



がん分野におけるアッヴィについて

アッヴィでは、複数の血液がんの標準治療の変革に取り組むとともに、多様ながん種に対する治験薬の開発を積極的に推進しています。献身的で経験豊富な当社のチームは、革新的なパートナーと協力し、画期的新薬となり得る製品の開発促進に努めています。当社は、世界で最も罹患者が多く、また最も消耗性が高いがん種に対し、20種類を超える治験薬を300件超の臨床試験で評価しています。当社の事業の目的は、人々の人生を豊かにすることです。そのため、患者さんが当社のがん治療薬にアクセスすることができるよう、ソリューションの探求にも取り組んでいます。詳細については、<http://www.abbvie.com/oncology> をご覧ください。

アッヴィについて

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Forward-Looking Statements

Some statements in this news release are, or may be considered, forward-looking statements for purposes of the Private Securities Litigation Reform Act of 1995. The words "believe," "expect," "anticipate," "project" and similar expressions, among others, generally identify forward-looking statements. AbbVie cautions that these forward-looking statements are subject to risks and uncertainties that may cause actual results to differ materially from those indicated in the forward-looking statements. Such risks and uncertainties include, but are not limited to, failure to realize the expected benefits from AbbVie's acquisition of Allergan plc ("Allergan"), failure to promptly and effectively integrate Allergan's businesses, competition from other products, challenges to intellectual property, difficulties inherent in the research and development process, adverse litigation or government action, changes to laws and regulations applicable to our industry and the impact of public health outbreaks, epidemics or pandemics, such as COVID-19. Additional information about the economic, competitive, governmental, technological and other factors that may affect AbbVie's operations is set forth in Item 1A, "Risk Factors," of AbbVie's 2019 Annual Report on Form 10-K, which has been filed with the Securities and Exchange Commission, as updated by its subsequent Quarterly Reports on Form 10-Q. AbbVie undertakes no obligation to release publicly any revisions to forward-looking statements as a result of subsequent events or developments, except as required by law.



1. Kater AP, et al. Abstract 125: Five-Year Analysis of Murano Study Demonstrates Enduring Undetectable Minimal Residual Disease (uMRD) in a Subset of Relapsed/Refractory Chronic Lymphocytic Leukemia (R/R CLL) Patients (Pts) Following Fixed-Duration Venetoclax-Rituximab (VenR) Therapy (Tx). Presented at the 2020 American Society of Hematology Annual Meeting & Exposition: December 5, 2020.
2. Al-Sawaf O, et al. Abstract 1310: Characteristics and Outcome of Patients with Chronic Lymphocytic Leukaemia and Partial Response to Venetoclax-Obinutuzumab. Presented at the 2020 American Society of Hematology Annual Meeting & Exposition: December 5, 2020.
3. Al-Sawaf O, et al. Abstract 127: Clonal Dynamics after Venetoclax-Obinutuzumab Therapy: Novel Insights from the Randomized, Phase 3 CLL14 Trial. Presented at the 2020 American Society of Hematology Annual Meeting & Exposition: December 5, 2020.
4. Leukemia and Lymphoma Society Minimal Residual Disease (MRD) Fact Sheet
https://www.lls.org/sites/default/files/National/USA/Pdf/Publications/FS35_MRD_Final_2019.pdf
5. Seymour JF, et al. Venetoclax-rituximab in relapsed or refractory chronic lymphocytic leukemia. *N Engl J Med.* 2018;378(12):1107-1120
6. Summary of Product Characteristics for VENCLYXTO (venetoclax). Ludwigshafen, Germany: AbbVie Deutschland GmbH & Co. KG.
7. VENCLEXTA (venetoclax) [Package Insert]. North Chicago, Ill.: AbbVie Inc.
8. Fischer K, et al. Venetoclax and Obinutuzumab in Patients with CLL and Coexisting Conditions. *N Engl J Med.* 2019;380:2225-2236.