Reducing liver disease-related deaths in the Asia-Pacific: the important role of decentralised and nonspecialist led hepatitis C treatment for cirrhotic patients

Bridget Draper,^{a,b,a,*} Win Lei Yee,^{c,2} Alisa Pedrana,^{a,b,4,3} Khin Pyone Kyi,^{e,4} Huma Qureshi,^{f,5} Hla Htay,^{c,6} Win Naing,^{e,7,8} Alexander J Thompson,^{h,i,9} Margaret Hellard,^{a,b,j,k,l,10} and Jessica Howell^{a,b,h,i,11}

Finding new ways to increase access to hepatitis C (HCV) treatment is integral to reducing liver disease-related deaths and achieving HCV elimination in the Asia-Pacific. Decentralised, non-specialist led care is essential to expand access to HCV treatment, both by increasing the number of providers and the geographical coverage. The vast majority of direct-acting antiviral (DAA) treatments for HCV do not need to be provided

including those with compensated cirrhosis - can be treated with DAAs by non-specialist practitioners (e.g. general practitioners (GPs)) in primary care settings. GPs treating HCV patients with compensated cirrhosis will be critical to ensuring adequate access to specialists for those who require specialist care for decompensated cirrhosis.

mary healthcare system.4,6,7

by a specialist. While all patients should have appropri-

ate liver disease assessment prior to treatment to deter-

mine the degree of liver fibrosis or cirrhosis and the

severity of liver function impairment, most patients -

For the first time there has been a reduction in HCV-related deaths globally, clearly demonstrating the impact and benefits of striving for HCV elimination. Despite this, treatment coverage remains low in the Asia-Pacific region, ranging from 5% in South East Asia to 10% in the Western Pacific.2 Only four countries in the Asia-Pacific (Australia, Japan, Mongolia, and Georgia) are on track to reach the World Health Organisation (WHO) elimination goals by 2030.3 Approaches adopted by these countries may be valuable for other countries pursuing elimination goals; common to all four is strong political commitment to achieving national targets through low-cost or subsidised testing and treatment programs.^{4,5} For example, Mongolia undertook epidemiological work to inform the national targets, pursued options for generic DAAs, included these low cost DAAs into the national health insurance scheme, launched a nation-wide screening program, and delivered HCV treatment through integrated services within the pri-

*Corresponding author: Bridget Draper, 85 Commercial Road, Melbourne, 3004 Victoria, Australia, +61 3 8506 2395.

E-mail address: bridget.draper@burnet.edu.au (B. Draper).

The Lancet Regional Health - Western Pacific 2022;20: 100359 Published online xxx https://doi.org/10.1016/j. lanwpc.2021.100359

^aDisease Elimination Program, Burnet Institute Melbourne, Australia

^bSchool of Public Health and Preventive Medicine, Monash University Melbourne, Australia

^cBurnet Institute Yangon, Myanmar

^dHealth Services Research and Implementation, Monash Partners, Melbourne, Australia

^eMyanmar Liver Foundation

fGastroenterologist, Doctors Plaza, Clifton, Karachi, Pakistan

^gYangon Specialty Hospital, Myanmar

^hSt Vincent's Hospital Melbourne, Australia

ⁱDepartment of Medicine, University of Melbourne, Australia

^jHepatitis Services, Department of Infectious Diseases Alfred Hospital Melbourne Australia

^kDoherty Institute, Melbourne, Australia

¹School of Population and Global Health, University of Melbourne, Australia

¹ Ms Bridget Draper, BHSc(Hons): 85 Commercial Road, Melbourne, 3004 Victoria Australia

² Dr Win Lei Yee, MBBS: No 226, 4th Floor, 226 Wizaya Plaza U Wisara Road, Bahan Township, Yangon, Myanmar

³ Dr Alisa Pedrana, PhD: 85 Commercial Road, Melbourne, 3004 Victoria Australia

⁴ Prof Khin Pyone Kyi, PhD: No. 33, 35 Pa Thein St, Yangon, Myanmar

⁵ Dr Huma Qureshi, MBBS: 5th Floor, Block 9 Clifton, The

Plaza, Karachi, Sindh 75600, Pakistan ⁶ Dr Hla Htay, MBBS: No 226, 4th Floor, 226 Wizaya Plaza U

Wisara Road, Bahan Township, Yangon, Myanmar 7 Prof Win Naing, MBBS: No. 33, 35 Pa Thein St, Yangon, Myanmar

⁸ Prof Alexander Thompson, PhD: 41 Victoria Parade, Fitzroy 3065 Victoria Australia

⁹ Prof Margaret Hellard, PhD: 85 Commercial Road, Melbourne, 3004 Victoria Australia

¹⁰ Dr Jessica Howell, PhD: 85 Commercial Road, Melbourne, 3004 Victoria Australia

More than half of the countries within the Asia-Pacific region are classified as low- or middle-income countries (LMICs).8 Many LMICs face multi-faceted barriers to scaling up access to HCV care: low public awareness of HCV, inadequate specialist workforce, limited implementation of decentralised care within existing health systems, and the unaffordable price of diagnostics and DAAs.4 In most LMICs, there are not enough specialists to provide care to those requiring their services. Furthermore, the specialist workforce is often concentrated in major urban health centres. In the case of HCV, the high workloads of specialists lead to long waiting times for patients, hampering timely treatment and leading to many becoming disengaged from care. 9,10 While the cost of diagnostics and DAAs remains a barrier to scaling up access to HCV care, the price of DAAs is continually declining, and some promising patent-free options are in the pipeline.4,II However, achieving global HCV elimination requires more than merely access to affordable DAAs.

Increasing access to treatment is integral to reducing liver disease-related deaths and achieving HCV elimination. One way to achieve this is to train non-specialist practitioners (e.g., GPs, nurse practitioners, pharmacists) to treat HCV with DAAs. We call for a renewed focus on treating people with compensated cirrhosis in primary care. This will allow for increased access to treatment for all and give priority access to specialists for those who require specialist management. General practitioners and other non-specialist providers in primary care settings can easily treat HCV in those with compensated cirrhosis, without the need of a specialist. 12 Referral to the specialist for further evaluation of ongoing liver disease can be done after the completion of DAA treatment. This approach will ensure that people with HCV cirrhosis are treated early, preserving liver function and reducing the risk of liver cancer. I,13 Evidence from one cohort of patients with confirmed liver cirrhosis through liver biopsy found that sustained virological response (SVR) was associated with reduced incidence of hepatic decompensation and liver cancer. 14 Evidence from a study using transient elastography to measure liver stiffness found that approximately one third of cirrhotic patients were reclassified as non-cirrhotic post-SVR.¹⁵ Another study found that those with Child-Pugh A cirrhosis had significantly reduced likelihood of clinical disease progression if they achieved SVR, compared to those who did not achieve a cure. 16 Evidence from these studies demonstrate how treatment induced cure reduces the risk of liver disease progression and the risk of developing liver cancer.

In addition, reducing the number of patients referred to specialists for DAA treatment will allow specialists more time to provide appropriate treatment to patients with advanced liver disease. Even in LMICs, strategies to reduce progression of cirrhosis among those with decompensated cirrhosis are available, for

example: use of beta-blockers to reduce portal hypertension, banding of oesophageal varices in bleeders and those at high risk of bleeding (requiring access to endoscopy), and the use of lactulose to prevent hepatic encephalopathy.^{13,17}

Key to expanding safe and effective HCV treatment by GPs is the development of clear HCV testing and treatment guidelines, and the simplified tools to assess the degree of fibrosis or cirrhosis and whether the cirrhosis is compensated or decompensated, prior to initiating treatment. Identifying whether a patient has decompensated cirrhosis and requires referral is a pillar of differentiated care, where specific groups receive different levels of care depending on need. Liver disease assessment can be performed by trained non-specialist practitioners in primary care settings. Transient elastography of the liver (e.g., FibroScan®) is currently used as best practice, as it provides a clear clinical picture of the liver fibrosis and aids with planning post-DAA therapy care. However, FibroScan® technology has accessibility and affordability issues for most people in the Asia-Pacific. Further, FibroScan® is a specialised technology for liver fibrosis assessment which requires frequent maintenance, and intensive training for operators to perform the scan and interpret the results. 18,19 Recognising this, the WHO Guidelines for HCV treatment (2018) recommend aspartate aminotransferase (AST) /platelet ratio index (APRI) score or FIB-4 test to assess liver fibrosis in resource limited settings. In many settings, APRI score is used to triage patients for FibroScan®. In others, it is used in conjunction with assessment for physical signs of liver decompensation. For example, the Myanmar National Guidelines require GPs to calculate the APRI score to decide treatment duration and to refer those with physical signs of hepatic decompensation to a specialist.20 The Pakistan HCV Testing and Treatment Guidelines also allow GPs to decide the duration of treatment based on APRI score, and only refer those with APRI >1.5 to a specialist.21 The simple blood tests (AST, platelets) required for APRI score are widely available in most settings. Furthermore, it is worth noting that the presence of compensated cirrhosis does not change treatment duration recommendations for some treatment regimens. DAAs are safe and effective in people with compensated cirrhosis, and HCV eradication through DAA therapy is itself associated with reduced risk of developing cirrhosis and liver cancer. As such, HCV treatment should not be unduly delayed by requirements for special tests for cirrhosis (e.g. FibroScan®, ultrasound); especially in settings where the specialist management for portal hypertension or liver cancer screening is not widely available or utilised.

Task-shifting to GPs and decentralisation of care into primary care settings is supported by growing evidence globally. ^{22,23} Systematic reviews have shown GPs can safely and effectively perform liver disease assessment

3

and initiate DAA therapy for compensated cirrhotic patients in community settings, resulting in few adverse outcomes and high sustained virological response (SVR12) rates.^{22,23} Ensuring patient safety is paramount, and this can be facilitated through use of quality-assured diagnostics and laboratories, and through adequate training and support for GPs. Clear clinical decision-making tools, including APRI score calculator, drug-drug interaction checkers, and summary tables of key pre-treatment assessments and treatment options, are available in many settings to support GPs to prescribe DAAs. For example, the EC toolkit in Australia was developed to collate available tools and develop extra tools to address unmet needs within primary care in Australia (see: https://ecpartnership.org.au/toolkit).

Task-shifting care to GPs and decentralisation into primary care settings will allow for increased access to HCV treatment outside of urban centres, ensuring more equitable access to care. Country-specific assessments on where to focus efforts, considering the epidemiology of disease burden, will be required to inform distribution of resources needed to serve the general population, or various key population groups including people who inject drugs, people in prison, or people living with HIV. Implementation of robust national data systems would improve the data available to inform strategies and to monitor progress. Such country-specific assessments can also help inform the testing strategy and assist practitioners to target testing, either through regular testing of key population groups and/ or to general population one-time screening; the approach will be country-specific based on estimated HCV prevalence among general population and key population groups, considering resource availability.24 With low-cost rapid diagnostics available for screening, it is unlikely that excessive screening of HCV will occur; however, it is important to ensure all those who screen positive have access to confirmatory nucleic acid testing to diagnose active infection.

Successful implementation of simplified clinical pathways in secondary and tertiary based hospitals, and some at decentralised HIV treatment sites, through national programs has been demonstrated in Myanmar, Cambodia, India, and Indonesia.²⁵ Our work in Myanmar and recent work from Cambodia showcase feasible and effective models of care for community-based treatment, using simplified clinical pathways.26,27 Both studies found that treatment of compensated cirrhotic patients by GPs at community-based sites was safe and effective, with high cure rates and few adverse events. Uptake of treatment was high and there was limited loss to follow up from care. 26,27 Task-shifting to GPs required minimal training. Setting clear referral criteria and pathways was important in the Myanmar model for ensuring GP confidence.²⁶ Decentralised and non-specialist led care has been successfully implemented in many countries in the Asia-Pacific region. In the

Western Pacific region, a recent review of progress towards HCV elimination found that ten countries have now expanded their hepatitis services into primary care. For example, in Malaysia following pilot of decentralised care in primary health care clinics, now 146 primary healthcare clinics are providing treatment, with care mostly provided by trained GPs.²⁸ In Australia, GP-led HCV treatment is common; almost half of patients initiated onto treatment from 2016 to 2020 had their prescription written by a GP.29 Initially, remote specialist consultation and approval was required by GPs without experience prescribing DAAs. This was facilitated through remote consultation forms and successful allowed treatment of cirrhotic patients by GPs where access to a specialist was limited.^{30,31} A similar process also occurs for the nurse-led prison treatment service in Victoria, Australia where nurses could provide treatment to those in prison via remote consultation form for those without cirrhosis or with compensated cirrhosis.32 Simplified care models with non-specialist practitioners prescribing DAAs could be scaled to reach the level of coverage we need to achieve elimination goals, especially in LMICs where there are very few specialists.

Our challenge now is to ensure that national guidelines allow GPs and other non-specialist practitioners to treat patients with compensated cirrhosis, to provide frequent training for these practitioners, and to develop clear, effective referral pathways for specialist review or advice. Training, mentorship, and support to GPs can be provided remotely through the Project ECHO (Extension for Community Healthcare Outcomes) model.33 Project ECHO is a tele-mentoring hub-and-spoke model where one hospital provides multi-disciplinary support for a network of primary care providers through use of frequent meetings for specialists to share best practices on HCV testing and treatment, and for community providers to share de-identified case presentations for discussion and recommendations for a treatment plan.33 Evidence from hepatitis C and other diseases / specialty areas has shown that a Project ECHO tele-mentoring model can be used successfully for training, ongoing support, and provision of continuous medical education.33-35 Another option for specialist support is through the use of remote consultation with a specialist; described in detail above, this process is widely used by GPs in Australia.^{30,31}

Other considerations for successfully embedding HCV care into primary care include: encouraging GPs to get trained and provide HCV care, accessibility of laboratories for HCV RNA testing and liver function tests, and navigating the competing workloads of GPs in primary care; these must be addressed within each local context. While ideally all cirrhotic patients treated by GPs will then receive ongoing post-DAA therapy liver care to monitor for liver disease progression, in particular liver cancer surveillance through biannual ultrasounds, this is

unlikely to be feasible or widely accessible in most resource-constrained settings and alternative more-pragmatic clinical guidelines are also needed. For example, ultrasounds are available in private clinics and hospitals in Myanmar, but access is limited in the public health system; biannual ultrasound, alfa feta-protein and platelets is recommended. Referral pathways for those with advanced disease or potentially liver cancer should be strengthened where possible, to ensure those who require specialist management can access it. By focusing on treating patients with cirrhosis early we can reduce the future burden of liver disease in resource-constrained settings across the Asia-Pacific region.

Decentralised care and non-specialist led HCV care will be key to expanding testing and treatment to all people affected by HCV. It is time to invest in scaling up decentralised care, along with training GPs, to ensure that all those living with HCV have access to a cure, particularly those with cirrhosis who are most likely to benefit from timely treatment to reduce their risk of liver disease progression.

Declaration of interests

MH has received investigator-initiated grant funding from Gilead Sciences and Abbvie for unrelated work. AP has received investigator-initiated grant funding from Gilead Sciences and Abbvie and honoraria from Gilead Sciences for unrelated work. AT has received consulting fees for advisory board participation and speaker fees from Gilead Sciences, Abbvie, Merck and BMS for unrelated work. AT also served on the board of directors for Gastroenterological Society of Australia (honorary position). WLY has received Gilead Sciences Public Health Award for unrelated work. KPK has received non-financial support from Mylan (support to attend AASLD Conference), Hetero for DAAs for Myanmar Liver Foundation Charity Clinic and Royal Ruby for donation liver supportive therapies for Myanmar Liver Foundation Charity Clinic. WN has received nonfinancial support from Mylan and Cipla to attend AASLD Conferences. JH has participated in Data Safety Monitoring Board for Gilead Sciences and has served on the Liver Faculty Australian Liver Association (Gastroenterological Society of Australia). All others declare no potential competing interests.

Role of the funding source

No specific funding source for this manuscript. The CT2 Study which informed this work was supported by Unitaid, through the FIND-led HEAD-Start project grant.

Author contributions

BD wrote the original draft, conducted literature search, and edited the manuscript to incorporate co-authors' inputs. JH conceptualised the idea, and reviewed and edited multiple drafts. WLY, AP, KPK, HQ, HH, WN, AT and MH provided input on evidence to include, reviewed and edited draft manuscripts. AP, MH and JH provided supervision. All authors revised the paper critically for intellectual content and approved the final version.

References

- I World Health Organization (WHO). Guidelines for the care and treatment of persons diagnosed with chronic hepatitis C virus infection. 2018. Available from: https://apps.who.int/iris/bitstream/handle/10665/273174/9789241550345-eng.pdf?ua=1
- 2 World Health Organization (WHO). Global progress report on HIV, viral hepatitis and sexually transmitted infections, 2021. 2021. Available from: https://www.who.int/publications/i/item/9789240027077
- 3 CDA Foundation. Just 12 countries worldwide on track to eliminate hepatitis C infection by 2030, with United Kingdom, Italy and Spain among those joining the list CDA Foundation. Available from: http://cdafound.org/just-12-countries-worldwide-on-track-to-eliminate-hepatitis-c-infection-by-2030-with-united-kingdom-italy-and-spain-among-those-joining-the-list/
- 4 World Health Organization (WHO). Progress report on access to hepatitis C treatment: focus on overcoming barriers in low- and middle-income countries. 2018. Available from: https://apps.who.int/iris/handle/10665/260445
- 5 World Health Organization (WHO). Japan's hepatitis programme frees people from disease and financial hardship. Available from: https://www.who.int/australia/news/feature-stories/detail/japan's-hepatitis-programme-frees-people-from-disease-and-financial-hardship
- Dore GJ, Martinello M. Global elimination of hepatitis C virus by 2030: the optimistic view. Clin Dilemmas Viral Liver Dis 2020: 238–43. Available from: https://onlinelibrary.wiley.com/doi/full/10.1002/9781119533481.ch40.
 Chan P-L, Le L-V, Ishikawa N, Easterbrook P. Regional progress
- 7 Chan P-L, Le L-V, Ishikawa N, Easterbrook P. Regional progress towards hepatitis C elimination in the Western Pacific Region, 2015-2020. Glob Heal Med 2021;3(5):253-61.
- 8 World Bank. The World Bank Country and Lending Groups. 2020. Available from: https://datahelpdesk.worldbank.org/knowledge-base/articles/906519-world-bank-country-and-lending-groups
- 9 Palmer AY, Wade AJ, Draper B, Howell J, Doyle JS, Petrie D, et al. A cost-effectiveness analysis of primary versus hospital-based specialist care for direct acting antiviral hepatitis C treatment. *Int J Drug Policy* 2020;76:102633.
- Io Wade AJ, Doyle JS, Gane E, Stedman C, Draper B, Iser D, et al. Outcomes of Treatment for Hepatitis C in Primary Care, Compared to Hospital-based Care: A Randomized, Controlled Trial in People Who Inject Drugs. Clin Infect Dis 2019.
- II Andrieux-Meyer I, Tan S-S, Thanprasertsuk S, Salvadori N, Menetrey C, Simon F, et al. Efficacy and safety of ravidasvir plus sofosbuvir in patients with chronic hepatitis C infection without cirrhosis or with compensated cirrhosis (STORM-C-I): interim analysis of a two-stage, open-label, multicentre, single arm, phase 2/3 trial. Lancet Gastroenterol Hepatol 2021.
- 12 Tran TT. Hepatitis C: Who should treat hepatitis C virus? The role of the primary care provider. *Clin Liver Dis* 2018 Mar 1;11(3):66–8.
- 13 Ginès P, Krag A, Abraldes JG, Solà E, Fabrellas N, Kamath PS. Liver cirrhosis. Lancet 2021 Oct 9;398(10308):1359-76.
- I4 Nahon P, Bourcier V, Layese R, Audureau E, Cagnot C, Marcellin P, et al. Eradication of Hepatitis C Virus Infection in Patients With Cirrhosis Reduces Risk of Liver and Non-Liver Complications. Gastroenterology 2017;152(1):142–56. e2.
- 15 Fernandes FF, Piedade J, Guimaraes L, Nunes EP, Chaves U, Goldenzon RV, et al. Effectiveness of direct-acting agents for hepatitis C and liver stiffness changing after sustained virological response. J Gastroenterol Hepatol 2019;34(12):2187–95.
- 16 Krassenburg LAP, Maan R, Ramji A, Manns MP, Cornberg M, Wedemeyer H, et al. Clinical outcomes following DAA therapy in patients with HCV-related cirrhosis depend on disease severity. J Hepatol 2021 May 1;74(5):1053–63.
- 17 EASL EASL. Clinical Practice Guidelines for the management of patients with decompensated cirrhosis. J Hepatol 2018 Aug 1;69 (2):406–60.
- 18 Londeix P. Diagnosis and monitoring of hepatitis C (HCV) in Morocco: Current status and strategies for universal access.

- Benchmarking HCV diagnostics, fibrosis evaluation, and treatment monitoring. 2018. Available from: https://www.alcs.ma/wp-content/uploads/2018/05/BenchmarkVA.pdf
- 19 FibroScan for assessing liver fibrosis and cirrhosis in primary care Medtech innovation briefing. 2020. Available from: www.nice.org. uk/guidance/mib216
- 20 Ministry of Health and Sports Myanmar. Myanmar National Simplified Treatment Guidelines for Hepatitis C Infection Second Edition, July 2019. 2019. Available from: https://mohs.gov.mm/page/3212
- 21 Pakistan HCV Treatment Guidelines, 2020. 2020. Available from: http://phrc.org.pk/assets/pakistan-national-hcv-treatment-guidelines-sample-2.pdf
- 22 Oru E, Trickey A, Shirali R, Kanters S, Easterbrook P. Decentralisation, integration, and task-shifting in hepatitis C virus infection testing and treatment: a global systematic review and meta-analysis. Lancet Glob Heal 2021.
- 23 Radley A, Robinson E, Aspinall EJ, Angus K, Tan L, Dillon JF. A systematic review and meta-analysis of community and primary-care-based hepatitis C testing and treatment services that employ direct acting antiviral drug treatments. *BMC Health Serv Res* 2019 Dec 28:19(1):765.
- 24 Easterbrook PJ. Who to test and how to test for chronic hepatitis C infection –2016 WHO testing guidance for low- and middle-income countries. *J Hepatol* 2016 Oct;65(1):S46–66.
- 25 Boeke CE, Adesigbin C, Agwuocha C, Anartati A, Aung HT, Aung KS, et al. Initial success from a public health approach to hepatitis C testing, treatment and cure in seven countries: The road to elimination. BMJ Global Health 2020;5:3767.
- 26 Draper B, Htay H, Pedrana A, Yee WL, Howell J, Kyi KP, et al. Outcomes of the CT2 Study: A 'one-stop-shop' for community-based hepatitis C testing and treatment in Yangon. Myanmar. Liver Int. 2021: liv.14983. Jun 21.
- 27 Zhang M, O'Keefe D, Craig J, Samley K, Bunreth V, Jolivet P, et al. Decentralised hepatitis C testing and treatment in rural Cambodia:

- evaluation of a simplified service model integrated in an existing public health system. *Lancet Gastroenterol Hepatol* 2021.
- 28 Sun J, Cheng H, Hassan MRA, Chan H-K, Piedagnel J-M. What China can learn from Malaysia to achieve the goal of "eliminate hepatitis C as a public health threat" by 2030 – a narrative review. Lancet Reg Heal - West Pacific 2021;16:100261.
- 29 Stafford F, Dore GJ, Clackett S, Martinello M, Matthews GV, Grebely J, et al. Prescribing of direct-acting antiviral therapy by general practitioners for people with hepatitis C in an unrestricted treatment program. Med J Aust 2021 Aug 23;215(7):332–3.
- Haridy J, Iyngkaran G, Nicoll A, Muller K, Wilson M, Wigg A, et al. Outcomes of community-based hepatitis C treatment by general practitioners and nurses in Australia through remote specialist consultation. *Intern Med J* 2021 Nov 1;51(11):1927–34.
 Wade AJ, McCormack A, Roder C, McDonald K, Davies M, Scott N,
- 31 Wade AJ, McCormack A, Roder C, McDonald K, Davies M, Scott N, et al. Aiming for elimination: Outcomes of a consultation pathway supporting regional general practitioners to prescribe direct-acting antiviral therapy for hepatitis C. J Viral Hepat 2018 Sep 1;25 (9):1089-98.
- 32 Papaluca T, McDonald L, Craigie A, Hellard M, Iser D, Thompson A, et al. Outcomes of treatment for hepatitis C in prisoners using a nurse-led, statewide model of care. J Hepatol 2019;70:839–46.
- 33 Arora S. Project ECHO: democratising knowledge for the elimination of viral hepatitis. Lancet Gastroenterol Hepatol 2019 Feb 1;4 (2):91-3.
- 34 Bikinesi L, O'Bryan G, Roscoe C, Mekonen T, Shoopala N, Mengistu AT, et al. Implementation and evaluation of a Project ECHO telementoring program for the Namibian HIV workforce. *Hum Resour Health* 2020 Sep 1;18(1):1–10.
- 35 Nhung LH, Dien TM, Lan NP, Thanh PQ, Cuong PV. Use of Project ECHO Telementoring Model in Continuing Medical Education for Pediatricians in Vietnam: Preliminary Results. Health Services Insights 2021 Aug 14;14.