial (MOVe-OUT; NCT04575597) [18, 37]. The trial predominantly (and exclusively in the phase III component) enrolled non-hospitalized adults with laboratory-confirmed mild to moderate COVID-19, at least one risk factor associated with poor disease outcomes and symptom onset within 5 days prior to randomization. Patients received molnupiravir 200, 400 or 800 mg in the phase II component [18] and 800 mg in the phase III component [37] or placebo twice daily for 5 days. The final dose selection for evaluation in the phase III component was based on the totality of clinical efficacy, virologic and safety outcomes across the molnupiravir clinical programme [36]. The primary endpoint of MOVe-OUT was the incidence of hospitalization for any cause or death through day 29 in the modified intent-to-treat population (NCT04575597) [18, 37]. In the phase II component (n = 302), 3.1% of patients in the combined molnupiravir group were hospitalized or died, compared with 5.4% in the placebo group [18]. Secondary endpoints of time to progression and time to sustained resolution or improvement of COVID-19 symptoms also favoured molnupiravir over placebo. At the end of treatment (day 5), molnupiravir 800 mg was associated with numerically higher rates of SARS-CoV-2 viral clearance versus placebo (12.7 vs 3.6% of patients). These results supported further evaluation of molnupiravir in the phase III component [18].

In a planned interim analysis of the phase III component of MOVe-OUT (n = 775), molnupiravir met the prespecified superiority criterion versus placebo; the primary endpoint incidence was 7.3 versus 14.1% (difference 6.8%; 95% CI − 11.3 to − 2.4; p = 0.001) [37]. Due to these positive results, the trial was stopped early based on recommendation by an independent data monitoring committee. In the final analysis (i.e. all randomized population; n = 1433), the corresponding incidence was 6.8 versus 9.7% (difference 3.0%; 95% CI – 5.9 to 0.1). A time-to-event analysis showed that molnupiravir reduced the risk of hospitalization or death by 31% versus placebo. The incidence of 29-day all-cause mortality was 0.1% in the molnupiravir group and 1.3% in the placebo group, corresponding to an 89% reduction in the risk of death. In most prespecified subgroups (based on race, age, days since symptom onset, baseline disease severity, SARS-CoV-2 nucleocapsid antibody status and qualitative assay at baseline and risk factors for severe disease), the point estimates for the difference in the primary endpoint incidence favoured molnupiravir over placebo. Consistent treatment benefit in terms of the primary endpoint was seen in subgroups based on causative SARS-CoV-2 variants of concern (delta, mu, gamma) based on data available at the time of the report. Molnupiravir also provided other clinical benefits, as assessed by the WHO Clinical Progression Scale and patient-reported COVID-19 symptoms. In exploratory analyses, based on the available data, molnupiravir reduced mean SARS-CoV-2 nasopharyngeal RNA titers from baseline at days 3 and 5 versus placebo (adjusted relative reduction 42 and 53%) [37].

2.3.2 Hospitalized Patients

Molnupiravir showed no clinical benefit in hospitalized adults with COVID-19 in a randomized, double-blind, placebo-controlled phase II trial (MOVe-IN; NCT04575584) [19]. Patients aged ≥ 18 years requiring in-hospital treatment for laboratory-confirmed COVID-19 with symptom onset within 10 days were randomized to molnupiravir 200, 400 or 800 mg, or placebo twice daily for 5 days (n = 304). The primary efficacy endpoint was sustained recovery rate, defined as the proportion of patients being alive and either not hospitalized or medically ready for hospital discharge. The sustained recovery rate was similar between molnupiravir and placebo groups (81.5–85.2% vs 84.7%) and the median time to sustained recovery was 9 days in all groups. All-cause mortality rates were also similar between the treatment groups. The lack of clinical benefit with molnupiravir in the hospitalized population was thought to be related to delayed treatment initiation in relation to the temporal pattern of COVID-19 symptom onset and illness severity [19].
### Key clinical trials of molnupiravir (Merck and/or Ridgeback biotherapeutics)

| Drug          | Indication                             | Phase | Status                     | Location(s)                        | Identifier                                      |
|---------------|----------------------------------------|-------|----------------------------|------------------------------------|------------------------------------------------|
| Molnupiravir  | COVID-19 prevention                    | III   | Recruiting                 | Multinational                      | MOVe-AHEAD; NCT04939428; 4482-013; EudraCT2021-000904-39; MK4482-013 |
| Molnupiravir  | Non-hospitalized adults with COVID-19   | II/III| Stopped early for efficacy | Multinational                      | MOVe-OUT; NCT04575597; EudraCT2020-003368-24; MK-4482-002; PHRR201209-003186; jRCT2031210148 |
| Molnupiravir  | Hospitalized adults with COVID-19       | II/III| Discontinued after phase II| Multinational                      | MOVe-IN; NCT04575584; EudraCT2020-003367-26; MK4482-001 |
| Molnupiravir  | Hospitalized adults with COVID-19       | II    | Recruiting                 | USA                                | END-COVID; NCT04405739; EIDD-2801-2004 |
| Molnupiravir  | COVID-19                               | II    | Completed                  | USA                                | NCT04405570; EIDD-2801-2003 |

### 2.4 Adverse Events

Molnupiravir was well tolerated in patients with COVID-19 in clinical trials [18, 19, 34, 37]. In a phase I trial (NCT04746183), outpatients with COVID-19 received molnupiravir 300–800 mg twice daily for 5 days plus standard-of-care or standard-of-care alone (control) [34]. A molnupiravir dose was deemed unsafe if the probability of ≥ 30% excess dose-limiting toxicity over controls was ≥ 25%. For the 800 mg dose, this probably was estimated at 0.9% [34].

In phase II trials, molnupiravir up to 800 mg twice daily for 5 days was not associated with any dose-limiting adverse events (AEs), specific safety signals or clinically relevant abnormal laboratory test results (including haematological toxicity) in non-hospitalized [18] or hospitalized [19] patients with COVID-19. The AE profile of molnupiravir was generally comparable to that of placebo [18, 19].

In the phase III component of MOVe-OUT, molnupiravir did not differ significantly from placebo with respect to the incidence of overall and treatment-related AEs [37]. At least one treatment-related AE occurred in 57 of 710 (8.0%) molnupiravir recipients and 59 of 701 (8.4%) placebo recipients; the most common (incidence ≥ 1%) of these were diarrhoea (1.7 vs 2.1%), nausea (1.4 vs 0.7%) and dizziness (1.0 vs 0.7%). No molnupiravir recipient and one placebo recipient experienced a treatment-related serious AE. Four molnupiravir and three placebo recipients discontinued study treatments because of treatment-related AEs. Two molnupiravir recipients and 12 placebo recipients died from AEs, none related to study treatment [37].

### 2.5 Ongoing Clinical Trials

The AGILE platform trial (NCT04746183) is still ongoing. The efficacy and safety of molnupiravir is being tested in a UK-based real-world study. A randomized, double-blind, placebo-controlled, multinational phase III trial (MOVe-AHEAD; NCT04939428) is evaluating molnupiravir for preventing the spread of COVID-19 within households. A randomized, double-blind, placebo-controlled, multicentre phase II trial (END-COVID; NCT04405739) is evaluating the efficacy and safety of molnupiravir in newly hospitalized adults with PCR-confirmed COVID-19.

### 3 Current Status

Molnupiravir received its first approval on 4 November 2021 in the UK for the treatment of mild-to-moderate COVID-19 in adults with a positive SARS-CoV-2 diagnostic test and who have at least one risk factor for developing severe illness [7].

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