

Mapping age- and sex-specific HIV prevalence in adults in sub-Saharan Africa, 2000–2018

HAEUSER, Emily, SERFES, Audrey L., CORK, Michael A., YANG, Mingyou, ABBASTABAR, Hedayat, ABHILASH, E. S., ADABI, Maryam, ADEBAYO, Oladimeji M., ADEKANMBI, Victor, ADEYINKA, Daniel Adedayo, AFZAL, Saira, AHINKORAH, Bright Opoku, AHMADI, Keivan, AHMED, Muktar Beshir, AKALU, Yonas, AKINYEMI, Rufus Olusola, AKUNNA, Chisom Joyqueenet, ALAHDAB, Fares, ALANEZI, Fahad Mashhour, ALANZI, Turki M., ALENE, Kefyalew Addis, ALHASSAN, Robert Kaba, ALIPOUR, Vahid, ALMASI-HASHIANI, Amir, ALVIS-GUZMAN, Nelson, AMEYAW, Edward Kwabena, AMINI, Saeed, AMUGSI, Dickson A., ANCUCEANU, Robert, ANVARI, Davood, APPIAH, Seth Christopher Yaw, ARABLOO, Jalal, AREMU, Olatunde, ASEMAHAGN, Mulusew A., JAFARABADI, Mohammad Asghari, AWEDEW, Atalel Fentahun, QUINTANILLA, Beatriz Paulina Ayala, AYANORE, Martin Amogre, AYNALEM, Yared Asmare, AZARI, Samad, AZENE, Zelalem Nigussie, DARSHAN, B. B., BABALOLA, Tesleem Kayode, BAIG, Atif Amin, BANACH, Maciej, BÄRNIGHAUSEN, Till Winfried, BELL, Arielle Wilder, BHAGAVATHULA, Akshaya Srikanth, BHARDWAJ, Nikha, BHARDWAJ, Pankaj, BHATTACHARYYA, Kritika, BIJANI, Ali, BITEW, Zebeay Workneh, BOHLOULI, Somayeh, BOLARINWA, Obasanjo Afolabi, BOLOOR, Archith, BOZICEVIC, Ivana, BUTT, Zahid A., CÁRDENAS, Rosario, CARVALHO, Felix, CHARAN, Jaykaran, CHATTU, Vijay Kumar, CHOWDHURY, Mohiuddin Ahsanul Kabir, CHU, Dinh-Toi, COWDEN, Richard G., DAHLAWI, Saad M. A., DAMIANI, Giovanni, DARTEH, Eugene Kofuor Maafo, DARWESH, Aso Mohammad, DAS NEVES, José, WEAVER, Nicole Davis, DE LEO, Diego, DE NEVE, Jan-Walter, DERIBE, Kebede, DEUBA, Keshab, DHARMARATNE, Samath, DIANATINASAB, Mostafa, DIAZ, Daniel, DIDARLOO, Alireza, DJALALINIA, Shirin, DOROSTKAR, Fariba, DUBLJANIN, Eleonora, DUKO, Bereket, EL TANTAWI, Maha, EL-JAAFARY, Shaimaa I., ESHRATI, Babak, ESKANDARIEH, Sharareh, EYAWO, Oghenowede, EZEONWUMELU, Ifeanyi Jude, EZZIKOURI, Sayeh, FARZADFAR, Farshad, FATTAHI, Nazir, FAUK, Nelsensius Klau, FERNANDES, Eduarda, FILIP, Irina, FISCHER, Florian, FOIGT, Nataliya A., FOROUTAN, Masoud, FUKUMOTO, Takeshi, GAD, Mohamed M., GAIDHANE, Abhay Motiramji, GEBREGIORGIS, Birhan Gebresillassie, GEBREMEDHIN, Ketema Bizuwork, GETACHER, Lemma, GHADIRI, Keyghobad, GHASHGHAE, Ahmad, GOLECHHA, Mahaveer, GUBARI, Mohammed Ibrahim Mohialdeen, GUGNANI, Harish Chander, GUIMARÃES, Rafael Alves, HAIDER, Mohammad Rifat, HAJ-MIRZAIAN, Arvin, HAMIDI, Samer, HASHI, Abdiwahab, HASSANIPOUR, Soheil, HASSANKHANI, Hadi, HAYAT, Khezar, HERTELIU, Claudiu, HO, Hung Chak,

HOLLA, Ramesh, HOSSEINI, Mostafa, HOSSEINZADEH, Mehdi, HWANG, Bing-Fang, IBITOYE, Segun Emmanuel, ILESANMI, Olayinka Stephen, ILIC, Irena M., ILIC, Milena D., ISLAM, Rakibul M., IWU, Chidozie C. D., JAKOVLJEVIC, Mihajlo, JHA, Ravi Prakash, JI, John S., JOHNSON, Kimberly B., JOSEPH, Nitin, JOSHUA, Vasna, JOUKAR, Farahnaz, JOZWIAK, Jacek Jerzy, KALANKESH, Leila R., KALHOR, Rohollah, KAMYARI, Naser, KANCHAN, Tanuj, MATIN, Behzad Karami, KARIMI, Salah Eddin, KAYODE, Gbenga A., KARYANI, Ali Kazemi, KERAMATI, Maryam, KHAN, Ejaz Ahmad, KHAN, Gulfaraz, KHAN, Md Nuruzzaman, KHATAB, Khaled <<http://orcid.org/0000-0002-8755-3964>>, KHUBCHANDANI, Jagdish, KIM, Yun Jin, KISA, Adnan, KISA, Sezer, KOPEC, Jacek A., KOSEN, Soewarta, LAXMINARAYANA, Sindhura Lakshmi Koulmane, KOYANAGI, Ai, KRISHAN, Kewal, DEFO, Barthelémy Kuate, KUGBEY, Nuworza, KULKARNI, Vaman, KUMAR, Manasi, KUMAR, Nithin, KUSUMA, Dian, LA VECCHIA, Carlo, LAL, Dharmesh Kumar, LANDIRES, Iván, LARSON, Heidi Jane, LASRADO, Savita, LEE, Paul H., LI, Shanshan, LIU, Xuefeng, MALEKI, Afshin, MALIK, Preeti, MANSOURNIA, Mohammad Ali, MARTINS-MELO, Francisco Rogerlândio, MENDOZA, Walter, MENEZES, Ritesh G., MENGESHA, Endalkachew Worku, MERETOJA, Tuomo J., MESTROVIC, Tomislav, MIRICA, Andreea, MOAZEN, Babak, MOHAMAD, Osama, MOHAMMAD, Yousef, MOHAMMADIAN-HAFSHEJANI, Abdollah, MOHAMMADPOURHODKI, Reza, MOHAMMED, Salahuddin, MOHAMMED, Shafiu, MOKDAD, Ali H., MORADI, Masoud, MORAGA, Paula, MUBARIK, Sumaira, MULU, Getaneh Baye B., MWANRI, Lillian, NAGARAJAN, Ahamarshan Jayaraman, NAIMZADA, Mukhammad David, NAVEED, Muhammad, NAZARI, Javad, NDEJJO, Rawlance, NEGOL, Ionut, NGALESONI, Frida N., NGUEFACK-TSAGUE, Georges, NGUNJIRI, Josephine W., NGUYEN, Cuong Tat, NGUYEN, Huong Lan Thi, NNAJI, Chukwudi A., NOUBIAP, Jean Jacques, NUÑEZ-SAMUDIO, Virginia, NWATAH, Vincent Ebuka, OANCEA, Bogdan, ODUKOYA, Oluwakemi Ololade, OLAGUNJU, Andrew T., OLAKUNDE, Babayemi Oluwaseun, OLUSANYA, Bolajoko Olubukunola, OLUSANYA, Jacob Olusegun, BALI, Ahmed Omar, ONWUJEKWE, Obinna E., ORISAKWE, Orish Ebere, OTSTAVNOV, Nikita, OTSTAVNOV, Stanislav S., OWOLABI, Mayowa O., MAHESH, P. A., PADUBIDRI, Jagadish Rao, PANA, Adrian, PANDEY, Ashok, PANDI-PERUMAL, Seithikurippu R., KAN, Fatemeh Pashazadeh, PATTON, George C., PAWAR, Shrikant, PEPRAH, Emmanuel K., POSTMA, Maarten J., PREOTESCU, Liliana, SYED, Zahiruddin Quazi, RABIEE, Navid, RADFAR, Amir, RAFIEI, Alireza, RAHIM, Fakher, RAHIMI-MOVAGHAR, Vafa, RAHMANI, Amir Masoud, RAMEZANZADEH, Kiana, RANA, Juwel, RANABHAT, Chhabi Lal, RAO, Sowmya J., RAWAF, David Laith, RAWAF, Salman, RAWASSIZADEH, Reza, REGASSA, Lemma Demissie, REZAEI, Nima, REZAPOUR, Aziz, RIAZ, Mavra A., RIBEIRO, Ana Isabel, ROSS, Jennifer M., RUBAGOTTI, Enrico, RUMISHA, Susan Fred, RWEGERERA, Godfrey M., MOGHADDAM, Sahar Saeedi, SAGAR, Rajesh, SAHILEDENGLE, Biniyam, SAHU, Maitreyi, SALEM, Marwa Rashad, KAFIL, Hossein Samadi, SAMY, Abdallah M., SARTORIUS, Benn, SATHIAN, Brijesh, SEIDU, Abdul-Aziz, SHAHEEN, Amira A., SHAIKH, Masood Ali, SHAMSIZADEH, Morteza, SHIFERAW, Wondimeneh Shibabaw, SHIN, Jae Il,

SHRESTHA, Roman, SINGH, Jasvinder A., SKRYABIN, Valentin Yurievich, SKRYABINA, Anna Aleksandrovna, SOLTANI, Shahin, SUFIYAN, Mu'awiyah Babale, TABUCHI, Takahiro, TADESSE, Eyayou Girma, TAVEIRA, Nuno, TEFAY, Fisaha Haile, THAPAR, Rekha, TOVANI-PALONE, Marcos Roberto, TSEGAYE, Gebiyaw Wudie, UMEOKONKWO, Chukwuma David, UNNIKRISHNAN, Bhaskaran, VILLAFANE, Jorge Hugo, VIOLANTE, Francesco S., VO, Bay, VU, Giang Thu, WADO, Yohannes Dibaba, WAHEED, Yasir, WAMAI, Richard G., WANG, Yanzhong, WARD, Paul, WICKRAMASINGHE, Nuwan Darshana, WILSON, Katherine, YAYA, Sanni, YIP, Paul, YONEMOTO, Naohiro, YU, Chuanhua, ZASTROZHIN, Mikhail Sergeevich, ZHANG, Yunquan, ZHANG, Zhi-Jiang, HAY, Simon I. and DWYER-LINDGREN, Laura

Available from Sheffield Hallam University Research Archive (SHURA) at:

<https://shura.shu.ac.uk/31185/>

This document is the Published Version [VoR]

Citation:

HAEUSER, Emily, SERFES, Audrey L., CORK, Michael A., YANG, Mingyou, ABBASTABAR, Hedayat, ABHILASH, E. S., ADABI, Maryam, ADEBAYO, Oladimeji M., ADEKANMBI, Victor, ADEYINKA, Daniel Adedayo, AFZAL, Saira, AHINKORAH, Bright Opoku, AHMADI, Keivan, AHMED, Muktar Beshir, AKALU, Yonas, AKINYEMI, Rufus Olusola, AKUNNA, Chisom Joyqueenet, ALAHDAB, Fares, ALANEZI, Fahad Mashhour, ALANZI, Turki M., ALENE, Kefyalew Addis, ALHASSAN, Robert Kaba, ALIPOUR, Vahid, ALMASI-HASHIANI, Amir, ALVIS-GUZMAN, Nelson, AMEYAW, Edward Kwabena, AMINI, Saeed, AMUGSI, Dickson A., ANCUCEANU, Robert, ANVARI, Davood, APPIAH, Seth Christopher Yaw, ARABLOO, Jalal, AREMU, Olatunde, ASEMAHAGN, Mulusew A., JAFARABADI, Mohammad Asghari, AWEDEW, Atalel Fentahun, QUINTANILLA, Beatriz Paulina Ayala, AYANORE, Martin Amogre, AYNALEM, Yared Asmare, AZARI, Samad, AZENE, Zelalem Nigussie, DARSHAN, B. B., BABALOLA, Tesleem Kayode, BAIG, Atif Amin, BANACH, Maciej, BÄRNIGHAUSEN, Till Winfried, BELL, Arielle Wilder, BHAGAVATHULA, Akshaya Srikanth, BHARDWAJ, Nikha, BHARDWAJ, Pankaj, BHATTACHARYYA, Kritika, BIJANI, Ali, BITEW, Zebeay Workneh, BOHLOULI, Somayeh, BOLARINWA, Obasanjo Afolabi, BOLOOR, Archith, BOZICEVIC, Ivana, BUTT, Zahid A., CÁRDENAS, Rosario, CARVALHO, Felix, CHARAN, Jaykaran, CHATTU, Vijay Kumar, CHOWDHURY, Mohiuddin Ahsanul Kabir, CHU, Dinh-Toi, COWDEN, Richard G., DAHLAWI, Saad M. A., DAMIANI, Giovanni, DARTEH, Eugene Kofuor Maafo, DARWESH, Aso Mohammad, DAS NEVES, José, WEAVER, Nicole Davis, DE LEO, Diego, DE NEVE, Jan-Walter, DERIBE, Kebede, DEUBA, Keshab, DHARMARATNE, Samath, DIANATINASAB, Mostafa, DIAZ, Daniel, DIDARLOO, Alireza, DJALALINIA, Shirin, DOROSTKAR, Fariba, DUBLJANIN,

Sheffield Hallam University Research Archive

<http://shura.shu.ac.uk>

Eleonora, DUKO, Bereket, EL TANTAWI, Maha, EL-JAAFARY, Shaimaa I.,
 ESHRATI, Babak, ESKANDARIEH, Sharareh, EYAWO, Oghenowede,
 EZEONWUMELU, Ifeanyi Jude, EZZIKOURI, Sayeh, FARZADFAR, Farshad,
 FATTAHI, Nazir, FAUK, Nelsensius Klau, FERNANDES, Eduarda, FILIP, Irina,
 FISCHER, Florian, FOIGT, Nataliya A., FOROUTAN, Masoud, FUKUMOTO, Takeshi,
 GAD, Mohamed M., GAIDHANE, Abhay Motiramji, GEBREGIORGIS, Birhan
 Gebresillassie, GEBREMEDHIN, Ketema Bizuwork, GETACHER, Lemma, GHADIRI,
 Keyghobad, GHASHGHAE, Ahmad, GOLECHHA, Mahaveer, GUBARI,
 Mohammed Ibrahim Mohialdeen, GUGNANI, Harish Chander, GUIMARÃES, Rafael
 Alves, HAIDER, Mohammad Rifat, HAJ-MIRZAIAN, Arvin, HAMIDI, Samer, HASHI,
 Abdiwahab, HASSANIPOUR, Soheil, HASSANKHANI, Hadi, HAYAT, Khezar,
 HERTELIU, Claudiu, HO, Hung Chak, HOLLA, Ramesh, HOSSEINI, Mostafa,
 HOSSEINZADEH, Mehdi, HWANG, Bing-Fang, IBITOYE, Segun Emmanuel,
 ILESANMI, Olayinka Stephen, ILIC, Irena M., ILIC, Milena D., ISLAM, Rakibul M.,
 IWU, Chidozie C. D., JAKOVLJEVIC, Mihajlo, JHA, Ravi Prakash, JI, John S.,
 JOHNSON, Kimberly B., JOSEPH, Nitin, JOSHUA, Vasna, JOUKAR, Farahnaz,
 JOZWIAK, Jacek Jerzy, KALANKESH, Leila R., KALHOR, Rohollah, KAMYARI,
 Naser, KANCHAN, Tanuj, MATIN, Behzad Karami, KARIMI, Salah Eddin, KAYODE,
 Gbenga A., KARYANI, Ali Kazemi, KERAMATI, Maryam, KHAN, Ejaz Ahmad, KHAN,
 Gulfaraz, KHAN, Md Nuruzzaman, KHATAB, Khaled, KHUBCHANDANI, Jagdish,
 KIM, Yun Jin, KISA, Adnan, KISA, Sezer, KOPEC, Jacek A., KOSEN, Soewarta,
 LAXMINARAYANA, Sindhura Lakshmi Koulmane, KOYANAGI, Ai, KRISHAN, Kewal,
 DEFO, Barthelemy Kuate, KUGBEY, Nuworza, KULKARNI, Vaman, KUMAR,
 Manasi, KUMAR, Nithin, KUSUMA, Dian, LA VECCHIA, Carlo, LAL, Dharmesh
 Kumar, LANDIRES, Iván, LARSON, Heidi Jane, LASRADO, Savita, LEE, Paul H., LI,
 Shanshan, LIU, Xuefeng, MALEKI, Afshin, MALIK, Preeti, MANSOURNIA,
 Mohammad Ali, MARTINS-MELO, Francisco Rogerlândio, MENDOZA, Walter,
 MENEZES, Ritesh G., MENGESHA, Endalkachew Worku, MERETOJA, Tuomo J.,
 MESTROVIC, Tomislav, MIRICA, Andreea, MOAZEN, Babak, MOHAMAD, Osama,
 MOHAMMAD, Yousef, MOHAMMADIAN-HAFSHEJANI, Abdollah,
 MOHAMMADPOURHODKI, Reza, MOHAMMED, Salahuddin, MOHAMMED, Shafiu,
 MOKDAD, Ali H., MORADI, Masoud, MORAGA, Paula, MUBARIK, Sumaira, MULU,
 Getaneh Baye B., MWANRI, Lillian, NAGARAJAN, Ahamarshan Jayaraman,
 NAIMZADA, Mukhammad David, NAVEED, Muhammad, NAZARI, Javad, NDEJJO,
 Rawlance, NEGOI, Ionut, NGALESONI, Frida N., NGUEFACK-TSAGUE, Georges,
 NGUNJIRI, Josephine W., NGUYEN, Cuong Tat, NGUYEN, Huong Lan Thi, NNAJI,
 Chukwudi A., NOUBIAP, Jean Jacques, NUÑEZ-SAMUDIO, Virginia, NWATAH,
 Vincent Ebuka, OANCEA, Bogdan, ODUKOYA, Oluwakemi Ololade, OLAGUNJU,
 Andrew T., OLAKUNDE, Babayemi Oluwaseun, OLUSANYA, Bolajoko Olubukunola,
 OLUSANYA, Jacob Olusegun, BALI, Ahmed Omar, ONWUJEKWE, Obinna E.,
 ORISAKWE, Orish Ebere, OTSTAVNOV, Nikita, OTSTAVNOV, Stanislav S.,
 OWOLABI, Mayowa O., MAHESH, P. A., PADUBIDRI, Jagadish Rao, PANA, Adrian,
 PANDEY, Ashok, PANDI-PERUMAL, Seithikurippu R., KAN, Fatemeh Pashazadeh,
 PATTON, George C., PAWAR, Shrikant, PEPRAH, Emmanuel K., POSTMA, Maarten
 J., PREOTESCU, Liliana, SYED, Zahiruddin Quazi, RABIEE, Navid, RADFAR, Amir,
 RAFIEI, Alireza, RAHIM, Fakher, RAHIMI-MOVAGHAR, Vafa, RAHMANI, Amir
 Masoud, RAMEZANZADEH, Kiana, RANA, Juwel, RANABHAT, Chhabi Lal, RAO,
 Sowmya J., RAWAF, David Laith, RAWAF, Salman, RAWASSIZADEH, Reza,
 REGASSA, Lemma Demissie, REZAEI, Nima, REZAPOUR, Aziz, RIAZ, Mavra A.,
 RIBEIRO, Ana Isabel, ROSS, Jennifer M., RUBAGOTTI, Enrico, RUMISHA, Susan
 Fred, RWEGERERA, Godfrey M., MOGHADDAM, Sahar Saeedi, SAGAR, Rajesh,
 SAHILEDENGLE, Biniyam, SAHU, Maitreyi, SALEM, Marwa Rashad, KAFIL,

Hossein Samadi, SAMY, Abdallah M., SARTORIUS, Benn, SATHIAN, Brijesh, SEIDU, Abdul-Aziz, SHAHEEN, Amira A., SHAIKH, Masood Ali, SHAMSIZADEH, Morteza, SHIFERAW, Wondimeneh Shibabaw, SHIN, Jae Il, SHRESTHA, Roman, SINGH, Jasvinder A., SKRYABIN, Valentin Yurievich, SKRYABINA, Anna Aleksandrovna, SOLTANI, Shahin, SUFIYAN, Mu'awiyah Babale, TABUCHI, Takahiro, TADESSE, Eyayou Girma, TAVEIRA, Nuno, TSEFAY, Fisaha Haile, THAPAR, Rekha, TOVANI-PALONE, Marcos Roberto, TSEGAYE, Gebiyaw Wudie, UMEOKONKWO, Chukwuma David, UNNIKRISHNAN, Bhaskaran, VILLAFANE, Jorge Hugo, VIOLANTE, Francesco S., VO, Bay, VU, Giang Thu, WADO, Yohannes Dibaba, WAHEED, Yasir, WAMAI, Richard G., WANG, Yanzhong, WARD, Paul, WICKRAMASINGHE, Nuwan Darshana, WILSON, Katherine, YAYA, Sanni, YIP, Paul, YONEMOTO, Naohiro, YU, Chuanhua, ZASTROZHIN, Mikhail Sergeevich, ZHANG, Yunquan, ZHANG, Zhi-Jiang, HAY, Simon I. and DWYER-LINDGREN, Laura (2022). Mapping age- and sex-specific HIV prevalence in adults in sub-Saharan Africa, 2000–2018. BMC Medicine, 20 (1): 488. [Article]

Copyright and re-use policy

See <http://shura.shu.ac.uk/information.html>

RESEARCH ARTICLE

Open Access



Mapping age- and sex-specific HIV prevalence in adults in sub-Saharan Africa, 2000–2018

Emily Haeuser^{1*}, Audrey L. Serfes¹, Michael A. Cork¹, Mingyou Yang¹, Hedayat Abbastabar², E. S. Abhilash³, Maryam Adabi⁴, Oladimeji M. Adebayo⁵, Victor Adekanmbi⁶, Daniel Adedayo Adeyinka^{7,8}, Saira Afzal^{9,10}, Bright Opoku Ahinkorah¹¹, Keivan Ahmadi¹², Muktar Beshir Ahmed^{13,14}, Yonas Akalu¹⁵, Rufus Olusola Akinyemi^{16,17}, Chisom Joyqueenet Akunna^{18,19}, Fares Alahdab²⁰, Fahad Mashhour Alanezi²¹, Turki M. Alanzi²², Kefyalew Addis Alene^{23,24}, Robert Kaba Alhassan²⁵, Vahid Alipour^{26,27}, Amir Almasi-Hashiani²⁸, Nelson Alvis-Guzman^{29,30}, Edward Kwabena Ameyaw¹¹, Saeed Amini³¹, Dickson A. Amugsi³², Robert Ancuceanu³³, Davood Anvari^{34,35}, Seth Christopher Yaw Appiah^{36,37}, Jalal Arabloo²⁶, Olatunde Aremu³⁸, Mulusew A. Asemahagn³⁹, Mohammad Asghari Jafarabadi^{40,41}, Atalel Fentahun Awedew⁴², Beatriz Paulina Ayala Quintanilla^{43,44}, Martin Amogre Ayanore^{45,46}, Yared Asmare Aynalem⁴⁷, Samad Azari⁴⁸, Zelalem Nigussie Azene⁴⁹, B. B. Darshan⁵⁰, Tesleem Kayode Babalola^{51,52}, Atif Amin Baig⁵³, Maciej Banach^{54,55}, Till Winfried Bärnighausen^{56,57}, Arielle Wilder Bell^{58,59}, Akshaya Srikanth Bhagavathula^{60,61}, Nikha Bhardwaj⁶², Pankaj Bhardwaj^{63,64}, Kritika Bhattacharyya^{65,66}, Ali Bijani⁶⁷, Zebeay Workneh Bitew^{68,69}, Somayeh Bohloul⁷⁰, Obasanjo Afolabi Bolarinwa⁵¹, Archith Boloor⁷¹, Ivana Bozicevic^{72,73}, Zahid A. Butt^{74,75}, Rosario Cárdenas⁷⁶, Felix Carvalho⁷⁷, Jaykaran Charan⁷⁸, Vijay Kumar Chattu^{79,80}, Mohiuddin Ahsanul Kabir Chowdhury^{81,82}, Dinh-Toi Chu⁸³, Richard G. Cowden⁸⁴, Saad M. A. Dahlawi⁸⁵, Giovanni Damiani^{86,87}, Eugene Kofuor Maafo Darteh⁸⁸, Aso Mohammad Darwesh⁸⁹, José das Neves^{90,91}, Nicole Davis Weaver¹, Diego De Leo⁹², Jan-Walter De Neve⁵⁶, Kebede Deribe^{93,94}, Keshab Deuba^{95,96}, Samath Dharmaratne^{1,97,98}, Mostafa Dianatinasab^{99,100}, Daniel Diaz^{101,102}, Alireza Didarloo¹⁰³, Shirin Djalalinia¹⁰⁴, Fariba Dorostkar¹⁰⁵, Eleonora Dubljanin¹⁰⁶, Bereket Duko^{107,108}, Maha El Tantawi¹⁰⁹, Shaimaa I. El-Jaafary¹¹⁰, Babak Eshraty¹¹¹, Sharareh Eskandarieh¹¹², Oghenowede Eyawo¹¹³, Ifeanyi Jude Ezeonwumelu^{114,115}, Sayeh Ezzikouri¹¹⁶, Farshad Farzadfar¹¹⁷, Nazir Fattahi¹¹⁸, Nelsensius Klau Fauk^{119,120}, Eduarda Fernandes¹²¹, Irina Filip^{122,123}, Florian Fischer¹²⁴, Nataliya A. Foigt¹²⁵, Masoud Foroutan^{126,127}, Takeshi Fukumoto¹²⁸, Mohamed M. Gad^{129,130}, Abhay Motiramji Gaidhane¹³¹, Birhan Gebresilassie Gebregiorgis⁴⁷, Ketema Bizuwork Gebremedhin¹³², Lemma Getacher¹³³, Keyghobad Ghadiri^{134,135}, Ahmad Ghashghae¹³⁶, Mahaveer Golechha¹³⁷, Mohammed Ibrahim Mohialdeen Gubari¹³⁸, Harish Chander Gugnani^{139,140}, Rafael Alves Guimarães¹⁴¹, Mohammad Rifat Haider¹⁴², Arvin Haj-Mirzaian^{143,144}, Samer Hamidi¹⁴⁵, Abdiwahab Hashi¹⁴⁶, Soheil Hassanipour^{147,148}, Hadi Hassankhani^{149,150}, Khezhar Hayat^{151,152}, Claudiu Herteliu^{153,154}, Hung Chak Ho¹⁵⁵,

*Correspondence: ehaeuser@uw.edu

¹ Institute for Health Metrics and Evaluation, University of Washington, Seattle, WA, USA

Full list of author information is available at the end of the article



© The Author(s) 2022. **Open Access** This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit <http://creativecommons.org/licenses/by/4.0/>. The Creative Commons Public Domain Dedication waiver (<http://creativecommons.org/publicdomain/zero/1.0/>) applies to the data made available in this article, unless otherwise stated in a credit line to the data.

Ramesh Holla⁵⁰, Mostafa Hosseini^{156,157}, Mehdi Hosseinzadeh^{158,159}, Bing-Fang Hwang¹⁶⁰, Segun Emmanuel Ibitoye¹⁶¹, Olayinka Stephen Ilesanmi^{162,163}, Irena M. Ilic¹⁶⁴, Milena D. Ilic¹⁶⁵, Rakibul M. Islam¹⁶⁶, Chidozie C. D. Iwu¹⁶⁷, Mihajlo Jakovljevic^{168,169}, Ravi Prakash Jha^{170,171}, John S. Ji¹⁷², Kimberly B. Johnson¹, Nitin Joseph¹⁷³, Vasna Joshua¹⁷⁴, Farahnaz Joukar^{147,148}, Jacek Jerzy Jozwiak¹⁷⁵, Leila R. Kalankesh¹⁷⁶, Rohollah Kalhor^{177,178}, Naser Kamyari¹⁷⁹, Tanuj Kanchan¹⁸⁰, Behzad Karami Matin¹¹⁸, Salah Eddin Karimi¹⁸¹, Gbenga A. Kayode^{182,183}, Ali Kazemi Karyani¹¹⁸, Maryam Keramati¹⁸⁴, Ejaz Ahmad Khan¹⁸⁵, Gulfaraz Khan¹⁸⁶, Md Nuruzzaman Khan¹⁸⁷, Khaled Khatab^{188,189}, Jagdish Khubchandani¹⁹⁰, Yun Jin Kim¹⁹¹, Adnan Kisa^{192,193}, Sezer Kisa¹⁹⁴, Jacek A. Kopec^{195,196}, Soewarta Kosen¹⁹⁷, Sindhura Lakshmi Koulmane Laxminarayana¹⁹⁸, Ai Koyanagi^{199,200}, Kewal Krishan²⁰¹, Barthelémy Kuate Defo^{202,203}, Nuworza Kugbey²⁰⁴, Vaman Kulkarni¹⁷³, Manasi Kumar^{205,206}, Nithin Kumar¹⁷³, Dian Kusuma^{207,208}, Carlo La Vecchia²⁰⁹, Dharmesh Kumar Lal²¹⁰, Iván Landires^{211,212}, Heidi Jane Larson^{1,213}, Savita Lasrado²¹⁴, Paul H. Lee²¹⁵, Shanshan Li²¹⁶, Xuefeng Liu^{217,218}, Afshin Maleki^{219,220}, Preeti Malik^{221,222}, Mohammad Ali Mansournia¹⁵⁶, Francisco Rogerlândio Martins-Melo²²³, Walter Mendoza²²⁴, Ritesh G. Menezes²²⁵, Endalkachew Worku Mengesha²²⁶, Tuomo J. Meretoja^{227,228}, Tomislav Mestrovic^{229,230}, Andreea Mirica¹⁵³, Babak Moazen^{56,231}, Osama Mohamad²³², Yousef Mohammad²³³, Abdollah Mohammadian-Hafshejani²³⁴, Reza Mohammadpourhodki²³⁵, Salahuddin Mohammed^{236,237}, Shafiu Mohammed^{238,239}, Ali H. Mokdad^{1,98}, Masoud Moradi¹¹⁸, Paula Moraga²⁴⁰, Sumaira Mubarik²⁴¹, Getaneh Baye B. Mulu²⁴², Lillian Mwanri²⁴³, Ahamarshan Jayaraman Nagarajan^{244,245}, Mukhammad David Naimzada^{246,247}, Muhammad Naveed²⁴⁸, Javad Nazari²⁴⁹, Rawlance Ndejjo²⁵⁰, Ionut Negoii^{251,252}, Frida N. Ngalesoni²⁵³, Georges Nguéfac-Tsague²⁵⁴, Josephine W. Ngunjiri²⁵⁵, Cuong Tat Nguyen²⁵⁶, Huong Lan Thi Nguyen²⁵⁶, Chukwudi A. Nnaji^{257,258}, Jean Jacques Noubiap²⁵⁹, Virginia Nuñez-Samudio^{260,261}, Vincent Ebuka Nwatah^{262,263}, Bogdan Oancea²⁶⁴, Oluwakemi Ololade Odukoya^{265,266}, Andrew T. Olagunju^{267,268}, Babayemi Oluwaseun Olakunde²⁶⁹, Bolajoko Olubukunola Olusanya²⁷⁰, Jacob Olusegun Olusanya²⁷⁰, Ahmed Omar Bali²⁷¹, Obinna E. Onwujekwe²⁷², Orish Ebere Orisakwe²⁷³, Nikita Otstavnov²⁴⁶, Stanislav S. Otstavnov^{246,274}, Mayowa O. Owolabi^{275,276}, P. A. Mahesh²⁷⁷, Jagadish Rao Padubidri²⁷⁸, Adrian Pana^{153,279}, Ashok Pandey^{280,281}, Seithikurippu R. Pandi-Perumal²⁸², Fatemeh Pashazadeh Kan²⁸³, George C. Patton^{284,285}, Shrikant Pawar²⁸⁶, Emmanuel K. Peprah²⁸⁷, Maarten J. Postma^{288,289}, Liliana Preotescu^{290,291}, Zahiruddin Quazi Syed¹³¹, Navid Rabiee^{292,293}, Amir Radfar²⁹⁴, Alireza Rafiei^{295,296}, Fakher Rahim²⁹⁷, Vafa Rahimi-Movaghar²⁹⁸, Amir Masoud Rahmani²⁹⁹, Kiana Ramezanzadeh¹⁴³, Juwel Rana^{300,301}, Chhabi Lal Ranabhat^{302,303}, Sowmya J. Rao³⁰⁴, David Laith Rawar^{305,306}, Salman Rawar^{307,308}, Reza Rawassizadeh³⁰⁹, Lemma Demissie Regassa³¹⁰, Nima Rezaei^{311,312}, Aziz Rezapour²⁶, Mavra A. Riaz³¹³, Ana Isabel Ribeiro³¹⁴, Jennifer M. Ross^{315,1,316}, Enrico Rubagotti^{317,318}, Susan Fred Rumisha^{319,320}, Godfrey M. Rwegerera³²¹, Sahar Saeedi Moghaddam¹¹⁷, Rajesh Sagar³²², Biniyam Sahiledengle³²³, Maitreyi Sahu¹, Marwa Rashad Salem³²⁴, Hossein Samadi Kafil³²⁵, Abdallah M. Samy³²⁶, Benn Sartorius^{98,327,328}, Brijesh Sathian^{329,330}, Abdul-Aziz Seidu^{88,331}, Amira A. Shaheen³³², Masood Ali Shaikh³³³, Morteza Shamsizadeh³³⁴, Wondimeneh Shibabaw Shiferaw⁴⁷, Jae Il Shin³³⁵, Roman Shrestha³³⁶, Jasvinder A. Singh^{337,338}, Valentin Yurievich Skryabin³³⁹, Anna Aleksandrovna Skryabina³⁴⁰, Shahin Soltani¹¹⁸, Mu'awiyah Babale Sufiyan³⁴¹, Takahiro Tabuchi³⁴², Eyayou Girma Tadesse³⁴³, Nuno Taveira^{344,345}, Fisaha Haile Tesfay^{346,347}, Rekha Thapar¹⁷³, Marcos Roberto Tovani-Palone^{348,349}, Gebiyaw Wudie Tsegaye³⁵⁰, Chukwuma David Umeokonkwo³⁵¹, Bhaskaran Unnikrishnan³⁵², Jorge Hugo Villafañe³⁵³, Francesco S. Violante^{354,355}, Bay Vo³⁵⁶, Giang Thu Vu³⁵⁷, Yohannes Dibaba Wado³⁵⁸, Yasir Waheed³⁵⁹, Richard G. Wamai^{360,361}, Yanzhong Wang³⁶², Paul Ward³⁶³, Nuwan Darshana Wickramasinghe³⁶⁴, Katherine Wilson³⁶⁵, Sanni Yaya^{366,367}, Paul Yip^{368,369}, Naohiro Yonemoto^{370,371}, Chuanhua Yu²⁴¹, Mikhail Sergeevich Zastrozhin^{372,373}, Yunquan Zhang^{374,375}, Zhi-Jiang Zhang³⁷⁶, Simon I. Hay^{1,98}, Laura Dwyer-Lindgren^{1,98} and on behalf of Local Burden of Disease sub-Saharan Africa HIV Prevalence

Abstract

Background: Human immunodeficiency virus and acquired immune deficiency syndrome (HIV/AIDS) is still among the leading causes of disease burden and mortality in sub-Saharan Africa (SSA), and the world is not on track to meet targets set for ending the epidemic by the Joint United Nations Programme on HIV/AIDS (UNAIDS) and the United Nations Sustainable Development Goals (SDGs). Precise HIV burden information is critical for effective geographic and epidemiological targeting of prevention and treatment interventions. Age- and sex-specific HIV prevalence estimates are widely available at the national level, and region-wide local estimates were recently published for adults overall. We add further dimensionality to previous analyses by estimating HIV prevalence at local scales, stratified into sex-specific 5-year age groups for adults ages 15–59 years across SSA.

Methods: We analyzed data from 91 seroprevalence surveys and sentinel surveillance among antenatal care clinic (ANC) attendees using model-based geostatistical methods to produce estimates of HIV prevalence across 43 countries in SSA, from years 2000 to 2018, at a 5 × 5-km resolution and presented among second administrative level (typically districts or counties) units.

Results: We found substantial variation in HIV prevalence across localities, ages, and sexes that have been masked in earlier analyses. Within-country variation in prevalence in 2018 was a median 3.5 times greater across ages and sexes, compared to for all adults combined. We note large within-district prevalence differences between age groups: for men, 50% of districts displayed at least a 14-fold difference between age groups with the highest and lowest prevalence, and at least a 9-fold difference for women. Prevalence trends also varied over time; between 2000 and 2018, 70% of all districts saw a reduction in prevalence greater than five percentage points in at least one sex and age group. Meanwhile, over 30% of all districts saw at least a five percentage point prevalence increase in one or more sex and age group.

Conclusions: As the HIV epidemic persists and evolves in SSA, geographic and demographic shifts in prevention and treatment efforts are necessary. These estimates offer epidemiologically informative detail to better guide more targeted interventions, vital for combating HIV in SSA.

Keywords: HIV, Mapping, Africa, Geostatistics, Spatial statistics, HIV prevalence, Demographics

Background

Four decades after its discovery, human immunodeficiency virus (HIV) continues to impact millions of people worldwide, remains one of the leading causes of morbidity and mortality globally [1, 2] and incurs billions of dollars annually in direct health care costs and indirect socioeconomic costs [3]. In sub-Saharan Africa (SSA) in 2019, an estimated 26 million people were living with HIV [2]. In recent years, international bodies have set goals to end the HIV epidemic: in 2014, the Joint United Nations Programme on HIV/AIDS (UNAIDS) introduced the “95-95-95” targets—that by 2030, 95% of people living with HIV globally would know their status, 95% of all people with diagnosed HIV infection would receive sustained antiretroviral therapy, and 95% of people living with HIV receiving antiretroviral therapy (ART) would be virally suppressed [4, 5]. The United Nations Sustainable Development Goals also call for an end to the AIDS epidemic by 2030 [6]. Unfortunately, despite a significant increase in ART coverage over the last 20 years and major progress in terms of reductions in HIV incidence and mortality [1], the latest estimates and projections indicate that

the world is not on track to meet these goals [2, 7, 8], and progress may stall further as a consequence of the COVID-19 pandemic [9].

Differences in HIV prevalence both within and between nations in SSA have been well-documented [10–14], as have differences between sexes [2, 12–14] and age groups [2]. These differences have also changed over time [1, 10], impacted in part by the onset, duration, location, and demographic targeting of different prevention and treatment interventions [15–17]. Epidemiologically targeted interventions are understood to be more effective compared to homogeneous interventions [18] and are increasingly important at a time when the future of funding for HIV prevention and treatment is both uncertain and highly variable [19, 20], particularly in the wake of disruptions related to the COVID-19 pandemic [21]. Evidence suggests that interventions are most effective when tailored to account for differences in the intensity of the epidemic by geographic location [14, 22], sex [23], and age [24]. Locally and demographically precise HIV prevalence information, however, is necessary in order to maximize the benefit of such methods; at present, such information in SSA is lacking.

HIV prevalence estimates stratified by age and sex are available at the national level through the Global Burden of Disease (GBD) [2] and from UNAIDS [25]. Both sources also provide subnational estimates at the first administrative level (e.g., province, state) in select countries. Recently, Dwyer-Lindgren et al. [10] presented aggregated adult HIV prevalence estimates for the years 2000–2017 at local scales in SSA, generalizing estimates for males and females combined, and across ages 15–49 years. Some studies have gone further to present subnational prevalence estimates separated by sex [26–29] or age [30]; however, these studies focused on single countries, and/or presented estimates for only one point in time, without describing any temporal trajectories in prevalence. To our knowledge, no previous studies have presented age- and sex-specific HIV prevalence estimates across SSA at local scales over time.

We built upon the HIV prevalence model from Dwyer-Lindgren et al. [10] to produce HIV prevalence estimates for 43 countries in SSA for males and females ages 15–59 years, stratified into nine 5-year age groups, for the years spanning 2000 to 2018. Countries, age groups, and time period were selected according to data availability. We expanded upon existing Bayesian spatiotemporal methods to model these estimates at a 5×5 -km resolution and present them here aggregated to the second administrative level (which varies by country but is typically equivalent to e.g., districts, municipalities), which is the level typically considered most relevant to policymakers and stakeholders. Prevalence estimates for all demographic groups at all levels of geographic aggregation, as well as number of people living with HIV (count estimates), are publicly available from the Global Health Data Exchange (<https://ghdx.healthdata.org/record/ihme-data/sub-saharan-africa-hiv-prevalence-geospatial-estimates-2000-2018>) and through a user-friendly data visualization tool (<http://vizhub.healthdata.org/lbd/hiv-prev-disagg>).

Methods

Overview

This ecological study follows the Guidelines for Accurate and Transparent Health Estimates Reporting (GATHER) [31] (Additional file 1: Section 1). This analysis relies secondary data sources to provide estimates of HIV prevalence on a 5×5 -km grid in 43 countries in SSA for males and females ages 15–59 years residing at each location, stratified into five-year age bins (i.e., ages 15–19, 20–24, 25–29, 30–34, 35–39, 40–44, 45–49, 50–54, 55–59), with annual resolution from year 2000 to 2018 inclusive, calibrated to national estimates from the GBD [2]. The period of 2000–2018 and the age range of 15–59 years were selected to

optimize the contemporaneousness of the estimates and to account for data availability—there were relatively few large-scale seroprevalence surveys conducted before 2000, and most seroprevalence surveys focus on adults, with little reporting outside the 15–59 years age range. We produced estimates for sex rather than gender binaries because sex is more predominantly reported in the available data sources. Due to data availability limitations we were unable to produce prevalence estimates for sex minority individuals outside the male/female binary. The 43 countries analyzed were also selected according to data availability—Mauritania was excluded as there were no HIV prevalence data available. We included six countries—Djibouti, Guinea-Bissau, Madagascar, Somalia, South Sudan and Sudan—where no seroprevalence survey data were available, but where sentinel surveillance data collected from antenatal care clinic (ANC) attendees (described below) were available. The implications of these and other limitations are expanded upon in the “Methodological advantages and limitations” section in the “Discussion” section.

The methodology used here largely parallels that previously used to map adult HIV prevalence in SSA [10], with the incorporation of modifications necessary to model by age and sex, and improvements related to the inclusion of spatially aggregated data and ANC data (Fig. 1). We used a 5×5 -km grid for consistency with this previous analysis; to align with the resolution available for pre-existing covariates incorporated in this analysis; and for flexibility in aggregating these estimates to other levels of interest (e.g., first- and second-level administrative subdivisions, such as states or districts, respectively, or more aggregated age groups such as reproductive ages [commonly 15–49]) using grid-cell-level estimates of age- and sex-specific population from Worldpop [32]. These population estimates were also used to estimate the number of people living with HIV in each demographic group. All analyses were conducted in R version 3.6.1 [33]. Figure 2 provides an overview of the analytic process, described in more depth below. Additional details are available in Additional file 1.

Data

HIV data

We compiled a geolocated dataset of 304,672 observations from 91 seroprevalence surveys from 37 countries and 10,351 observations from sentinel surveillance among antenatal care clinic attendees (ANC data) in 43 countries (Additional file 2: Tables S1–S2; Fig. 1). Data from seroprevalence surveys were originally in the form of survey microdata (that is, individual-level survey responses) or survey reports (Additional file 2:

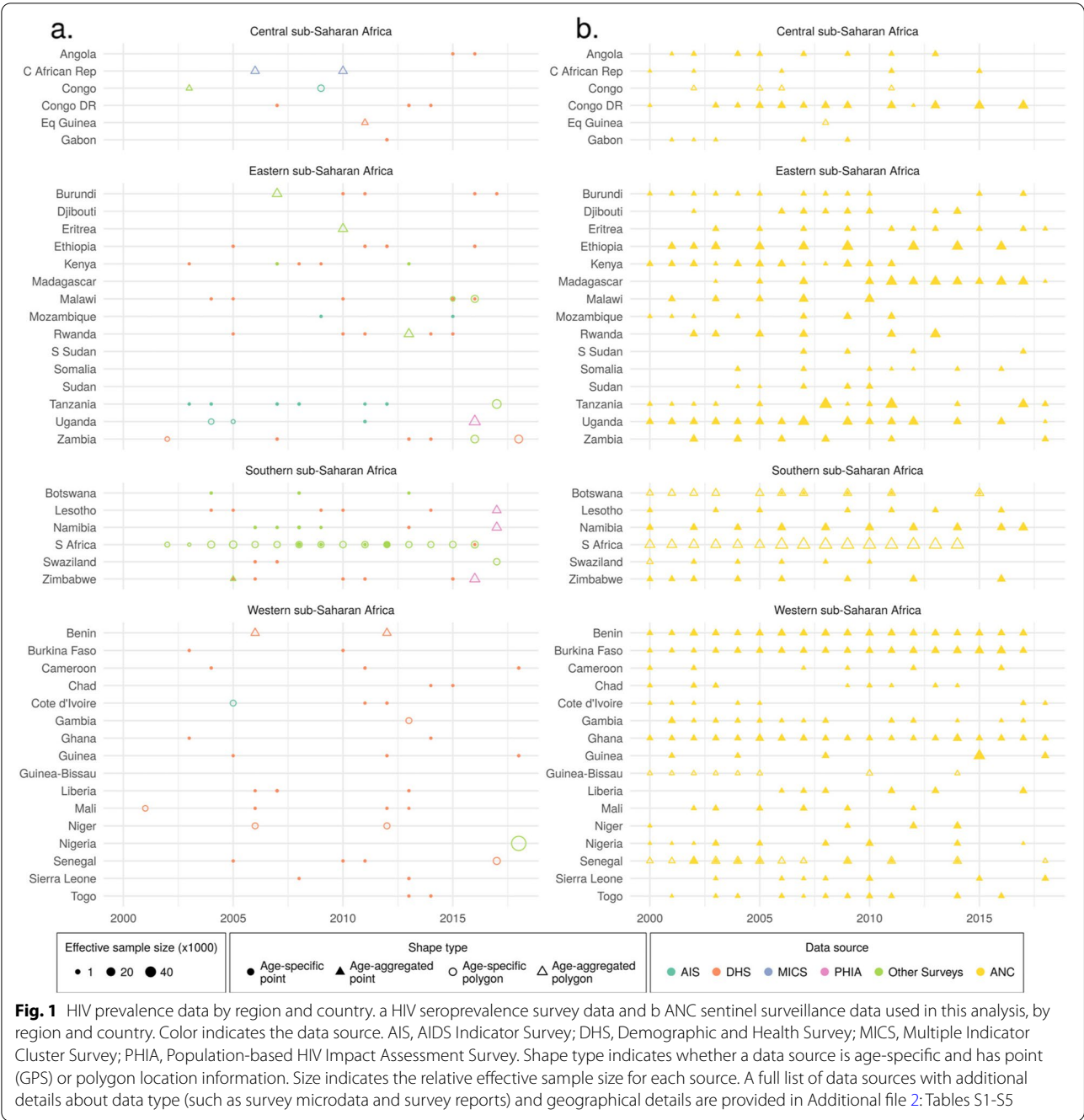
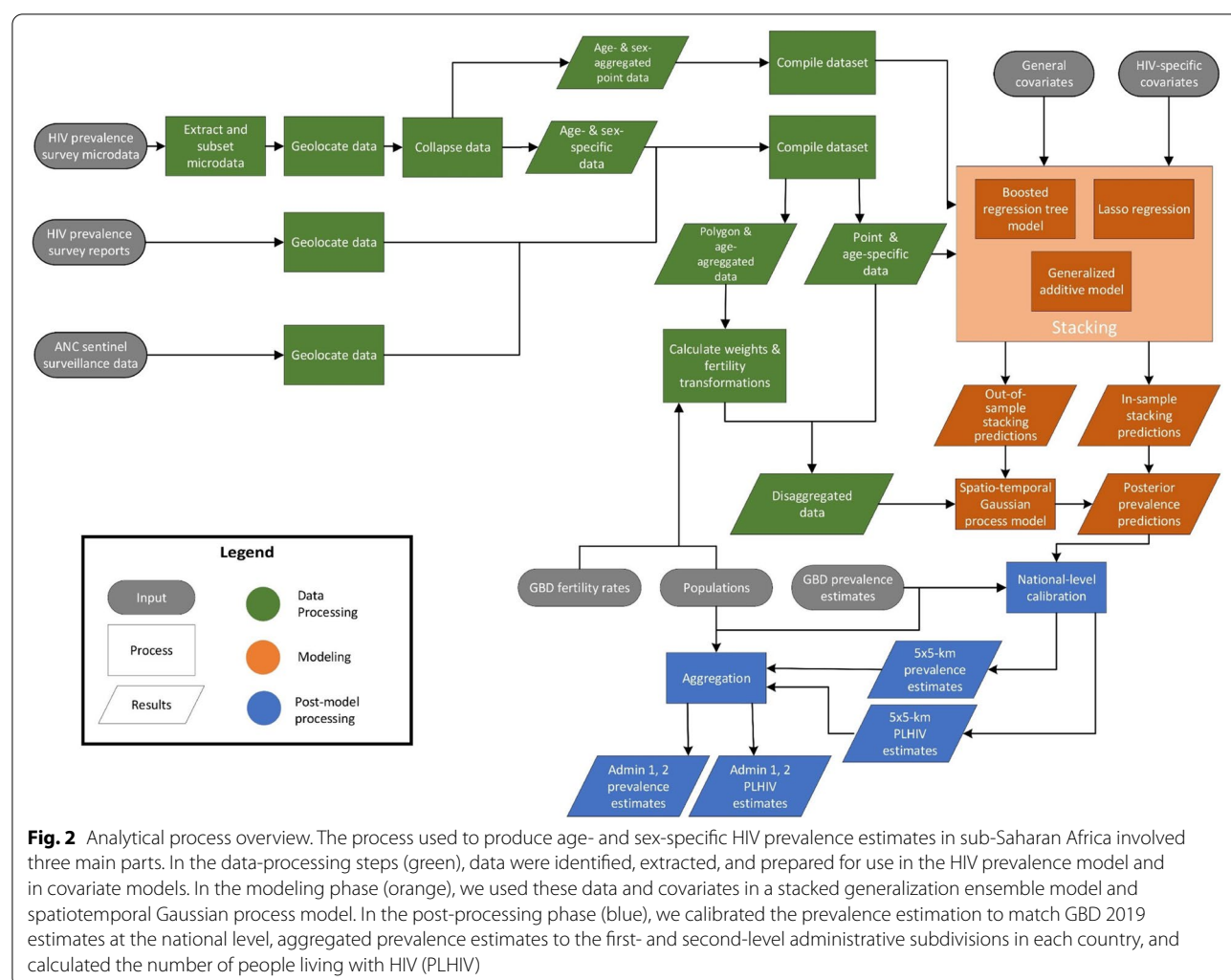


Table S1). For surveys with available microdata, we extracted variables related to age, sex, HIV blood test result, location, and year, as well as survey weights, where available. We excluded rows with missing information on any of these variables, and subset the data to ages 15–59 years. For data coded by gender rather than sex, we treated these data as if they were sex-specific rather than gender-specific. We recognize that sex and gender are not interchangeable: sex is a biological variable, while gender is a fluid social construct. In the absence of quality data, however, we could not disaggregate estimates by gender at this time. After subsetting by age, we collapsed the age-specific data into 5-year age bins (hereafter referred to as “ages”) by sex. We did this by calculating the weighted age- and sex-specific HIV prevalence at the finest spatial resolution available. Ideally, this was at the level of global positioning system (GPS) coordinates that represent the location of a survey cluster. In most



surveys, GPS coordinates are randomly displaced (typically by 2–5 km depending on the setting and the survey series [34]) in order to protect respondent's confidentiality. In instances where GPS coordinates were not available, the smallest areal unit (termed a “polygon”) possible was used instead. These typically represented an administrative subdivision. For surveys without microdata but for which estimates with some subnational resolution were provided in a report, we extracted these estimates with information about the sample size and location. GPS coordinates were not available for these reports, so these data were exclusively matched to polygons. In most reports, age ranges larger than 5 years were reported. Among these, we retained data reported for age ranges that corresponded exactly to one or more of the 5-year age bins used in this model; for example, we included surveys covering age ranges 15–49 years, or 15–24 years, but excluded those covering age ranges such as 18–24 years. For age-aggregated data, we retained

information regarding the age range covered, to be used in our modeling process as described below. We also only included sex-specific data. For more information on excluded surveys see Additional file 2: Table S3.

Data that were spatially aggregated (i.e., polygon data) and/or age-aggregated required additional processing. Although we ultimately modeled HIV prevalence at the level of the observation, be it point or polygon, age-specific or age-aggregated, our modeling process initially specified HIV prevalence at the point-, time-, age-, and sex-specific level. Because of this, it was necessary that we disaggregate the age-aggregated and polygon survey data to be location- and age-specific. We did this by distributing polygon data to pixels proportional to population. Specifically, for each polygon, we generated points at the centroid of each 5×5 -km pixel falling within that polygon and replicated that observation's HIV prevalence and sample size at the location of each of those centroids. Age-aggregated point data were

similarly disaggregated by replicating the HIV prevalence and sample size once for each year-age group covered in the overall age range. In the cases of age-aggregated polygon data, these two processes were combined. Next, each of the disaggregated, location- and age-specific rows of data associated with a given aggregated observation were assigned weights proportional to the age- and sex-specific population residing at that location for the given year, derived from WorldPop [32]. Weights per observation all summed to one. This process substantially increased the size of the dataset. To reduce the associated computational burden when fitting the model, in cases where at least one row within an observation was given a weight of less than half of one divided by the number of locations and/or ages in that observation, we successively dropped the lowest-weighted locations and/or ages until reaching a maximum of 1% of the observation's weight dropped. Remaining locations and/or ages within that observation were then reweighted to maintain a total weight of one. Data that were not aggregated (i.e., age-specific point observations) were each assigned a weight of one.

ANC data were primarily derived from national HIV estimate files developed by national teams and compiled and shared via UNAIDS [35] and supplemented with data derived from sentinel surveillance country reports (Additional file 2: Table S2). We extracted information from these sources on HIV prevalence and sample size by site and year. Sites were geolocated to specific GPS coordinates where possible and otherwise to a polygon that represents an administrative subdivision. The ANC data available for this analysis were not age-specific. Because ANC data included only pregnant females, we assumed the age range of these data to be that of females with non-zero fertility rates in SSA according to GBD 2019 [36], that is, females ages 15–54 years. We disaggregated ANC data to the age and location level as we did for age-aggregated or polygon survey data. However, specific locations and ages were weighted by number of births rather than population size. The number of births for a given age and location was estimated as the product of the location-, age-, and sex-specific population, again derived from WorldPop [32], and the national fertility rate, derived from GBD 2019 estimates [36].

Covariates

This analysis included the same covariates as the previous analysis [10]. This included five pre-existing covariates: (1) travel time to the nearest settlement of more than 50,000 inhabitants; (2) total population; (3) night-time lights; (4) urbanicity; and (5) malaria incidence (Additional file 2: Table S4). In addition, eight

covariates were constructed explicitly for this analysis owing to their known association with HIV prevalence and data availability: (1) prevalence of male circumcision (all forms); (2) prevalence of self-reported sexually transmitted infection (STI) symptoms; (3) prevalence of marriage or living with a partner as married; (4) prevalence of one's current partner living elsewhere among females; (5) prevalence of condom use at last sexual encounter; (6) prevalence of reporting ever having had intercourse among young females; and (7) and (8) prevalence of multiple partners in the past year for males and for females, respectively. We updated the covariates constructed for this analysis to incorporate newly available data but utilized the original statistical methods (Additional file 1: Section 3.2; Additional file 2: Table S5; Additional file 3: Figs. S1–S8).

Model and estimation

Covariate stacking

An ensemble covariate modeling approach (“stacking”) was implemented to capture possible nonlinear interactions among the covariates across space and time [37]. In this approach, three sub-models were fitted to the HIV survey data with the covariates as explanatory predictors: generalized additive models [38], boosted regression trees [39], and lasso regression [40]. Each sub-model was fitted using fivefold cross-validation to avoid overfitting, and the out-of-sample predictions from across the five folds were compiled into a single set of predictions that were used to fit the geostatistical model described below. In addition, each sub-model was also fitted to the full dataset to generate a complete set of in-sample predictions that were subsequently used when generating predictions from the geostatistical model (Additional file 3: Figs. S9–S11). Because the covariates used here were neither age-specific nor (for most) sex-specific, we fit these sub-models at that same age- and sex-aggregated level as the HIV-specific covariates, modeling HIV prevalence data aggregated across ages 15–49 and males and females. The age range 15–49 years was used in this case because of its more common usage in seroprevalence surveys compared to the 15–59 years range, allowing us to retain more data for the stacking model. Polygon data were excluded from stacking models due to their incongruity with the configurations needed for the different sub-models. The ANC data were also excluded due to known sampling biases, which are described in the Additional file 1: Section 4.2.

Geostatistical model

This model was fit in Template Model Builder (TMB) [41]. Owing to computational constraints, and to

allow for regional differences in the relationships between covariates and HIV prevalence, as well as differences in the temporal, spatial, and demographic autocorrelation in HIV prevalence, separate models were fitted for four regions (Additional file 3: Fig. S12). We modeled HIV prevalence stratified by space, time, age, and sex using a generalized linear mixed-effects model. To simultaneously model point- and polygon-level observations, as well as both age-specific and age-aggregated observations, we specified the data likelihood at the observation level (i), which accommodated all of these. We modeled the number of HIV-positive individuals (Y_i) among a sample (N_i) for a given observation as a binomial variable:

$$Y_i \sim \text{Binomial}(N_i, p_i)$$

Logit-transformed prevalence was however first specified at the space, time, age, and sex-disaggregated level (j):

$$\begin{aligned} \text{logit}(p_j) &= \beta_0 + \beta_1 X_j + Z_{1,j} + Z_{2,j} + Z_{3,c[j]} \\ Z_{1,j} &\sim \text{GP}(0, \Sigma_{1,space} \otimes \Sigma_{1,time}) \\ Z_{2,j} &\sim \text{GMRF}(0, \Sigma_{2,time} \otimes \Sigma_{2,age} \otimes \Sigma_{2,sex}) \\ Z_{3,c[j]} &\sim \text{GMRF}(0, \Sigma_{3,c}) \end{aligned}$$

We specified logit-transformed prevalence at the disaggregated level (p_j) as a linear combination of:

- A regional intercept (β_0);
- Covariates and associated regression parameters ($\beta_1 X_j$);
- Random effects correlated across space and time, ($Z_{1,j}$);
- Random effects correlated across time, age, and sex, ($Z_{2,j}$);
- Country-specific (c) random effects correlated across age, ($Z_{3,c[j]}$).

The random effects capturing correlations between space, time, age, and sex included:

- $Z_{1,j}$: a Gaussian process with mean 0 and a covariance matrix given by the Kronecker product of a spatial Matérn covariance function [42] ($\Sigma_{1,space}$) and a temporal first-order autoregressive covariance function ($\Sigma_{1,time}$);
- $Z_{2,j}$: a Gaussian Markov Random Field with mean 0 and a covariance matrix given by the Kronecker product of first-order autoregressive covariance functions for time ($\Sigma_{2,time}$), age ($\Sigma_{2,age}$), and sex ($\Sigma_{2,sex}$);
- $Z_{3,c[j]}$: a Gaussian Markov Random Field with mean 0 and a covariance matrix given by country-specific first-order autoregressive covariance functions for age ($\Sigma_{3,c}$).

We used the stochastic partial differential equation [43] approach to approximate the continuous spatiotemporal Gaussian random field ($Z_{1,j}$). Sensitivity analyses were carried out to compare this model configuration to others with differing p_j specification configurations, as well as to several other model and data specifications, and are described in detail in the Additional file 1: Section 4.3, Additional file 3: Figs. S13–S15, and the “Discussion” section. We then specified observation-level (i) prevalence:

$$p_i = \text{logit}^{-1} \left(\text{logit} \left(\sum (p_{transformed,j} \cdot w_j) \right) + (\beta_2 + U_{s[i]}) \cdot I_{ANC} + \epsilon_i \right)$$

p_i was calculated as the sum of disaggregated prevalence ($p_{transformed,j}$) estimates multiplied by their respective population (or in the case of ANC data, birth) weights (w_j), plus the incorporation of additional ANC-related transformations and bias corrections (β_2 , $U_{s[i]}$, and I_{ANC} described below), and an observation-level uncorrelated error term (ϵ_i):

$$\epsilon_i \sim \text{Normal}(0, \sigma_i^2)$$

In cases where data were already disaggregated spatially and by age, $w_j = 1$.

HIV prevalence as measured by sentinel surveillance of ANC clinic attendees is known to be biased as a measure of HIV prevalence in the general adult female population [44], because it only covers pregnant females who attend ANC, compared to all adult females [45, 46]. Additionally, fertility rates differ between HIV⁺ and HIV⁻ females, with the exact relationship varying by age [47], thereby impacting age-specific ANC clinic visitation rates. To address this, for ANC data we transformed prevalence among pregnant females based on the underlying prevalence among all females and the age-specific fertility-rate ratio (HIV⁺ fertility/HIV⁻ fertility). For ANC data,

$$p_{transformed,j} = \frac{(p_j \cdot FRR_j)}{(p_j \cdot FRR_j) + 1 - p_j}$$

Fertility rate ratios (FRR_j) were derived from GBD 2019 fertility estimates [36], taken at the national level except in cases where subnational estimates were available (in Ethiopia, Nigeria, and South Africa). For survey data,

$$p_{transformed,j} = p_j$$

To allow for additional ANC-related bias at the observation level (i), in instances where data in our model were derived from ANC sentinel surveillance (where $I_{ANC} = 1$ for ANC data, and $I_{ANC} = 0$ for all other data) our model incorporated a fixed term (β_2) that captured overall mean

bias in the ANC data, and a random effect ($U_{s[i]}$) for a given ANC site s that captured spatial differences in the extent of this bias:

$$U_{s[i]} \sim \text{Normal}(0, \sigma_{site[i]}^2)$$

Fitted model parameters are detailed in Additional file 2: Table S6. From each fitted model, we generated 1000 draws from the approximated joint posterior distribution of all model parameters and used these to construct 1000 draws of p_j , setting I_{ANC} to 0. Fivefold cross-validation was used to assess model performance and to compare a number of alternative models (Additional file 3: Figs. S13–S15). We also compared the re-aggregated adult-level estimates from our final model to those from the results of an age- and sex-aggregated counterpart (Additional file 3: Fig. S16).

Post-estimation

To take advantage of the more structured modeling approach and additional national-level data used by GBD 2019 [2], we performed post hoc calibration of our estimates to the corresponding national-level GBD estimates. For each country, year, age bin, and sex in our analysis, we defined a “raking factor” equal to the ratio of the GBD estimate for this country-year-age-sex to the population-weighted posterior mean HIV prevalence in all corresponding grid cells (Additional file 3: Figs. S17–S18). These raking factors were then used to scale each draw of HIV prevalence for each grid cell within that GBD geography, year, age, and sex. Point estimates for each grid cell were calculated as the mean of the scaled draws, and 95% uncertainty intervals were calculated as the 2.5th and 97.5th percentiles of the scaled draws. Grid cells that crossed international borders within modeling regions were fractionally allocated to multiple countries in proportion to the covered area during this process. In cases where subnational (i.e., first administrative level) estimates were available from the GBD, that is, for Ethiopia, Nigeria and South Africa, we calibrated to those estimates rather than those at the national level. Uncertainty in GBD estimates was not accounted for in this calibration.

In addition to estimates of HIV prevalence on a 5 × 5-km grid, we constructed estimates of HIV prevalence for first- and second-level administrative subdivisions. We did this by calculating age- and sex-specific population-weighted averages of prevalence for all grid cells within a given area. This process was carried out for each of the 1000 posterior draws (after calibration to GBD), with final point estimates derived from the mean of these draws and uncertainty intervals from the 2.5th and 97.5th percentiles. Additionally, estimates of the

number of people living with HIV for a given age and sex in each grid cell were derived by multiplying estimated prevalence in each grid cell by the corresponding population estimate from WorldPop [32], which was also calibrated to match GBD 2019 [36] (Additional file 1: Section 4.4; complete estimates of people living with HIV are available along with all prevalence estimates at (<https://ghdx.healthdata.org/record/ihme-data/sub-saharan-africa-hiv-prevalence-geospatial-estimates-2000-2018>)).

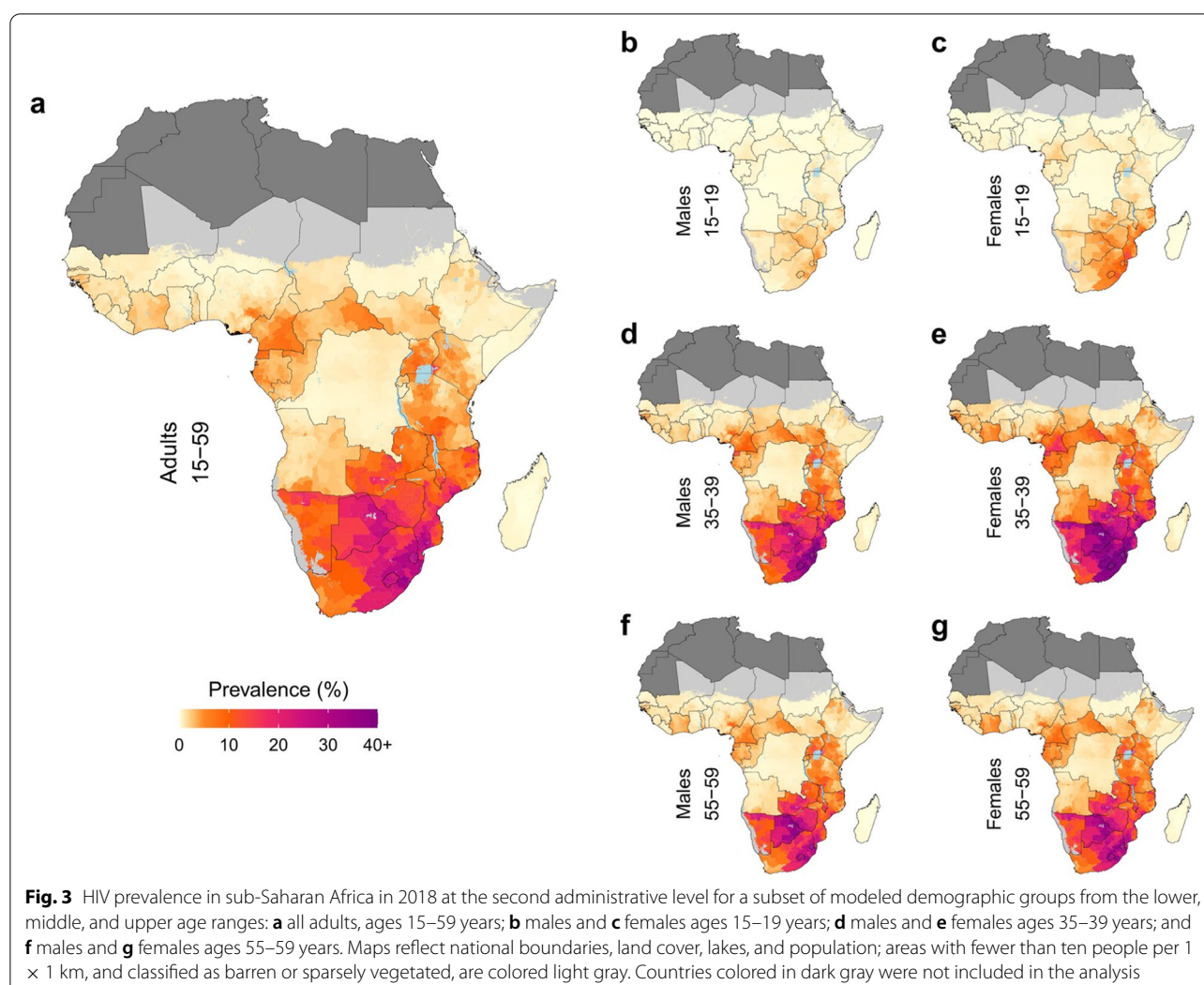
Although the model makes predictions for all locations covered by available covariates, all final model outputs for which land cover was classified as barren or sparsely vegetated according to European Space Agency Climate Change Initiative satellite data [48] and for which total population density was less than 10 individuals per 1 × 1-km in 2015 were masked for improved clarity when communicating with data specialists and policymakers. Maps were generated in R using the *ggplot2* [49] package version 3.3.0.

Results

Geographic variation

We found large differences in the spatial and demographic distribution of estimated HIV prevalence in SSA that were masked in demographically aggregated estimates (Figs. 3 and 4; Additional file 3: Figs. S19–S34). This was particularly striking among middle and older age groups. For example, in the year 2018, the maximum estimated HIV prevalence in any second-level administrative unit for adults ages 15–59 years was 35.4% in Umgungundlovu in the Kwazulu Natal province, South Africa (95% uncertainty interval (UI), 22.3–46.3%). However, estimated prevalence reached up to 59.4% [46.5–71.2%], almost 1.7 times higher, for females ages 35–39 years within that same location. Across all second-level administrative units, age groups, and sexes, females ages 35–39 in Nkilingo in Lubombo, Eswatini, had the highest estimated HIV prevalence in the year 2018, at 62.5% [50.1–74.5%].

Geographic variation within countries was also more dramatic in our demographically disaggregated results. Across SSA countries, the median absolute difference between second-level administrative units with the lowest and highest estimated prevalence within a given country in 2018 was 3.5 times greater when considered across ages and sexes, than when estimated for all adults combined (11.2 percentage points versus 3.2 percentage points). This difference in within-country prevalence range between demographically aggregated versus disaggregated estimates varied greatly between countries. For example, in Mozambique, this range across second-level administrative units was 30.1 percentage points [16.7–46.3] for combined adults and

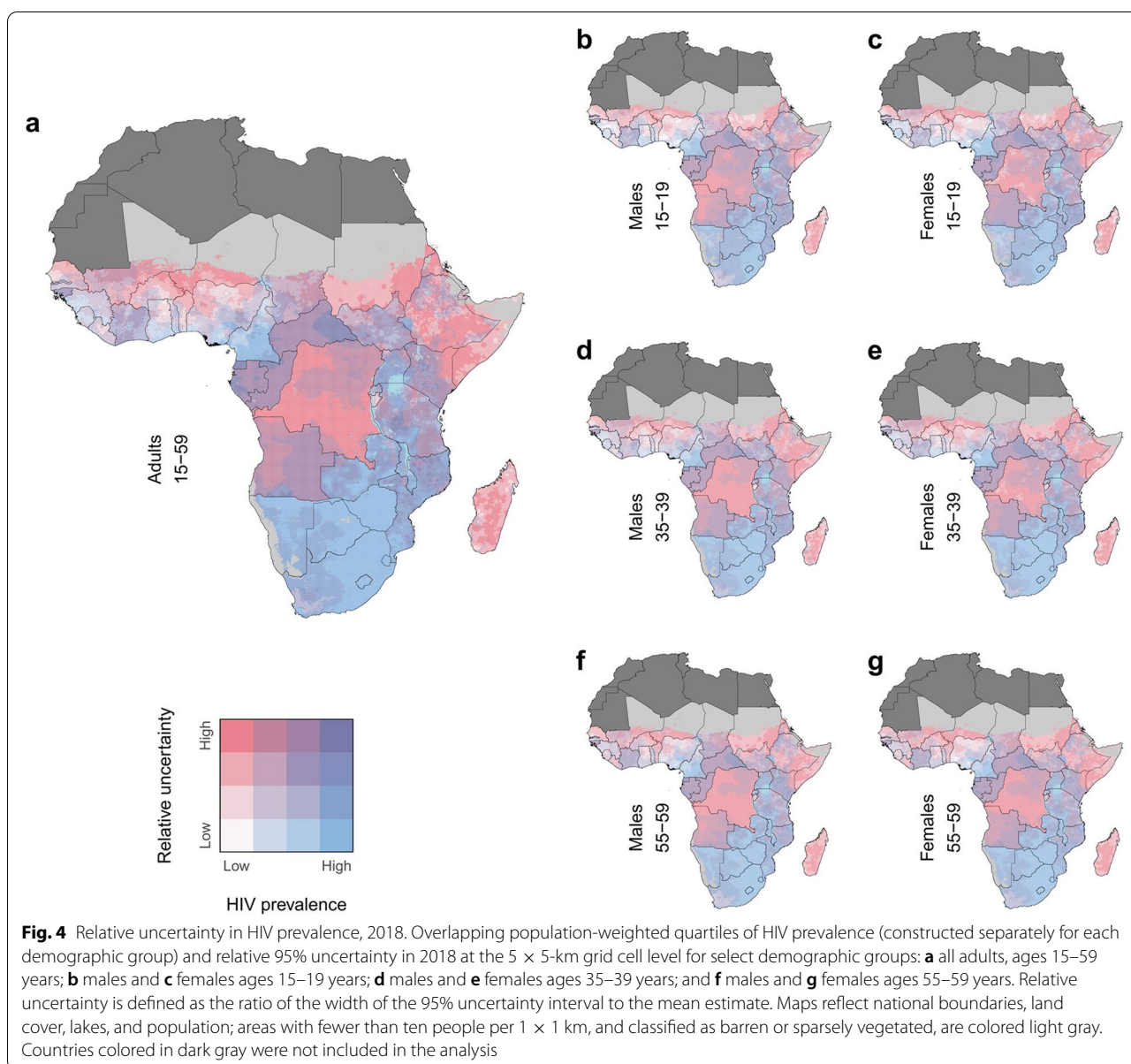


56.9 percentage points [37.4–78.2] (or 1.9 times larger) for estimates across ages and sexes. In Lesotho, on the other hand, this range was 8.2 times larger for estimates across ages and sexes compared to adults combined (51.6 percentage points [40.1–63.5] versus 6.3 percentage points [1.4–11.5]). Overall, countries in Eastern SSA tended to see greater such discrepancies compared to other regions; here, the median absolute difference between second-level administrative units was 4.4 times greater when considered across ages and sexes than for all adults combined (14.0 versus 3.2 percentage points). For complete geographic variation comparisons within each country, including uncertainty estimates, see Additional file 4.

Variation between males and females

Across SSA and across the years 2000–2018, estimated HIV prevalence was generally higher among females

than males (Fig. 5). In 2018, for prevalence aggregated across ages 15–59 years, in no second-level administrative units was estimated prevalence higher among males compared to females. The absolute difference in estimated prevalence in 2018 between females and males reached a maximum of 15.0 percentage points (in Umkhanyakude, in KwaZulu-Natal, South Africa, with 36.3% [24.7–46.8%] estimated prevalence in females compared to 21.3% [13.1–28.7%] estimated prevalence in males), for a female to male prevalence ratio of 1.7 [1.5–1.9]. Countries in Central SSA, where overall prevalence was lower than in other SSA regions, tended to see the largest disparity between females and males in terms of relative differences. Estimated prevalence among females in Central SSA ranged up to a maximum of 2.7 [1.84–4.2] times greater than estimated prevalence in males in 2018 (in San Antonio de Palé, in Annobón, Equatorial Guinea, with 8.3%

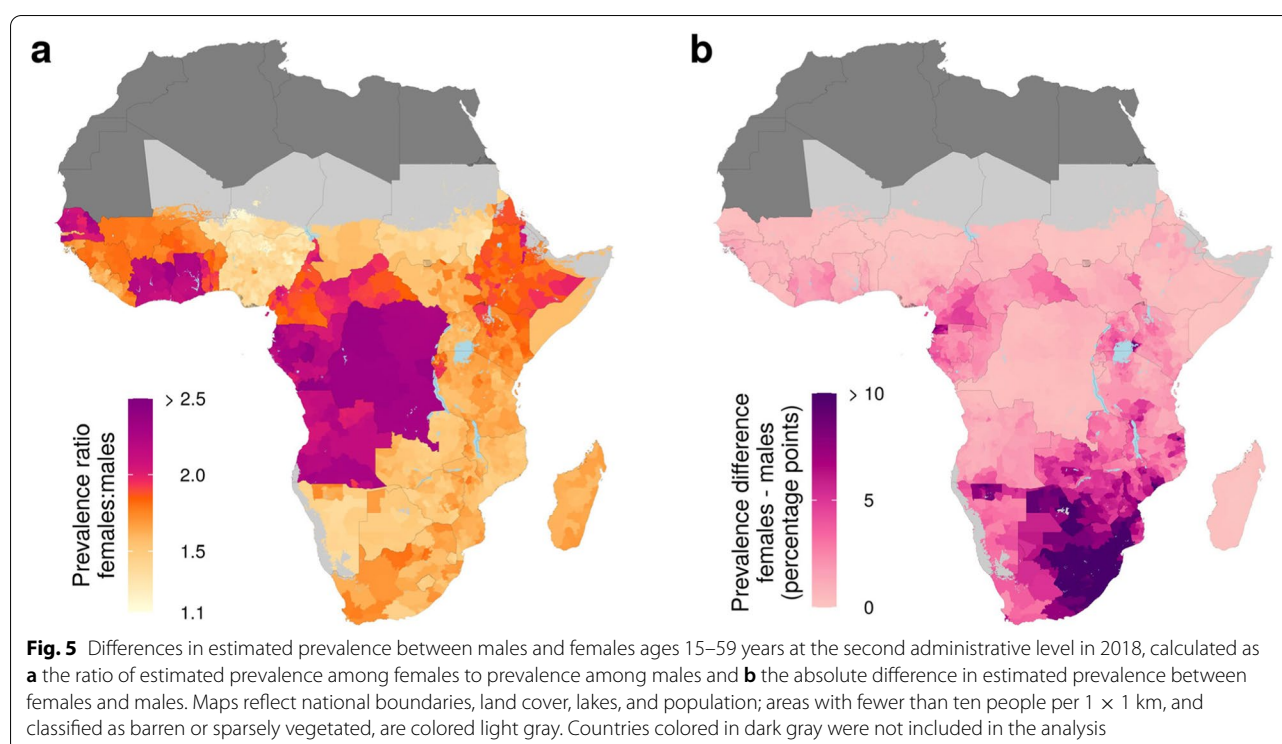


[2.1–21.4%] prevalence in females compared to 3.1% [0.8–8.1%] prevalence in males). Across Central SSA second-level administrative units, the median ratio between female and male estimated prevalence was 2.2, compared to the all-SSA median ratio of 1.6. The greatest absolute differences were seen in Eastern SSA, where the median absolute difference between female and male estimated prevalence was 1.9 percentage points in 2018, compared to the all-SSA median absolute difference of 0.9 percentage points. These differences between female and male prevalence in 2018 were less than those observed in the year 2000, when the median ratio between female and male estimated

prevalence was 1.5, and the median absolute difference was 1.5 percentage points. We did not note substantial differences in within-country variations in prevalence between females and males in either 2000 or 2018 in any region. For complete comparisons between sexes by second-level administrative unit, including uncertainty estimates, see Additional file 4.

Variation between age groups

Prevalence within second-level administrative units was also highly variable across age groups (Fig. 6), and relative variation in prevalence between age groups in 2018 tended to be higher in males. Comparing estimated



prevalence across age groups within a given second-level administrative unit in 2018, the ratio between highest and lowest prevalence among age groups tended to be larger among males compared to females (median ratio across all SSA second-level administrative units of 14.4 for males, and 9.3 for females). For males, this ratio between highest and lowest estimated prevalence among age groups was smaller in Central SSA compared to other regions (median ratio of 8.3) and was largest in Western SSA (median ratio of 21.7). There was little regional difference for females. The sexes also differed in changes in this ratio between years, where it decreased over time for males (with a median ratio in 2000 of 52.7) but increased over time for females (median ratio in 2000 of 5.6). For complete age variation comparisons by second-level administrative unit, including uncertainty estimates, see Additional file 4.

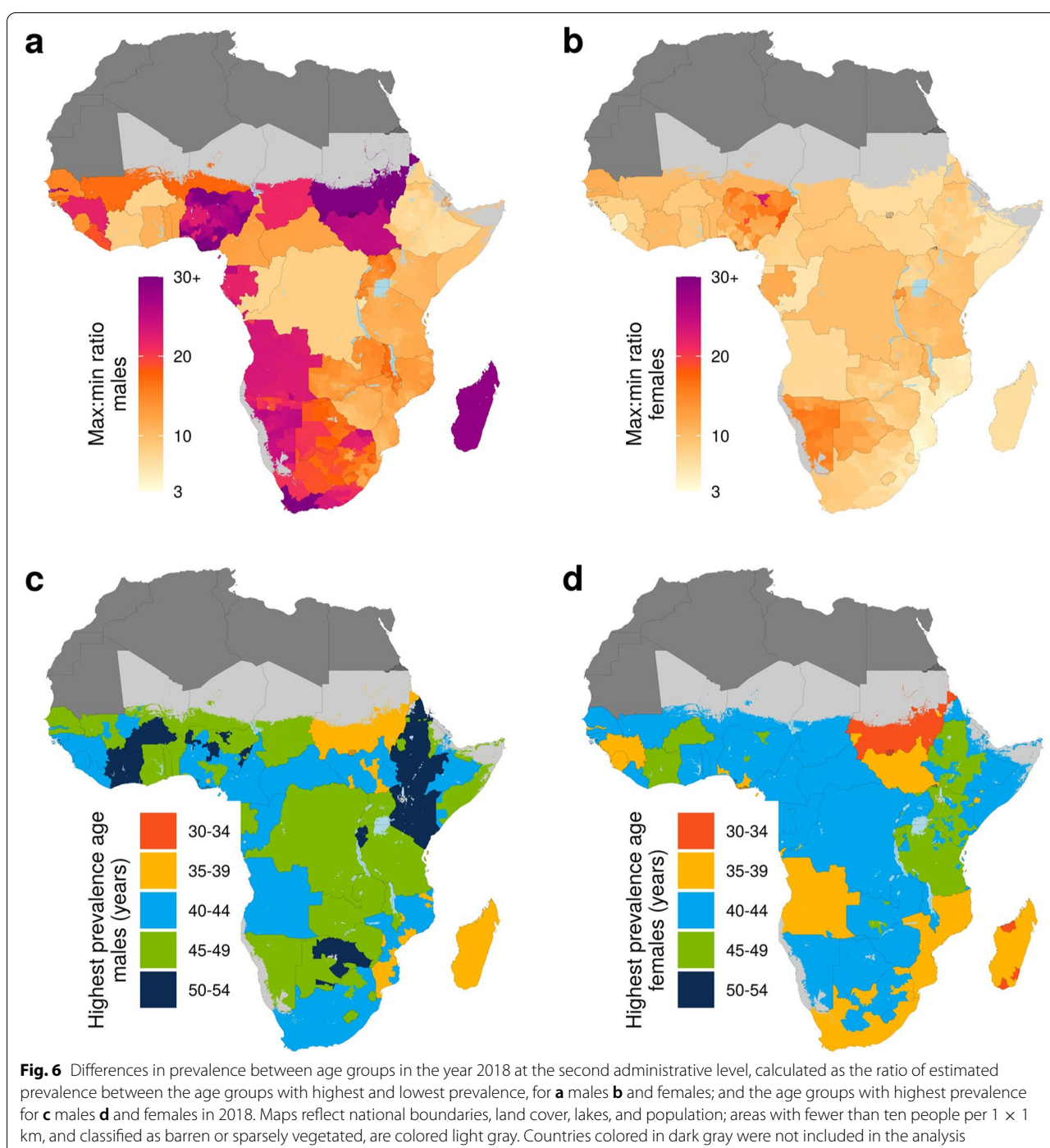
Across SSA, the age group with the highest estimated prevalence in any given second-level administrative unit in 2018 was always between ages 35 and 54 years for males and between 30 and 49 years for females (Fig. 6). In 2018, males ages 45–49 years most commonly had the highest estimated prevalence across all age groups in a given second-level administrative unit, at 46.8% of second-level administrative units (1894 of 4043) from within 23 of 43 countries. Females ages 40–44 years had the highest estimated prevalence across age groups in 63.8% of second-level administrative units (2581 of

4043) in 31 of 43 countries. For both males and females, the age group with the highest estimated prevalence tended to vary more across Eastern SSA compared to other regions.

Within-country variation between second-level administrative units was relatively consistent across age groups. The ratio of maximum to minimum estimated prevalence among districts within each country was lowest for ages 35–39 years (median ratio of 4.3 across countries) and highest for ages 15–19 years (median ratio of 4.8 across countries) in 2018. Slightly larger differences were seen between age groups in Eastern and Southern SSA, with lower variation in middle-age groups and greater within-country variation in younger age groups. The maximum-to-minimum within-country prevalence ratio in Eastern SSA was lowest for adults ages 40–44 years (median ratio of 5.4 across Eastern SSA countries) and highest for adults ages 15–19 years (median ratio of 6.7 across Eastern SSA countries). These same age groups also represented the highest and lowest ratios in Southern SSA countries, with median values of 2.0 in adults ages 40–44 years and 2.8 in adults ages 15–19 years.

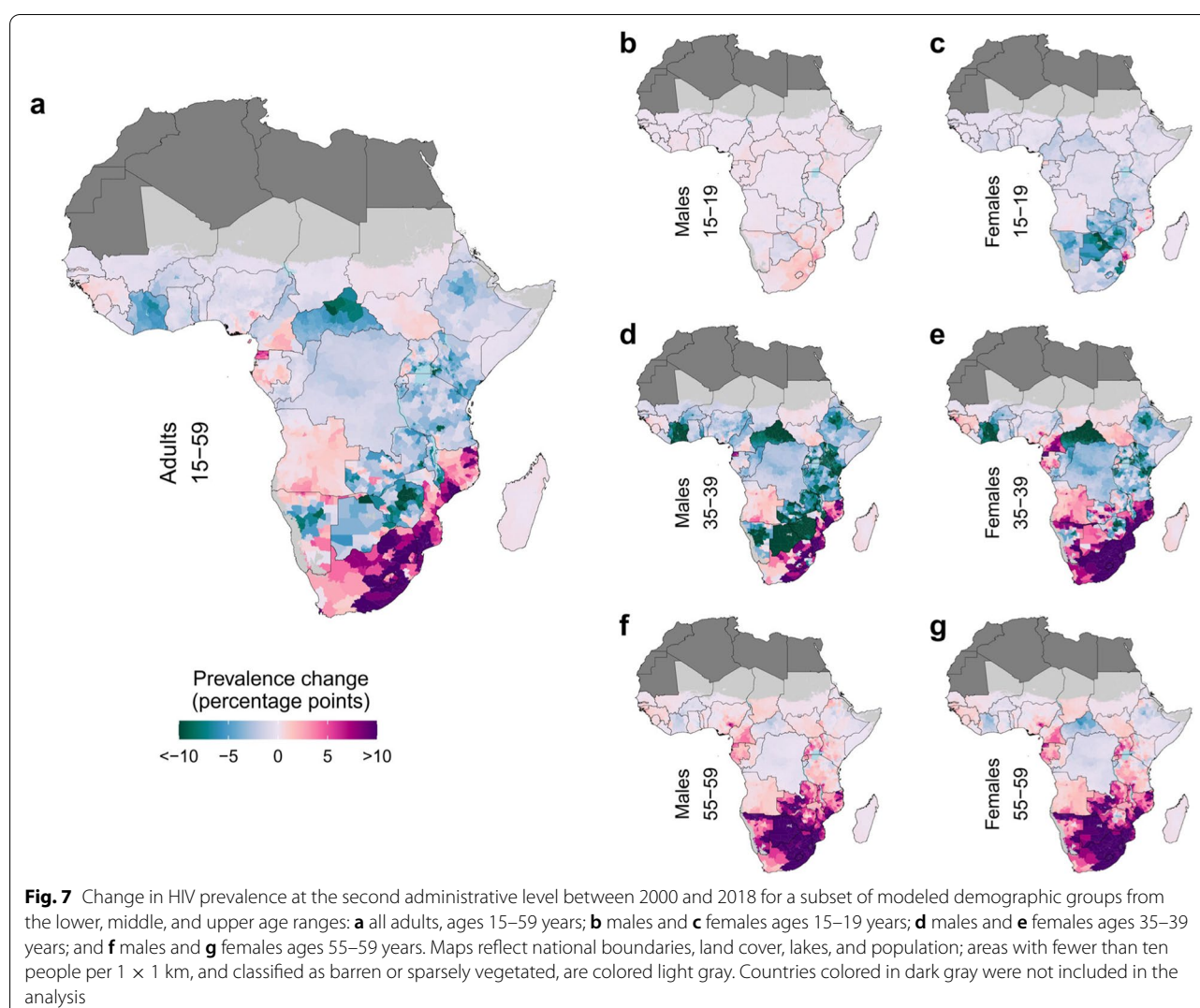
Variation over time

Estimated change in prevalence over time among all adults masked broad differences between specific age and sex groups (Fig. 7; Additional file 3: Figs. S35–S40). Large temporal changes were much more common



when considering sexes and age groups, compared to all adults combined. Between the years 2000 and 2018, among all adults ages 15–59 years, estimated HIV prevalence increased by more than 5.0 percentage points in only 3.7% (151 out of 4043) of second-level administrative units across SSA and decreased by more than 5.0 percentage points in 7.9% (321 of 4043) of second-level

administrative units. On the other hand, 37.7% (1523 of 4043) of second-level administrative units experienced an increase in estimated HIV prevalence greater than 5.0 percentage points in that timeframe in at least one sex and age group, and 70.9% (2867 of 4043) of second-level administrative units saw a decrease greater than 5.0 percentage points in at least one sex and age group.



The distribution of districts with large increases or decreases in prevalence over time also varied greatly by region. All regions saw a decrease of greater than 5.0 percentage points in estimated prevalence for at least one sex and age group in a majority of second-level administrative units between 2000 and 2018: 61.2% for Central SSA, (393 out of 642), 70.9% (1160 out of 1635) for Western SSA, 71.0% (1032 out of 1452) for Eastern SSA, and 90.1% (283 out of 314) for Southern SSA. However, Southern SSA also had a very high proportion of second-level administrative units seeing an increase of greater than 5.0 percentage points in that same time frame, at 92.0% (289 of 314), while only a minority of second-level administrative units saw similar increases in the other regions.

We found diverging overall trends between age groups over time, with greater decreases over time among

younger age groups, and greater increases among older age groups. For example, for females ages 25–29 years, we found that estimated prevalence decreased by at least 1.0 percentage point in the year 2018 compared to 2000 in more than 73.3% of second-level administrative units in SSA (2965 of 4043) and increased by at least 1.0 percentage point in only 2.4% (99 of 4043) of all second-level administrative units. Conversely, among females ages 50–54 years, estimated prevalence decreased between 2000 and 2018 by at least 1.0 percentage point in just 11.8% (477 of 4043) of second-level administrative units but increased by at least 1.0 percentage point in 40.1% (1622 of 4043) of second-level administrative units. We found this trend to be similar across regions. For complete comparisons of prevalence over time for each second-level administrative unit, age, and sex, including uncertainty estimates, see Additional file 4.

Discussion

The results of this study, the first to present age- and sex-specific HIV prevalence estimates across sub-Saharan Africa at local scales, emphasize the interactions of geographic and demographic differences in HIV prevalence, going beyond previous research focused on either aspect individually. Just as previous work demonstrated how much geographic variability is masked in national prevalence estimates [10], we show here that demographically aggregated estimates mask important variation in the age and sex distributions of HIV prevalence at a local level, which in turn provide much clearer insights into the evolution of the HIV epidemic in SSA.

Many intervention methods are commonly used in the fight against the HIV epidemic, and variation in their efficacy and implementation has likely contributed to the prevalence trends presented here. Cost-efficiency is a consistent priority and is generally maximized by using targeted, integrated interventions [50]. For example, HIV prevention via behavioral and biomedical interventions based on local prevalence rates, HIV testing, and treatment initiation may be priorities for some age groups [51], while long-term ART retention and comorbidity care may require more emphasis for others [52]. Barriers to access to care often differ between geographic and demographic groups, where in some cases barriers may be logistical (e.g., geographic isolation and programmatic fragmentation [53]) or social (e.g., lack of information, stigmatization, homophobia [54]), and require different intervention methods. Males and females are also often targeted using different points of contact. For example, HIV testing has been recommended for all females attending antenatal care clinics [55], whereas for males the provision of self-, home-based, and mobile testing compared to facility-based testing may be more useful for testing and subsequent uptake of care [56–58]. Effective targeting of these interventions requires local, demographically specific HIV burden information, such as provided in the estimates presented here. Countries may similarly use this burden information to prioritize subnational and demographically specific treatment needs. This resource may also be useful in program evaluation efforts and thus aid the development of more successfully tailored interventions.

Variation in the social determinants driving HIV incidence and mortality, and thus HIV prevalence, are also an important consideration when assessing inequalities in HIV prevalence between locations and demographic groups. While prevalence among females is consistently higher than prevalence among males, for example, these differences can be attributed to different exposure to risk factors (such as age at first sex between

males and females, marital status) in different countries [59]. In addition to understanding local patterns in HIV prevalence, effective interventions also need to consider, if not focus directly on, locally important risk factors and determinants of HIV infection and mortality [60, 61].

Our estimates point to many local shifts in HIV prevalence over time. A multitude of factors can affect HIV prevalence trends at the local level over time, from local changes in prevention interventions to shifts in the overall demographics of an area, but one particularly important factor is local scale-up of ART [62, 63]. Increases in ART coverage and reduced treatment costs have repeatedly been associated with large demographic shifts among people living with HIV [64] due to its success in reducing HIV mortality, leading to greatly increasing numbers of people living with HIV over the age of 50 years; our results reflect this trend. Given evidence pointing to differences between younger and older ART patients in rates of CD4 cell count decline [65], immune reconstitution rates [66], and risk of associated non-communicable diseases [67, 68], among other health metrics [69], it is necessary that treatment plans for older patients be specifically tailored for their age group. Our results highlight those locations with large existing populations of people living with HIV for ages 50–59 years, and those seeing rapid growth of HIV prevalence in that demographic group. At the same time, the minimal change in estimated prevalence over time among the youngest age groups suggests that continued and even expanded efforts in HIV prevention for adolescents and young adults still need to be maintained as a priority across the continent.

Despite the significant progress made through this analysis in describing HIV burden in SSA, prevalence estimates mask complex and varied relationships between HIV incidence and mortality, as well as migration and seasonal mobility. It is difficult to determine, for example, if a dramatic decrease in HIV prevalence in an area is due to reduced incidence, increased mortality, or differences in the immigration and emigration rates of HIV⁺ and HIV⁻ individuals. Primary data for all three of these metrics are not widely available for SSA, adding additional complexity to the interpretation of our estimates. Importantly, no estimates of these indicators are consistently available at local scales for specific demographic groups. Furthermore, local data related to diagnosis, treatment, and viral suppression rates are also limited, despite these metrics lying at the heart of the UNAIDS 95-95-95 goals [4]. While very informative, difficulties can still arise in intervention decision-making built around HIV prevalence estimates alone, without understanding their underlying drivers. Improved surveillance of HIV prevalence, incidence,

and mortality, combined with reliable population and migration estimates and information on local programs, are necessary to fully understand the complexities of the region's HIV epidemic. Clearly, even with the development of more comprehensive burden information, any modeled estimates should only be used for intervention purposes in conjunction with local program knowledge.

Methodological advantages and limitations

The methods used in this analysis build upon those previously used by Dwyer-Lindgren et al. to model adult HIV prevalence [10]. While this analysis does improve upon and have advantages over the previous methods in some ways, it faces some of the same, as well as some new limitations. As with the previous study, and as with all modeling studies, the quality of our estimates is highly dependent on the quality and coverage of our input data. Despite constructing a large database of HIV prevalence data, coverage gaps and small sample sizes in some locations can be associated with imprecision and/or large uncertainty intervals in some of our prevalence estimates (Additional file 3: Figs. S27–S34). Additionally, the location information associated with the data compiled for this analysis is subject to some error. In order to protect respondent confidentiality, most surveys that collect GPS coordinates perform some type of random displacement on those coordinates prior to releasing data for secondary analysis: for example, GPS coordinates for Demographic and Health Surveys (DHS) are displaced by up to 2 km for urban clusters, up to 5 km for most rural clusters, and up to 10 km in a random 1% of rural clusters [34]. Past research has found that displacement can degrade the predictive power of a geostatistical model, however this effect was found to be modest, and researchers concluded that relatively accurate mapping can be undertaken at a 5×5 -km resolution even with GPS displacement [70].

The approximate integration method we use in this analysis better handles uncertainty estimation and easily accommodates not only polygon data but age-aggregated data as well, compared to the polygon resampling method that has been used elsewhere [10, 71, 72]. At the same time, given the large number of dimensions being modeled, as well as the high data input count produced by our data disaggregation technique, we found that current matrix packages, as well as our computational facilities, could not accommodate a Gaussian process that accounted for the covariance of a complete space-time-age-sex Kronecker product. We therefore focused on the interactions between space, time, age, and sex that we believed would be most relevant in terms of capturing important variability in these dimensions, within our

computational abilities. Our modeling strategy also assumed no difference in the probability that an HIV⁺ versus an HIV⁻ pregnant woman would access antenatal care and therefore be included in ANC surveillance.

Due to limited data availability, we delineated estimates in this analysis using a male/female binary. We recognize that this approach does not allow for investigation of HIV prevalence among gender and sex diverse people, despite the disproportionate burden of HIV commonly seen among these populations [73]. Further, we recognize that many data sources do not provide the option to select a sex other than “male” or “female,” gender options beyond “man” or “woman,” and often conflate gender with sex. In the future, we hope that high-quality data on HIV prevalence for gender and sexual diverse people will be more widely available, so we can produce estimates beyond females and males.

We note that our results include unprecedentedly high prevalence estimates for certain population subsets. In most cases, we do not believe these estimates are implausible. For example, we estimated prevalence among middle- and older-aged females to be up to 59.2% [45.9–73.0%] in Umgungundlovu in KwaZulu-Natal, South Africa in 2018. Previous research has estimated prevalence for females adults of all ages combined in Umgungundlovu in 2017 to be 46.6% [43.8–49.5%] [74]. As we have shown that prevalence in middle- and older-aged females tended to be higher than all-ages prevalence, we believe our estimates for middle- and older-aged females during this time period in this location to be reasonable, especially with uncertainty intervals taken into consideration. In rare cases, however, our methods yielded estimates which we were unable to support through the literature. For example, for males ages 35–39 and 40–44 years in Nyatike in Migori, Kenya, we estimated prevalence in the year 2000 to be 77.8% [50.2–100.0%] and 78.7% [50.0–100.0%], respectively. It is unlikely true prevalence in that area and year was this high (though given the large uncertainty intervals associated with these values, it is probable that true prevalence does fall within those ranges). We note, however, that the high estimates in this area and surrounding second-level administrative units were predominantly associated with the earlier years in our time series—we believe the more recent estimates in Nyatike to be more realistic [75]. In these locations, decreases in prevalence over time may therefore also be overestimated. These instances were rare.

A combination of data limitations and model complexity ultimately led to large uncertainty intervals around our estimates. Given that our 95% coverage estimates in model validation were consistently higher than expected (Additional file 3: Figs. S14–S16), this

indicates that these uncertainty intervals may be larger than appropriate. Wide uncertainty can limit the utility of our estimates in terms of informing HIV policies, and reducing this uncertainty through improved data coverage will be an important consideration in future iterations of this model. We were also unable to account for all sources of uncertainty such as uncertainty in the WorldPop estimates used in many stages of our modeling and estimation processes and uncertainty in covariates.

Conclusions

HIV continues to impose enormous human and financial costs [3] on SSA, decades since its emergence. Financial and logistical disruptions and discontinuities due to the impacts of COVID-19, as well as changes in ART adherence, are likely to present new barriers [21, 76] to the UNAIDS 95-95-95 goals [4]. This analysis provides important insight into the nuances of HIV burden in SSA, offering information that is critical to the development of targeted interventions.

Abbreviations

AIDS: Acquired immune deficiency syndrome; ANC: Antenatal care; ART: Antiretroviral therapy; GATHER: Guidelines for Accurate and Transparent Health Estimates Reporting; GBD: Global Burden of Disease; GPS: Global positioning system; HIV: Human immunodeficiency virus; SDGs: Sustainable Development Goals; SSA: Sub-Saharan Africa; TMB: Template Model Builder; UNAIDS: Joint United Nations Programme on HIV/AIDS.

Supplementary Information

The online version contains supplementary material available at <https://doi.org/10.1186/s12916-022-02639-z>.

Additional file 1: Supplemental information. 1. Compliance with the Guidelines for Accurate and Transparent Health Estimates Reporting (GATHER). 2. HIV data sources and data processing. 3. Covariate and auxiliary data. 4. Statistical model. 5. References.

Additional file 2: Supplemental tables. Table S1. HIV seroprevalence survey data. Table S2. ANC sentinel surveillance data. Table S3. HIV and covariates surveys excluded from this analysis. Table S4. Sources for pre-existing covariates. Table S5. HIV covariate survey data. Table S6. Fitted model parameters.

Additional file 3: Supplemental figures. Figure S1. Prevalence of male circumcision. Figure S2. Prevalence of signs and symptoms of sexually transmitted infections. Figure S3. Prevalence of marriage or living as married. Figure S4. Prevalence of partner living elsewhere among females. Figure S5. Prevalence of condom use during most recent sexual encounter. Figure S6. Prevalence of sexual activity among young females. Figure S7. Prevalence of multiple partners among males in the past year. Figure S8. Prevalence of multiple partners among females in the past year. Figure S9. HIV prevalence predictions from the boosted regression tree model. Figure S10. HIV prevalence predictions from the generalized additive model. Figure S11. HIV prevalence predictions from the lasso regression model. Figure S12. Modeling regions. Figure S13. Age- and sex-specific vs. adult prevalence modeling. Figure S14. Data sensitivity. Figure S15. Model specification validation. Figure S16. Modeled and re-aggregated adult prevalence comparison. Figure S17. HIV prevalence raking factors for males. Figure S18. HIV prevalence raking factors for females. Figure S19. Age-specific HIV prevalence in males, 2000. Figure

S20. Age-specific HIV prevalence in females, 2000. Figure S21. Age-specific HIV prevalence in males, 2005. Figure S22. Age-specific HIV prevalence in females, 2005. Figure S23. Age-specific HIV prevalence in males, 2010. Figure S24. Age-specific HIV prevalence in females, 2010. Figure S25. Age-specific HIV prevalence in males, 2018. Figure S26. Age-specific HIV prevalence in females, 2018. Figure S27. Age-specific uncertainty interval range estimates in males, 2000. Figure S28. Age-specific uncertainty interval range estimates in females, 2000. Figure S29. Age-specific uncertainty interval range estimates in males, 2005. Figure S30. Age-specific uncertainty interval range estimates in females, 2005. Figure S31. Age-specific uncertainty interval range estimates in males, 2010. Figure S32. Age-specific uncertainty interval range estimates in females, 2010. Figure S33. Age-specific uncertainty interval range estimates in males, 2018. Figure S34. Age-specific uncertainty interval range estimates in females, 2018. Figure S35. Change in HIV prevalence in males, 2000-2005. Figure S36. Change in HIV prevalence in females, 2000-2005. Figure S37. Change in HIV prevalence in males, 2005-2010. Figure S38. Change in HIV prevalence in females, 2005-2010. Figure S39. Change in HIV prevalence in males, 2010-2018. Figure S40. Change in HIV prevalence in females, 2010-2018. Figure S41. Space mesh for geostatistical models.

Additional file 4: Supplemental results. 1. README. 2. Prevalence range across districts. 3. Prevalence range between sexes. 4. Prevalence range between ages. 5. Age-specific district ranges.

Acknowledgements

LBD sub-Saharan Africa HIV Prevalence Collaborators
S Afzal acknowledges support of the Pakistan Society of Medical Infectious Diseases and King Edward Medical University to access the relevant data of HIV from various sources. T W Bärnighausen was supported by the Alexander von Humboldt Foundation through the Alexander von Humboldt Professor award, funded by the German Federal Ministry of Education and Research. F Carvalho and E Fernandes acknowledge support from Fundação para a Ciência e a Tecnologia (FCT), I.P., in the scope of the project UIDP/04378/2020 and UIDB/04378/2020 of the Research Unit on Applied Molecular Biosciences - UCIBIO and the project LA/P/0140/2020 of the Associate Laboratory Institute for Health and Bioeconomy - i4HB; FCT/MCTES (Ministério da Ciência, Tecnologia e Ensino Superior) through the project UIDB/50006/2020. K Deribe acknowledges support by the Wellcome Trust [grant number 201900/Z/16/Z] as part of his International Intermediate Fellowship. C Herteliu and A Pana are partially supported by a grant of the Romanian National Authority for Scientific Research and Innovation, CNDS-UEFISCDI, project number PN-III-P4-ID-PCCF-2016-0084. Claudiu Herteliu is partially supported by a grant of the Romanian Ministry of Research Innovation and Digitalization, MCID, project number ID-585-CTR-42-PFE-2021. Y J Kim acknowledges support by the Research Management Centre, Xiamen University Malaysia [No. XMUMRF/2020-C6/ITCM/0004]. S L Koulmane Laxminarayana acknowledges institutional support by the Manipal Academy of Higher Education. K Krishan acknowledges non-financial support from UGC Centre of Advanced Study, CAS II, Department of Anthropology, Panjab University, Chandigarh, India. M Kumar would like to acknowledge NIH/FIC K43 TW010716-04. I Landires is a member of the Sistema Nacional de Investigación (SNI), supported by the Secretaría Nacional de Ciencia, Tecnología e Innovación (SENACYT), Panama. V Nuñez-Samudio is a member of the Sistema Nacional de Investigación (SNI), which is supported by Panama's Secretaría Nacional de Ciencia, Tecnología e Innovación (SENACYT). O O Odukeya was supported by the Fogarty International Center of the National Institutes of Health under the Award Number K43TW010704. The content is solely the responsibility of the authors and does not necessarily represent the official views of the National Institutes of Health. Z Quazi Syed acknowledges support from JNMC, Datta Meghe Institute of Medical Sciences. A I Ribeiro was supported by National Funds through FCT, under the 'Stimulus of Scientific Employment - Individual Support' program within the contract CEECIND/02386/2018. A M Samy acknowledges the support from a fellowship of the Egyptian Fulbright Mission program and Ain Shams University. R Shrestha acknowledges support from NIDA K01 Award: K01DA051346. N Taveira acknowledges support from FCT and Aga Khan Development Network (AKDN) - Portugal Collaborative Research Network in Portuguese speaking countries in Africa (project reference: 332821690), and by the European & Developing Countries

Clinical Trials Partnership (EDCTP), UE (project reference: RIA2016MC-1615). B Unnikrishnan acknowledges support from Kasturba Medical College, Mangalore, Manipal Academy of Higher Education, Manipal.

Authors' contributions

Individual author contributions are divided into the following categories: managing the estimation or publication process; writing the first draft of the manuscript; primary responsibility for applying analytical methods to produce estimates; primary responsibility for seeking, cataloging, extracting, or cleaning data; designing or coding figures and tables; providing data or critical feedback on data sources; developing methods or computational machinery; providing critical feedback on methods or results; drafting the manuscript or revising it critically for important intellectual content; and managing the overall research process. Members of the core research team for this topic area had full access to the underlying data used to generate estimates presented in this article. All other authors had access to and reviewed estimates as part of the research evaluation process, which includes additional stages of formal review. All authors read and approved the final manuscript. Managing the estimation or publications process: LDL, EH, SIH. Writing the first draft of the manuscript: EH. Primary responsibility for applying analytical methods to produce estimates: EH, MAC. Primary responsibility for seeking, cataloging, extracting, or cleaning data; designing or coding figures and tables: ALS, MY. Providing data or critical feedback on data sources: ALS, ESA, OMA, VA1, SA1, BOA, MBA, ROA, CJA, FA, FMA, TMA, RKA, AAH, NAG, DAA2, DA, SCYA, JA, OA, MAA1, BPAQ, YAA, SA3, TKB, AAB, MB, TWB, ASB, SB, OAB, AB2, VKC, DTC, RGC, GD, AMD, JDN, MD, AD, FD, BD, BE, SE1, IF, TF, MMG, BGG, LG, KG, AG, MG, HCG, AHM1, SH2, HH, CH, HCH, MH2, SEI, CCDI, MJ, KBJ, FJ, JJJ, LRK, RK, NK1, SEK, GAK, MK1, MNK, KK1, JK, YJK, AK1, SK1, SLKL, KK2, BKD, NK2, MK2, DK, DKL, SL1, XL, MAM, FRMM, WM, RGM, EWM, BM, AMH, SM1, SM2, AHM2, MM, SM3, GBBM, AJN, MDN, MN, IN, FNN, GNT, JWN, CTN, HLTN, CAN, JJN, BO, ATO, BOO1, BOO2, JOO, AOB, OEO1, OEO2, MOO, MPA, JRP, AP1, AP2, FPK, SP, MJP, LP, ZQS, NR1, AR1, AR2, FR, VRM, AMR, KR, JR, CLR, SJR, DLR, SR, RR, LDR, NR2, MAR, ER, GMR, RS1, MS1, MRS, HSK, AMS, BS2, BS3, AAS3, MAS, JIS, RS2, JAS, VYS, AAS2, TT, NT, FHT, MRTP, CDU, BU, BV, GTV, YDW, YW1, SY, NY, CY, ZJZ, SIH. Developing methods or computational machinery: MY, DAA1, SA1, MBA, TMA, AAH, DA, ASB, SB, DTC, AMD, MD, FD, BD, MH2, SEI, NK1, MK1, AK1, SM1, AHM2, AOB, ZQS, NR1, AMR, CLR, RR, LDR, ER, AMS, BS2, TT, BV, KW, MSZ, SIH, LDL. Providing critical feedback on methods or results: MY, HA, ESA, MA, OMA, VA1, DAA1, SA1, BOA, KA, MBA, YA, ROA, CJA, FA, FMA, TMA, KAA, RKA, VA2, AAH, NAG, EKA, SA2, DAA2, DA, SCYA, JA, OA, MAA1, MAJ, BPAQ, MAA2, SA3, ZNA, DBB, TKB, AAB, MB, TWB, AWB, ASB, NB, PB, KB, AB1, ZWB, SB, OAB, AB2, IB, ZAB, JC, VKC, MAK, DTC, RGC, SMAD, GD, EKMD, AMD, JDN, NDW1, JWDN, KD1, KD2, MD, DD, AD, SD2, FD, ED, BD, SE1, OE, IJE, FF1, NF, IF, FF2, MF, TF, MMG, AMG, BGG, KBG, LG, AG, MG, MIMG, HCG, RAG, MRH, SH1, AH, SH2, HH, KH, CH, HCH, RH, MH2, BFH, SEI, OSI, IMI, MDI, RMI, CCDI, MJ, RPJ, JSJ, VJ, FJ, JJJ, LRK, RK, NK1, TK, BKM, SEK, GAK, AKK, EAK, MNK, KK1, JK, YJK, AK1, SK1, JAK, SK2, SLKL, AK2, KK2, BKD, NK2, YK, MK2, NK3, DK, CLV, DKL, HJL, SL1, SL2, XL, PM1, MAM, FRMM, WM, RGM, EWM, TM, AM2, BM, OM, YM, AMH, RM, SM1, SM2, AHM2, SM3, GBBM, LM, AJN, MDN, MN, JN, RN, IN, FNN, GNT, JWN, CTN, HLTN, CAN, JJN, VEN, BO, ATO, BOO1, BOO2, JOO, AOB, OEO1, OEO2, NO, SSO, MOO, MPA, JRP, AP1, SRPP, FPK, SP, EKP, MJP, LP, ZQS, NR1, AR1, AR2, FR, VRM, AMR, JR, CLR, SJR, DLR, SR, RR, LDR, NR2, AR3, AIR, JMR, ER, SFR, GMR, SSM, RS1, BS1, MS1, MRS, HSK, AMS, BS2, BS3, AAS3, AAS1, MAS, MS2, WSS, JIS, RS2, JAS, VYS, AAS2, SS, MBS, TT, EGT, NT, FHT, RT, MRTP, GWT, CDU, BU, JHV, FSV, BV, GTV, YDW, YW1, RGW, YW2, PW, NDW2, SY, PY, NY, CY, MSZ, YZ, SIH, LDL. Drafting the work or revising it critically for important intellectual content: ALS, MY, HA, OMA, VA1, DAA1, SA1, BOA, MBA, YA, ROA, FA, KAA, RKA, AAH, NAG, SA2, DAA2, RA, JA, OA, MAA1, AFA, BPAQ, MAA2, ZNA, AAB, MB, TWB, AWB, ASB, KB, SB, IB, RC, FC, VKC, DTC, RGC, GD, EKMD, JDN, NDW1, JWDN, KD1, SDD, MD, DD, AD, FD, MET, SIEJ, SE1, OE, SE2, NKF, EF, IF, FF2, NAF, MF, TF, MMG, AG, MIMG, RAG, AH, CH, HCH, RH, MH1, SEI, OSI, IMI, MDI, RMI, CCDI, MJ, RPJ, NJ, JJJ, GAK, EAK, GK, MNK, KK1, JK, YJK, AK1, SK1, SLKL, AK2, KK2, BKD, NK2, DK, CLV, IL, SL1, PHL, SL2, AM1, PM1, FRMM, WM, RGM, EWM, TJM, TM, BM, OM, YM, RM, SM1, SM2, AHM2, PM2, LM, AJN, JN, IN, FNN, GNT, JWN, CTN, HLTN, JJN, VNS, VEN, OOO, ATO, BOO1, BOO2, JOO, OEO1, NO, MOO, MPA, JRP, AP1, FPK, GCP, SP, MJP, LP, ZQS, NR1, AR1, AR2, FR, VRM, CLR, SJR, DLR, SR, NR2, MAR, AIR, JMR, ER, GMR, RS1, BS1, MS1, MRS, HSK, AMS, RS2, JAS, VYS, AAS2, MBS, TT, EGT, NT, MRTP, BU, JHV, FSV, GTV, YW1, RGW, YW2, PW, NDW2, SY, NY, MSZ, ZJZ, SIH, LDL. Managing the overall research enterprise: LDL, SIH, AHM2, GCP, and BS2.

Funding

This work was primarily supported by grant OPP1132415 from the Bill & Melinda Gates Foundation. The funder of the study had no role in study design, data collection, data analysis, data interpretation, writing of the report, or decision to publish. The corresponding authors had full access to all the data in the study and had final responsibility for the decision to submit for publication.

Availability of data and materials

The findings of this study are supported by data available in public online repositories and data publicly available upon request of the data provider. Details regarding the data sources used and their availability can be found in Additional file 2: Supplemental Tables 1-5 and online via the Global Health Data Exchange (<https://ghdx.healthdata.org/record/ihme-data/sub-saharan-africa-hiv-prevalence-geospatial-estimates-2000-2018>). Estimates can also be further explored through the Global Health Data Exchange, as well as via our online visualization tool (<http://vizhub.healthdata.org/lbd/hiv-prev-disagg>). Administrative boundaries were modified from the Database for Global Administrative Areas (GADM) dataset [77]. Populations were retrieved from WorldPop [32]. This study complies with the Guidelines for Accurate and Transparent Health Estimates Reporting (GATHER) recommendations [31]. All maps and figures presented in this study are generated by the authors; no permissions are required for publication. All computer code is available online and can be found at (https://github.com/ihmeuw/lbd/tree/hiv_prev-africa-2020).

Declarations

Ethics approval and consent to participate

Not applicable.

Consent for publication

Not applicable.

Competing interests

S Afzal reports leadership or fiduciary role in other board, society, committee or advocacy group, unpaid, with the Pakistan society of Community Medicine & Public Health, the Pakistan Association of Medical Editors, and the Pakistan Society of Medical Infectious Diseases, all outside the submitted work. R Ancuceanu reports 5 payment or honoraria for lectures, presentations, speakers bureaus, manuscript writing, or educational events from Avvie, Sandoz, and B Braun, all outside the submitted work. T W Bärnighausen reports research grants from the European Union (Horizon 2020 and EIT Health), German Research Foundation (DFG), US National Institutes of Health, German Ministry of Education and Research, Alexander von Humboldt Foundation, Else-Kröner-Fresenius-Foundation, Wellcome Trust, Bill & Melinda Gates Foundation, KfW, UNAIDS, and WHO; consulting fees from KfW on the OSCAR initiative in Vietnam; participation on a Data Safety Monitoring Board or Advisory Board with the NIH-funded study "Healthy Options" (PIs: Smith Fawzi, Kaaya), Chair, Data Safety and Monitoring Board (DSMB), German National Committee on the "Future of Public Health Research and Education," Chair of the scientific advisory board to the EDCTP Evaluation, Member of the UNAIDS Evaluation Expert Advisory Committee, National Institutes of Health Study Section Member on Population and Public Health Approaches to HIV/AIDS (PPAH), US National Academies of Sciences, Engineering, and Medicine's Committee for the "Evaluation of Human Resources for Health in the Republic of Rwanda under the President's Emergency Plan for AIDS Relief (PEPFAR)," University of Pennsylvania (UPenn) Population Aging Research Center (PARC) External Advisory Board Member; leadership or fiduciary role in other board, society, committee or advocacy group, paid or unpaid, as co-chair of the Global Health Hub Germany (which was initiated by the German Ministry of Health); all outside the submitted work. J das Neves reports grants or contracts from Ref. 13605 – Programa GENESE, Gilead Portugal (PGG/002/2016 – Programa GENESE, Gilead Portugal) outside the submitted work. L Dwyer-Lindgren reports support for the present manuscript from the Bill & Melinda Gates Foundation through grant OPP1132415. I Filip reports other financial or non-financial interests from Avicenna Medical and Clinical Research Institute, outside the submitted work. E Haeuser reports support for

the present manuscript from the Bill & Melinda Gates Foundation through grant OPP1132415. C Herteliu reports grants from Romanian Ministry of Research Innovation and Digitalization, MCID, for project number ID-585-CTR-42-PFE-2021 (Jan 2022-Jun 2023) "Enhancing institutional performance through development of infrastructure and transdisciplinary research ecosystem within socio-economic domain – PERFECTIS," from Romanian National Authority for Scientific Research and Innovation, CNDS-UEFISCDI, for project number PN-III-P4-ID-PCCF-2016-0084 (Oct 2018-Sep 2022) "Understanding and modelling time-space patterns of psychology-related inequalities and polarization," and project number PN-III-P2-2.1-SOL-2020-2-0351 (Jun 2020-Oct 2020) "Approaches within public health management in the context of COVID-19 pandemic," and from the Ministry of Labour and Social Justice, Romania for project number "Agenda for skills Romania 2020-2025"; all outside the submitted work. J J Jozwiak reports payment or honoraria for lectures, presentations, speakers bureaus, manuscript writing, or educational events from Teva, Amgen, Synexus, Boehringer Ingelheim, Zentiva, and Sanofi as personal fees, all outside the submitted work. J Khubchandani reports other financial interests from Teva Pharmaceuticals, all outside the submitted work. K Krishnan reports other non-financial support from UGC Centre of Advanced Study, CAS II, Department of Anthropology, Panjab University, Chandigarh, India, outside the submitted work. H J Larson reports grants or contracts from the MacArthur Foundation and Merck to London School of Hygiene and Tropical Medicine, and from the Vaccine Confidence Fund to the University of Washington; payment or honoraria for lectures, presentations, speakers bureaus, manuscript writing, or educational events from Center for Strategic and International Studies as payment to LSHTM for co-chairing HighLevel Panel and from GSK as personal payment for developing training sessions and lectures; leadership or fiduciary role in other board, society, committee or advocacy group, pair, with the ApiJect Advisory Board; all outside the submitted work. O O Odukoya reports support for the present manuscript from the Fogarty International Center of the National Institutes of Health under the Award Number K43TW010704. The content is solely the responsibility of the authors and does not necessarily represent the official views of the National Institutes of Health. A Pans reports grants from Romanian National Authority for Scientific Research and Innovation, CNDS-UEFISCDI, for project number PN-III-P4-ID-PCCF-2016-0084 (Oct 2018-Sep 2022) "Understanding and modelling time-space patterns of psychology-related inequalities and polarization," and project number PN-III-P2-2.1-SOL-2020-2-0351 (Jun 2020-Oct 2020) "Approaches within public health management in the context of COVID-19 pandemic," outside the submitted work. S R Pandi-Perumal reports royalties from Springer for editing services; stock or stock options in Somnogen Canada Inc as the President and Chief Executive Officer; all outside the submitted work. A Radfar reports other financial or non-financial interests from Avicenna Medical and Clinical Research Institute, outside the submitted work. A I Ribeiro reports grants or contracts from National Funds through FCT, under the 'Stimulus of Scientific Employment – Individual Support' program within the contract CEECIND/02386/2018, outside the submitted work. J M Ross reports support for the present manuscript from the Bill & Melinda Gates Foundation through grant OPP1132415; grants or contracts from National Institutes of Health and Firland Foundation as payments to their institution; consulting fees from United States Agency for International Development as personal payments, and from KNCV Tuberculosis Foundation as payments to their institution; all outside the submitted work. E Rubagotti reports payment or honoraria for lectures, presentations, speakers bureaus, manuscript writing, or educational events from the Greenwich China Office and University Prince Mohammad VI, Morocco, all outside the submitted work. B Sartorius reports grants or contracts from DHSC – GRAM Project; Leadership or fiduciary role in other board, society, committee or advocacy group, paid or unpaid, as a member of the GBD Scientific Council and a Member of WHO RGHS; all outside the submitted work. J A Singh reports consulting fees from Crealta/Horizon, Medisys, Fidia, PK Med, Two labs Inc, Adept Field Solutions, Clinical Care options, Clearview healthcare partners, Putnam associates, Focus forward, Navigant consulting, Spherix, MedIQ, Jupiter Life Science LLC, UBM LLC, Trio Health, Medscape, WebMD, and Practice Point communications, and the National Institutes of Health and the American College of Rheumatology; payment or honoraria for participating in the speakers bureau for Simply Speaking; support for attending meetings and/or travel from the steering committee of OMERACT, to attend their meeting every 2 years; participation on a Data Safety Monitoring Board or Advisory Board as an unpaid member of

the FDA Arthritis Advisory Committee; leadership or fiduciary role in other board, society, committee or advocacy group, paid or unpaid, as a member of the steering committee of OMERACT, an international organization that develops measures for clinical trials and receives arm's length funding from 12 pharmaceutical companies, with the Veterans Affairs Rheumatology Field Advisory Committee as Chair, and with the UAB Cochrane Musculoskeletal Group Satellite Center on Network Meta-analysis as a director and editor; stock or stock options in TPT Global Tech, Vaxart pharmaceuticals, Atyu Biopharma, Adaptimmune Therapeutics, GeoVax Labs, Pieris Pharmaceuticals, EnzoLytics Inc, Series Therapeutics, Tonix Pharmaceuticals, and Charlotte's Web Holdings Inc. and previously owned stock options in Amarin, Viking, and Moderna pharmaceuticals; all outside the submitted work. N Taveira reports grants or contracts from FCT and Aga Khan Development Network (AKDN) – Portugal Collaborative Research Network in Portuguese speaking countries in Africa (Project reference: 332821690) and from European & Developing Countries Clinical Trials Partnership (EDCTP), UE (Project reference: RIA2016MC-1615), as payments made to their institution, all outside the submitted work.

Author details

¹Institute for Health Metrics and Evaluation, University of Washington, Seattle, WA, USA. ²Advanced Diagnostic and Interventional Radiology Research Center, Tehran University of Medical Sciences, Tehran, Iran. ³Department of Botany, Sree Narayana Guru College Chelannur, Kozhikode, India. ⁴Hamadan University of Medical Sciences, Hamadan, Iran. ⁵College of Medicine, University College Hospital, Ibadan, Ibadan, Nigeria. ⁶Department of Population Medicine, Cardiff University, Cardiff, UK. ⁷Department of Community Health and Epidemiology, University of Saskatchewan, Saskatoon, SK, Canada. ⁸Department of Public Health, Federal Ministry of Health, Abuja, Nigeria. ⁹Department of Community Medicine, King Edward Memorial Hospital, Lahore, Pakistan. ¹⁰Department of Public Health, Public Health Institute, Lahore, Pakistan. ¹¹The Australian Centre for Public and Population Health Research (ACPPHR), University of Technology Sydney, Sydney, NSW, Australia. ¹²School of Public Health, Imperial College London, London, UK. ¹³Department of Epidemiology, Jimma University, Jimma, Ethiopia. ¹⁴Australian Center for Precision Health, University of South Australia, Adelaide, SA, Australia. ¹⁵Department of Medical Physiology, University of Gondar, Gondar, Ethiopia. ¹⁶Institute for Advanced Medical Research and Training, University of Ibadan, Ibadan, Nigeria. ¹⁷Institute of Neuroscience, Newcastle University, Newcastle upon Tyne, UK. ¹⁸Department of Public Health, The Intercountry Centre for Oral Health (ICOH) for Africa, Jos, Nigeria. ¹⁹Department of Public Health, Federal Ministry of Health, Garki, Nigeria. ²⁰Mayo Evidence-based Practice Center, Mayo Clinic Foundation for Medical Education and Research, Rochester, MN, USA. ²¹Imam Abdulrahman Bin Faisal University, Dammam, Saudi Arabia. ²²Health Information Management and Technology Department, Imam Abdulrahman Bin Faisal University, Dammam, Saudi Arabia. ²³Faculty of Health Sciences, Curtin University, Perth, WA, Australia. ²⁴Wesfarmers Centre of Vaccines and Infectious Diseases, Telethon Kids Institute, Perth, WA, Australia. ²⁵Institute of Health Research, University of Health and Allied Sciences, Ho, Ghana. ²⁶Health Management and Economics Research Center, Iran University of Medical Sciences, Tehran, Iran. ²⁷Department of Health Economics, Iran University of Medical Sciences, Tehran, Iran. ²⁸Department of Epidemiology, Arak University of Medical Sciences, Arak, Iran. ²⁹Research Group in Hospital Management and Health Policies, Universidad de la Costa (University of the Coast), Barranquilla, Colombia. ³⁰Research Group in Health Economics, University of Cartagena, Cartagena, Colombia. ³¹Department of Health Services Management, Khomoin University of Medical Sciences, Khomoin, Iran. ³²Department of Maternal and Child Wellbeing, African Population and Health Research Center, Nairobi, Kenya. ³³Pharmacy Department, Carol Davila University of Medicine and Pharmacy, Bucharest, Romania. ³⁴Department of Parasitology, Mazandaran University of Medical Sciences, Sari, Iran. ³⁵Department of Parasitology, Ilam University of Medical Sciences, Ilam, Iran. ³⁶Department of Sociology and Social Work, Kwame Nkrumah University of Science and Technology, Kumasi, Ghana. ³⁷Center for International Health, Ludwig Maximilians University, Munich, Germany. ³⁸Department of Public Health, Birmingham City University, Birmingham, UK. ³⁹School of Public Health, Bahir Dar University, Bahir Dar, Ethiopia. ⁴⁰Department of Biostatistics and Epidemiology, Tabriz University of Medical Sciences, Tabriz, Iran. ⁴¹Department of Biostatistics and Epidemiology, Zanjan University of Medical Sciences, Zanjan, Iran. ⁴²Department of Surgery, Addis Ababa University, Addis Ababa, Ethiopia. ⁴³The Judith Lumley Centre, La Trobe University, Melbourne, Australia.

- VIC, Australia. ⁴⁴San Martin de Porres University, Lima, Peru. ⁴⁵Department of Health Policy Planning and Management, University of Health and Allied Sciences, Ho, Ghana. ⁴⁶Department of Health Economics, Centre for Health Policy Advocacy Innovation & Research in Africa (CHPAIR-Africa), Accra, Ghana. ⁴⁷Department of Nursing, Debre Berhan University, Debre Berhan, Ethiopia. ⁴⁸Hospital Management Research Center, Iran University of Medical Sciences, Tehran, Iran. ⁴⁹Department of Reproductive Health, University of Gondar, Gondar, Ethiopia. ⁵⁰Kasturba Medical College, Mangalore, Manipal Academy of Higher Education, Manipal, India. ⁵¹Discipline of Public Health Medicine, University of KwaZulu-Natal, Durban, South Africa. ⁵²Department of Community Health and Primary Care, University of Lagos, Lagos, Nigeria. ⁵³Unit of Biochemistry, Universiti Sultan Zainal Abidin (Sultan Zainal Abidin University), Kuala Terengganu, Malaysia. ⁵⁴Department of Hypertension, Medical University of Lodz, Lodz, Poland. ⁵⁵Polish Mothers' Memorial Hospital Research Institute, Lodz, Poland. ⁵⁶Heidelberg Institute of Global Health (HIGH), Heidelberg University, Heidelberg, Germany. ⁵⁷T.H. Chan School of Public Health, Harvard University, Boston, MA, USA. ⁵⁸Department of Global Health and Social Medicine, Harvard University, Boston, MA, USA. ⁵⁹Department of Social Services, Tufts Medical Center, Boston, MA, USA. ⁶⁰Department of Social and Clinical Pharmacy, Charles University, Hradec Kralova, Czech Republic. ⁶¹Institute of Public Health, United Arab Emirates University, Al Ain, United Arab Emirates. ⁶²Department of Anatomy, All India Institute of Medical Sciences, Jodhpur, India. ⁶³Department of Community Medicine and Family Medicine, All India Institute of Medical Sciences, Jodhpur, India. ⁶⁴School of Public Health, All India Institute of Medical Sciences, Jodhpur, India. ⁶⁵Department of Statistical and Computational Genomics, National Institute of Biomedical Genomics, Kalyani, India. ⁶⁶Department of Statistics, University of Calcutta, Kolkata, India. ⁶⁷Social Determinants of Health Research Center, Babol University of Medical Sciences, Babol, Iran. ⁶⁸Nutrition Department, St. Paul's Hospital Millennium Medical College, Addis Ababa, Ethiopia. ⁶⁹St. Paul's Hospital Millennium Medical College, Addis Ababa, Ethiopia. ⁷⁰Department of Veterinary Medicine, Islamic Azad University, Kermanshah, Iran. ⁷¹Department of Internal Medicine, Manipal Academy of Higher Education, Mangalore, India. ⁷²WHO Collaborating Centre for HIV Strategic Information, University of Zagreb, Zagreb, Croatia. ⁷³Faculty of Public Health and Policy, London School of Hygiene & Tropical Medicine, London, UK. ⁷⁴School of Public Health and Health Systems, University of Waterloo, Waterloo, ON, Canada. ⁷⁵Al Shifa School of Public Health, Al Shifa Trust Eye Hospital, Rawalpindi, Pakistan. ⁷⁶Department of Health Care, Metropolitan Autonomous University, Mexico City, Mexico. ⁷⁷Research Unit on Applied Molecular Biosciences (UCIBIO), University of Porto, Porto, Portugal. ⁷⁸Department of Pharmacology, All India Institute of Medical Sciences, Jodhpur, India. ⁷⁹Department of Community Medicine, Datta Meghe Institute of Medical Sciences, Sawangi, India. ⁸⁰Saveetha Medical College, Saveetha University, Chennai, India. ⁸¹James P Grant School of Public Health, BRAC University, Dhaka, Bangladesh. ⁸²Department of Epidemiology and Biostatistics, University of South Carolina, Columbia, SC, USA. ⁸³Center for Biomedicine and Community Health, VNU-International School, Hanoi, Vietnam. ⁸⁴Department of Psychology, University of the Free State, Park West, South Africa. ⁸⁵Environmental Health Department, Imam Abdulrahman Bin Faisal University, Dammam, Saudi Arabia. ⁸⁶IRCCS Istituto Ortopedico Galeazzi (Galeazzi Orthopedic Institute IRCCS), University of Milan, Milan, Italy. ⁸⁷Department of Dermatology, Case Western Reserve University, Cleveland, OH, USA. ⁸⁸Department of Population and Health, University of Cape Coast, Cape Coast, Ghana. ⁸⁹Department of Information Technology, University of Human Development, Sulaymaniyah, Iraq. ⁹⁰Institute for Research and Innovation in Health, University of Porto, Porto, Portugal. ⁹¹Institute of Biomedical Engineering (INEB), University of Porto, Porto, Portugal. ⁹²Australian Institute for Suicide Research and Prevention, Griffith University, Mount Gravatt, QLD, Australia. ⁹³Wellcome Trust Brighton and Sussex Centre for Global Health Research, Brighton and Sussex Medical School, Brighton, UK. ⁹⁴School of Public Health, Addis Ababa University, Addis Ababa, Ethiopia. ⁹⁵National Centre for AIDS and STD Control, Save the Children, Kathmandu, Nepal. ⁹⁶Department of Global Public Health, Karolinska Institute, Stockholm, Sweden. ⁹⁷Department of Community Medicine, University of Peradeniya, Peradeniya, Sri Lanka. ⁹⁸Department of Health Metrics Sciences, School of Medicine, University of Washington, Seattle, WA, USA. ⁹⁹Department of Epidemiology, Maastricht University, Maastricht, Netherlands. ¹⁰⁰Department of Epidemiology, Shiraz University of Medical Sciences, Shiraz, Iran. ¹⁰¹Center of Complexity Sciences, National Autonomous University of Mexico, Mexico City, Mexico. ¹⁰²Faculty of Veterinary Medicine and Zootechnics, Autonomous University of Sinaloa, Rosales, Culiacán, Mexico. ¹⁰³Department of Community Medicine and Public Health, Urmia University of Medical Science, Urmia, Iran. ¹⁰⁴Development of Research and Technology Center, Ministry of Health and Medical Education, Tehran, Iran. ¹⁰⁵Department of Medical Laboratory Sciences, Iran University of Medical Sciences, Tehran, Iran. ¹⁰⁶Institute of Microbiology and Immunology, University of Belgrade, Belgrade, Serbia. ¹⁰⁷School of Public Health, Hawassa University, Hawassa, Ethiopia. ¹⁰⁸School of Public Health, Curtin University, Perth, WA, Australia. ¹⁰⁹Pediatric Dentistry and Dental Public Health Department, Alexandria University, Alexandria, Egypt. ¹¹⁰Department of Neurology, Cairo University, Cairo, Egypt. ¹¹¹Preventive Medicine and Public Health Research Center, Iran University of Medical Sciences, Tehran, Iran. ¹¹²Multiple Sclerosis Research Center, Tehran University of Medical Sciences, Tehran, Iran. ¹¹³Faculty of Health, York University, Toronto, BC, Canada. ¹¹⁴Institute for Health Science Research Germans Trias i Pujol, Autonomous University of Barcelona, Badalona, Spain. ¹¹⁵IrsiCaixa AIDS Research Institute, Badalona, Spain. ¹¹⁶Department of Virology, Pasteur Institute of Morocco, Casablanca, Morocco. ¹¹⁷Non-communicable Diseases Research Center, Tehran University of Medical Sciences, Tehran, Iran. ¹¹⁸Research Center for Environmental Determinants of Health, Kermanshah University of Medical Sciences, Kermanshah, Iran. ¹¹⁹Torrens University Australia, Adelaide, SA, Australia. ¹²⁰Institute of Resource Governance and Social Change, Kupang, Indonesia. ¹²¹Associated Laboratory for Green Chemistry (LAQV), University of Porto, Porto, Portugal. ¹²²Psychiatry Department, Kaiser Permanente, Fontana, CA, USA. ¹²³School of Health Sciences, A.T. Still University, Mesa, AZ, USA. ¹²⁴Institute of Public Health, Charité Universitätsmedizin Berlin (Charité Medical University Berlin), Berlin, Germany. ¹²⁵Institute of Gerontology, National Academy of Medical Sciences of Ukraine, Kyiv, Ukraine. ¹²⁶Department of Medical Parasitology, Abadan University of Medical Sciences, Abadan, Iran. ¹²⁷Faculty of Medicine, Abadan University of Medical Sciences, Abadan, Iran. ¹²⁸Department of Dermatology, Kobe University, Kobe, Japan. ¹²⁹Department of Cardiovascular Medicine, Cleveland Clinic, Cleveland, OH, USA. ¹³⁰Gillings School of Global Public Health, University of North Carolina Chapel Hill, Chapel Hill, NC, USA. ¹³¹Department of Community Medicine, Datta Meghe Institute of Medical Sciences, Wardha, India. ¹³²Department of Nursing and Midwifery, Addis Ababa University, Addis Ababa, Ethiopia. ¹³³Department of Public Health, Debre Berhan University, Debre Berhan, Ethiopia. ¹³⁴Infectious Disease Research Center, Kermanshah University of Medical Sciences, Kermanshah, Iran. ¹³⁵Pediatric Department, Kermanshah University of Medical Sciences, Kermanshah, Iran. ¹³⁶School of Public Health, Qazvin University of Medical Sciences, Qazvin, Iran. ¹³⁷Health Systems and Policy Research, Indian Institute of Public Health, Gandhinagar, India. ¹³⁸Department of Family and Community Medicine, University Of Sulaimani, Sulaimani, Iraq. ¹³⁹Department of Microbiology, Saint James School of Medicine, The Valley, Anguilla. ¹⁴⁰Department of Epidemiology, Saint James School of Medicine, The Valley, Anguilla. ¹⁴¹Faculty of Nursing, Federal University of Goiás, Goiânia, Brazil. ¹⁴²Department of Health Policy and Management, University of Georgia, Athens, GA, USA. ¹⁴³Department of Pharmacology, Shahid Beheshti University of Medical Sciences, Tehran, Iran. ¹⁴⁴Obesity Research Center, Shahid Beheshti University of Medical Sciences, Tehran, Iran. ¹⁴⁵School of Health and Environmental Studies, Hamdan Bin Mohammed Smart University, Dubai, United Arab Emirates. ¹⁴⁶Department of Public Health, Jigjiga University, Jigjiga, Ethiopia. ¹⁴⁷Gastrointestinal and Liver Diseases Research Center, Guilan University of Medical Sciences, Rasht, Iran. ¹⁴⁸Caspian Digestive Disease Research Center, Guilan University of Medical Sciences, Rasht, Iran. ¹⁴⁹School of Nursing and Midwifery, Tabriz University of Medical Sciences, Tabriz, Iran. ¹⁵⁰Independent Consultant, Tabriz, Iran. ¹⁵¹Institute of Pharmaceutical Sciences, University of Veterinary and Animal Sciences, Lahore, Pakistan. ¹⁵²Department of Pharmacy Administration and Clinical Pharmacy, Xian Jiaotong University, Xian, China. ¹⁵³Department of Statistics and Econometrics, Bucharest University of Economic Studies, Bucharest, Romania. ¹⁵⁴School of Business, London South Bank University, London, UK. ¹⁵⁵Department of Urban Planning and Design, University of Hong Kong, Hong Kong, China. ¹⁵⁶Department of Epidemiology and Biostatistics, Tehran University of Medical Sciences, Tehran, Iran. ¹⁵⁷Pediatric Chronic Kidney Disease Research Center, Tehran University of Medical Sciences, Tehran, Iran. ¹⁵⁸Institute of Research and Development, Duy Tan University, Da Nang, Vietnam. ¹⁵⁹Department of Computer Science, University of Human Development, Sulaymaniyah, Iraq. ¹⁶⁰Department of Occupational Safety and Health, China Medical University, Taichung, Taiwan. ¹⁶¹Department of Health Promotion and Education, University of Ibadan, Ibadan, Nigeria. ¹⁶²Department of Community Medicine,

- University of Ibadan, Ibadan, Nigeria. ¹⁶³Department of Community Medicine, University College Hospital, Ibadan, Ibadan, Nigeria. ¹⁶⁴Faculty of Medicine, University of Belgrade, Belgrade, Serbia. ¹⁶⁵Department of Epidemiology, University of Kragujevac, Kragujevac, Serbia. ¹⁶⁶Department of Epidemiology and Preventive Medicine, Monash University, Melbourne, VIC, Australia. ¹⁶⁷School of Health Systems and Public Health, University of Pretoria, Pretoria, South Africa. ¹⁶⁸Institute of Advanced Manufacturing Technologies, Peter the Great St. Petersburg Polytechnic University, St. Petersburg, Russia. ¹⁶⁹Institute of Comparative Economic Studies, Hosei University, Tokyo, Japan. ¹⁷⁰Department of Community Medicine, Dr. Baba Saheb Ambedkar Medical College & Hospital, Delhi, India. ¹⁷¹Department of Community Medicine, Banaras Hindu University, Varanasi, India. ¹⁷²Vanke School of Public Health, Tsinghua University, Beijing, China. ¹⁷³Department of Community Medicine, Manipal Academy of Higher Education, Mangalore, India. ¹⁷⁴National Institute of Epidemiology, Indian Council of Medical Research, Chennai, India. ¹⁷⁵Department of Family Medicine and Public Health, University of Opole, Opole, Poland. ¹⁷⁶School of Management and Medical Informatics, Tabriz University of Medical Sciences, Tabriz, Iran. ¹⁷⁷Institute for Prevention of Non-communicable Diseases, Qazvin University of Medical Sciences, Qazvin, Iran. ¹⁷⁸Health Services Management Department, Qazvin University of Medical Sciences, Qazvin, Iran. ¹⁷⁹Department of Biostatistics, Abadan University of Medical Sciences, Abadan, Iran. ¹⁸⁰Department of Forensic Medicine and Toxicology, All India Institute of Medical Sciences, Jodhpur, India. ¹⁸¹Social Determinants of Health Research Center, Tabriz University of Medical Sciences, Tabriz, Iran. ¹⁸²International Research Center of Excellence, Institute of Human Virology Nigeria, Abuja, Nigeria. ¹⁸³Julius Centre for Health Sciences and Primary Care, Utrecht University, Utrecht, Netherlands. ¹⁸⁴Mashhad University of Medical Sciences, Mashhad, Iran. ¹⁸⁵Department of Epidemiology and Biostatistics, Health Services Academy, Islamabad, Pakistan. ¹⁸⁶Department of Medical Microbiology & Immunology, United Arab Emirates University, Al Ain, United Arab Emirates. ¹⁸⁷Department of Population Science, Jatiya Kabi Kazi Nazrul Islam University, Mymensingh, Bangladesh. ¹⁸⁸Faculty of Health and Wellbeing, Sheffield Hallam University, Sheffield, UK. ¹⁸⁹College of Arts and Sciences, Ohio University, Zanesville, OH, USA. ¹⁹⁰Department of Public Health, New Mexico State University, Las Cruces, NM, USA. ¹⁹¹School of Traditional Chinese Medicine, Xiamen University Malaysia, Sepang, Malaysia. ¹⁹²School of Health Sciences, Kristiania University College, Oslo, Norway. ¹⁹³Department of Global Community Health and Behavioral Sciences, Tulane University, New Orleans, LA, USA. ¹⁹⁴Department of Nursing and Health Promotion, Oslo Metropolitan University, Oslo, Norway. ¹⁹⁵School of Population and Public Health, University of British Columbia, Vancouver, BC, Canada. ¹⁹⁶Arthritis Research Canada, Richmond, Canada. ¹⁹⁷Independent Consultant, Jakarta, Indonesia. ¹⁹⁸Kasturba Medical College, Udupi, India. ¹⁹⁹Biomedical Research Networking Center for Mental Health Network (CIBERSAM), San Juan de Dios Sanitary Park, Sant Boi de Llobregat, Spain. ²⁰⁰Catalan Institution for Research and Advanced Studies (ICREA), Barcelona, Spain. ²⁰¹Department of Anthropology, Panjab University, Chandigarh, India. ²⁰²Department of Demography, University of Montreal, Montreal, QC, Canada. ²⁰³Department of Social and Preventive Medicine, University of Montreal, Montreal, Canada. ²⁰⁴University of Environment and Sustainable Development, Somanya, Ghana. ²⁰⁵Department of Psychiatry, University of Nairobi, Nairobi, Kenya. ²⁰⁶Division of Psychology and Language Sciences, University College London, London, UK. ²⁰⁷Imperial College Business School, Imperial College London, London, UK. ²⁰⁸Faculty of Public Health, University of Indonesia, Depok, Indonesia. ²⁰⁹Department of Clinical Sciences and Community Health, University of Milan, Milan, Italy. ²¹⁰Public Health Foundation of India, Gurugram, India. ²¹¹Unit of Genetics and Public Health, Institute of Medical Sciences, Las Tablas, Panama. ²¹²Ministry of Health, Herrera, Panama. ²¹³Department of Infectious Disease Epidemiology, London School of Hygiene & Tropical Medicine, London, UK. ²¹⁴Department of Otorhinolaryngology, Father Muller Medical College, Mangalore, India. ²¹⁵Department of Health Sciences, University of Leicester, Leicester, UK. ²¹⁶School of Public Health and Preventive Medicine, Monash University, Melbourne, VIC, Australia. ²¹⁷Lerner Research Institute, Cleveland Clinic, Cleveland, OH, USA. ²¹⁸Department of Quantitative Health Science, Case Western Reserve University, Cleveland, OH, USA. ²¹⁹Department of Environmental Health Engineering, Tehran University of Medical Sciences, Tehran, Iran. ²²⁰Environmental Health Research Center, Kurdistan University of Medical Sciences, Sanandaj, Iran. ²²¹Department of Pediatrics, Montefiore Medical Center, New York, NY, USA. ²²²Department of Environmental Medicine and Public Health, Icahn School of Medicine at Mount Sinai, New York, NY, USA. ²²³Campus Caucaia, Federal Institute of Education, Science and Technology of Ceará, Caucaia, Brazil. ²²⁴Peru Country Office, United Nations Population Fund (UNFPA), Lima, Peru. ²²⁵Forensic Medicine Division, Imam Abdulrahman Bin Faisal University, Dammam, Saudi Arabia. ²²⁶Department of Reproductive Health and Population Studies, Bahir Dar University, Bahir Dar, Ethiopia. ²²⁷Breast Surgery Unit, Helsinki University Hospital, Helsinki, Finland. ²²⁸University of Helsinki, Helsinki, Finland. ²²⁹Clinical Microbiology and Parasitology Unit, Dr. Zora Profozic Polyclinic, Zagreb, Croatia. ²³⁰University Centre Varazdin, University North, Varazdin, Croatia. ²³¹Institute of Addiction Research (ISFF), Frankfurt University of Applied Sciences, Frankfurt, Germany. ²³²Department of Radiation Oncology, University of California San Francisco, San Francisco, CA, USA. ²³³Internal Medicine Department, King Saud University, Riyadh, Saudi Arabia. ²³⁴Department of Epidemiology and Biostatistics, Shahrekord University of Medical Sciences, Shahrekord, Iran. ²³⁵Department of Nursing, Mashhad University of Medical Sciences, Mashhad, Iran. ²³⁶Department of Biomolecular Sciences, University of Mississippi, Oxford, MS, USA. ²³⁷Department of Pharmacy, Mizan-Tepi University, Mizan, Ethiopia. ²³⁸Health Systems and Policy Research Unit, Ahmadu Bello University, Zaria, Nigeria. ²³⁹Department of Health Care Management, Technical University of Berlin, Berlin, Germany. ²⁴⁰Computer, Electrical, and Mathematical Sciences and Engineering Division, King Abdullah University of Science and Technology, Thuwal, Saudi Arabia. ²⁴¹Department of Epidemiology and Biostatistics, Wuhan University, Wuhan, China. ²⁴²Department of Pediatrics and Child Health, Debre Berhan University, Debre Berhan, Ethiopia. ²⁴³College of Medicine and Public Health, Flinders University, Adelaide, SA, Australia. ²⁴⁴Research and Analytics Department, Initiative for Financing Health and Human Development, Chennai, India. ²⁴⁵Department of Research and Analytics, Bioinsilico Technologies, Chennai, India. ²⁴⁶Laboratory of Public Health Indicators Analysis and Health Digitalization, Moscow Institute of Physics and Technology, Dolgoprudny, Russia. ²⁴⁷Experimental Surgery and Oncology Laboratory, Kursk State Medical University, Kursk, Russia. ²⁴⁸Department of Biotechnology, University of Central Punjab, Lahore, Pakistan. ²⁴⁹Department of Pediatrics, Arak University of Medical Sciences, Arak, Iran. ²⁵⁰Department of Disease Control and Environmental Health, Makerere University, Kampala, Uganda. ²⁵¹Department of General Surgery, Carol Davila University of Medicine and Pharmacy, Bucharest, Romania. ²⁵²Department of General Surgery, Emergency Hospital of Bucharest, Bucharest, Romania. ²⁵³Ministry of Health, Community Development, Gender, Elderly and Children, Dar es Salaam, Tanzania. ²⁵⁴Department of Public Health, University of Yaoundé I, Yaoundé, Cameroon. ²⁵⁵Department of Biological Sciences, University of Embu, Embu, Kenya. ²⁵⁶Institute for Global Health Innovations, Duy Tan University, Hanoi, Vietnam. ²⁵⁷South African Medical Research Council, Cape Town, South Africa. ²⁵⁸School of Public Health and Family Medicine, University of Cape Town, Cape Town, South Africa. ²⁵⁹Centre for Heart Rhythm Disorders, University of Adelaide, Adelaide, SA, Australia. ²⁶⁰Unit of Microbiology and Public Health, Institute of Medical Sciences, Las Tablas, Panama. ²⁶¹Department of Public Health, Ministry of Health, Herrera, Panama. ²⁶²Department of Pediatrics, National Hospital, Abuja, Nigeria. ²⁶³Department of International Public Health, University of Liverpool, Liverpool, UK. ²⁶⁴Administrative and Economic Sciences Department, University of Bucharest, Bucharest, Romania. ²⁶⁵Department of Community Health and Primary Care, University of Lagos, Idi Araba, Nigeria. ²⁶⁶Department of Family and Preventive Medicine, University of Utah, Salt Lake City, UT, USA. ²⁶⁷Department of Psychiatry and Behavioural Neurosciences, McMaster University, Hamilton, ON, Canada. ²⁶⁸Department of Psychiatry, University of Lagos, Lagos, Nigeria. ²⁶⁹Community Prevention and Care Services, National AIDS Control Committee, Abuja, Nigeria. ²⁷⁰Centre for Healthy Start Initiative, Lagos, Nigeria. ²⁷¹Diplomacy and Public Relations Department, University of Human Development, Sulaymaniyah, Iraq. ²⁷²Department of Pharmacology and Therapeutics, University of Nigeria Nsukka, Enugu, Nigeria. ²⁷³University of Port Harcourt, Port Harcourt, Nigeria. ²⁷⁴Department of Project Management, National Research University Higher School of Economics, Moscow, Russia. ²⁷⁵Department of Medicine, University of Ibadan, Ibadan, Nigeria. ²⁷⁶Department of Medicine, University College Hospital, Ibadan, Ibadan, Nigeria. ²⁷⁷Department of Respiratory Medicine, Jagadguru Sri Shivarathreeswara Academy of Health Education and Research, Mysore, India. ²⁷⁸Department of Forensic Medicine and Toxicology, Manipal Academy of Higher Education, Manipal, India. ²⁷⁹Department of Health Metrics, Center for Health Outcomes & Evaluation, Bucharest, Romania. ²⁸⁰Research Department, Nepal Health Research Council, Kathmandu, Nepal. ²⁸¹Research Department, Public Health Research Society Nepal, Kathmandu,

- Nepal. ^{282c}Saveetha Medical College and Hospitals, Saveetha University, Chennai, India. ²⁸³Iran University of Medical Sciences, Tehran, Iran.
- ²⁸⁴Department of Pediatrics, University of Melbourne, Melbourne, VIC, Australia. ²⁸⁵Population Health Theme, Murdoch Children's Research Institute, Melbourne, VIC, Australia. ²⁸⁶Department of Genetics, Yale University, New Haven, CT, USA. ²⁸⁷School of Global Public Health, New York University, New York, NY, USA. ²⁸⁸University Medical Center Groningen, University of Groningen, Groningen, Netherlands. ²⁸⁹School of Economics and Business, University of Groningen, Groningen, Netherlands. ²⁹⁰National Institute of Infectious Diseases, Bucuresti, Romania. ²⁹¹Department of Infectious Diseases, Carol Davila University of Medicine and Pharmacy, Bucharest, Romania. ²⁹²School of Engineering, Macquarie University, Sydney, NSW, Australia. ²⁹³Pohang University of Science and Technology, Pohang, South Korea. ²⁹⁴College of Medicine, University of Central Florida, Orlando, FL, USA. ²⁹⁵Department of Immunology, Mazandaran University of Medical Sciences, Sari, Iran. ²⁹⁶Molecular and Cell Biology Research Center, Mazandaran University of Medical Sciences, Sari, Iran. ²⁹⁷Metabolomics and Genomics Research Center, Tehran University of Medical Sciences, Tehran, Iran. ²⁹⁸Sina Trauma and Surgery Research Center, Tehran University of Medical Sciences, Tehran, Iran. ²⁹⁹Future Technology Research Center, National Yunlin University of Science and Technology, Yunlin, Taiwan. ³⁰⁰Department of Epidemiology, Biostatistics and Occupational Health, McGill University, Montreal, QC, Canada. ³⁰¹Research and Innovation Division, South Asian Institute for Social Transformation (SAIST), Dhaka, Bangladesh. ³⁰²Research Department, Policy Research Institute, Kathmandu, Nepal. ³⁰³Health and Public Policy Department, Global Center for Research and Development, Kathmandu, Nepal. ³⁰⁴Department of Oral Pathology, Sharavathi Dental College and Hospital, Shimogga, India. ³⁰⁵WHO Collaborating Centre for Public Health Education and Training, Imperial College London, London, UK. ³⁰⁶University College London Hospitals, London, UK. ³⁰⁷Department of Primary Care and Public Health, Imperial College London, London, UK. ³⁰⁸Academic Public Health England, Public Health England, London, UK. ³⁰⁹Department of Computer Science, Boston University, Boston, MA, USA. ³¹⁰Department of Epidemiology and Biostatistics, Haramaya University, Harar, Ethiopia. ³¹¹Research Center for Immunodeficiencies, Tehran University of Medical Sciences, Tehran, Iran. ³¹²Network of Immunity in Infection, Malignancy and Autoimmunity (NIIMA), Universal Scientific Education and Research Network (USERN), Tehran, Iran. ³¹³Faculty of Business and Management, Universiti Sultan Zainal Abidin (Sultan Zainal Abidin University), Kuala Terengganu, Malaysia. ³¹⁴Epidemiology Research Unit (EPIUnit), University of Porto, Porto, Portugal. ³¹⁵Department of Global Health, University of Washington, Seattle, WA, USA. ³¹⁶Department of Medicine, University of Washington, Seattle, WA, USA. ³¹⁷African Genome Center, Mohammed VI Polytechnic University (UM6P), Ben Guerir, Morocco. ³¹⁸Centro de Investigaciones en Anomalías Congénitas y Enfermedades Raras (Center for Research in Congenital Anomalies and Rare Diseases), Universidad ICESI (ICESI University), Cali, Colombia. ³¹⁹Malaria Atlas Project, Telethon Kids Institute, Perth, Australia. ³²⁰Department of Health Statistics, National Institute for Medical Research, Dar es Salaam, Tanzania. ³²¹Department of Internal Medicine, University of Botswana, Gaborone, Botswana. ³²²Department of Psychiatry, All India Institute of Medical Sciences, New Delhi, India. ³²³Department of Public Health, Madda Walabu University, Bale Robe, Ethiopia. ³²⁴Public Health and Community Medicine Department, Cairo University, Giza, Egypt. ³²⁵Drug Applied Research Center, Tabriz University of Medical Sciences, Tabriz, Iran. ³²⁶Department of Entomology, Ain Shams University, Cairo, Egypt. ³²⁷Centre for Tropical Medicine and Global Health, University of Oxford, Oxford, UK. ³²⁸Nuffield Department of Medicine, University of Oxford, Oxford, UK. ³²⁹Geriatric and Long Term Care Department, Hamad Medical Corporation, Doha, Qatar. ³³⁰Faculty of Health & Social Sciences, Bournemouth University, Bournemouth, UK. ³³¹College of Public Health, Medical and Veterinary Sciences, James Cook University, QLD, Townsville, Australia. ³³²Public Health Division, An-Najah National University, Nablus, Palestine. ³³³Independent Consultant, Karachi, Pakistan. ³³⁴Faculty of Caring Science, Work Life, and Social Welfare, University of Borås, Borås, Sweden. ³³⁵College of Medicine, Yonsei University, Seoul, South Korea. ³³⁶Department of Internal Medicine, Yale University, New Haven, CT, USA. ³³⁷School of Medicine, University of Alabama at Birmingham, Birmingham, AL, USA. ³³⁸Medicine Service, US Department of Veterans Affairs (VA), Birmingham, AL, USA. ³³⁹Department No.16, Moscow Research and Practical Centre on Addictions, Moscow, Russia. ³⁴⁰Department of Infectious Diseases and Epidemiology, Pirogov Russian National Research Medical University, Moscow, Russia. ³⁴¹Department of Community Medicine, Ahmadu Bello University, Zaria, Nigeria. ³⁴²Cancer Control Center, Osaka International Cancer Institute, Osaka, Japan. ³⁴³Department of Biomedical Sciences, Arba Minch University, Arba Minch, Ethiopia. ³⁴⁴University Institute "Egas Moniz", Monte da Caparica, Portugal. ³⁴⁵Research Institute for Medicines, University of Lisbon, Lisbon, Portugal. ³⁴⁶School of Public Health, Mekelle University, Mekelle, Ethiopia. ³⁴⁷Southgate Institute for Health and Society, Flinders University, Adelaide, SA, Australia. ³⁴⁸Department of Pathology and Legal Medicine, University of São Paulo, Ribeirão Preto, Brazil. ³⁴⁹Modestum LTD, London, UK. ³⁵⁰College of Medicine and Health Sciences, Bahir Dar University, Bahir Dar, Ethiopia. ³⁵¹Department of Community Medicine, Alex Ekwueme Federal University Teaching Hospital Abakaliki, Abakaliki, Nigeria. ³⁵²Kasturba Medical College, Manipal Academy of Higher Education, Mangalore, India. ³⁵³Clinical Research Department, IRCCS Fondazione Don Carlo Gnocchi, Milan, Italy. ³⁵⁴Department of Medical and Surgical Sciences, University of Bologna, Bologna, Italy. ³⁵⁵Occupational Health Unit, Sant'Orsola Malpighi Hospital, Bologna, Italy. ³⁵⁶Faculty of Information Technology, HUTECH University, Ho Chi Minh City, Vietnam. ³⁵⁷Center of Excellence in Behavioral Medicine, Nguyen Tat Thanh University, Ho Chi Minh City, Vietnam. ³⁵⁸Population Dynamics and Sexual and Reproductive Health, African Population and Health Research Center, Nairobi, Kenya. ³⁵⁹Foundation University Medical College, Foundation University Islamabad, Islamabad, Pakistan. ³⁶⁰Department of Cultures, Societies and Global Studies, Northeastern University, Boston, MA, USA. ³⁶¹School of Public Health, University of Nairobi, Nairobi, Kenya. ³⁶²School of Population Health and Environmental Sciences, King's College London, London, UK. ³⁶³Centre for Health Policy Research, Torrens University Australia, Adelaide, SA, Australia. ³⁶⁴Department of community Medicine, Rajarata University of Sri Lanka, Anuradhapura, Sri Lanka. ³⁶⁵Department of Biostatistics, University of Washington, Seattle, WA, USA. ³⁶⁶School of International Development and Global Studies, University of Ottawa, Ottawa, ON, Canada. ³⁶⁷The George Institute for Global Health, University of Oxford, Oxford, UK. ³⁶⁸Centre for Suicide Research and Prevention, University of Hong Kong, Hong Kong, China. ³⁶⁹Department of Social Work and Social Administration, University of Hong Kong, Hong Kong, China. ³⁷⁰Department of Neuropsychopharmacology, National Center of Neurology and Psychiatry, Kodaira, Japan. ³⁷¹Department of Public Health, Juntendo University, Tokyo, Japan. ³⁷²Department of Bioengineering and Therapeutic Sciences, University of California San Francisco, San Francisco, CA, USA. ³⁷³Addictology Department, Russian Medical Academy of Continuous Professional Education, Moscow, Russia. ³⁷⁴School of Public Health, Wuhan University of Science and Technology, Wuhan, China. ³⁷⁵Hubei Province Key Laboratory of Occupational Hazard Identification and Control, Wuhan University of Science and Technology, Wuhan, China. ³⁷⁶School of Medicine, Wuhan University, Wuhan, China.

Received: 26 April 2022 Accepted: 26 October 2022

Published online: 19 December 2022

References

- Frank TD, Carter A, Jahagirdar D, Biehle MH, Douwes-Schultz D, Larson SL, et al. Global, regional, and national incidence, prevalence, and mortality of HIV, 1980–2017, and forecasts to 2030, for 195 countries and territories: a systematic analysis for the Global Burden of Diseases, Injuries, and Risk Factors Study 2017. *Lancet HIV*. 2019;6:e831–59.
- GBD 2019 Diseases and Injuries Collaborators. Global burden of 369 diseases and injuries in 204 countries and territories, 1990–2019: a systematic analysis for the Global Burden of Disease Study 2019. *Lancet*. 2020;396:1204–22.
- Micah AE, Su Y, Bachmeier SD, Chapin A, Cogswell IE, Crosby SW, et al. Health sector spending and spending on HIV/AIDS, tuberculosis, and malaria, and development assistance for health: progress towards Sustainable Development Goal 3. *Lancet*. 2020;396:693–724.
- UNAIDS. Joint United Nations Programme on HIV/AIDS. 2020. <https://www.unaids.org/en>.
- UNAIDS. Understanding Fast Track: accelerating action to end the AIDS epidemic by 2030. 2015. http://www.unaids.org/sites/default/files/media_asset/201506_JC2743_Understanding_FastTrack_en.pdf.

6. United Nations. Transforming our world: the 2030 agenda for sustainable development. New York: United Nations; 2015.
7. Bekker L-G, Alleyne G, Baral S, Cepeda J, Daskalakis D, Dowdy D, et al. Advancing global health and strengthening the HIV response in the era of the Sustainable Development Goals: the International AIDS Society—Lancet Commission. *Lancet*. 2018;392:312–58.
8. Jones J, Sullivan PS, Curran JW. Progress in the HIV epidemic: identifying goals and measuring success. *PLoS Med*. 2019;16:e1002729.
9. Hogan AB, Jewell BL, Sherrard-Smith E, Vesga JF, Watson OJ, Whittaker C, et al. Potential impact of the COVID-19 pandemic on HIV, tuberculosis, and malaria in low-income and middle-income countries: a modelling study. *Lancet Glob Health*. 2020;8:e1132–41.
10. Dwyer-Lindgren L, Cork MA, Sligar A, Steuben KM, Wilson KF, Provost NR, et al. Mapping HIV prevalence in sub-Saharan Africa between 2000 and 2017. *Nature*. 2019;570:189–93.
11. Larmarange J, Bendaub V. HIV estimates at second subnational level from national population-based surveys. *AIDS*. 2014;28(Suppl 4):S469–76.
12. Meyer-Rath G, McGillen JB, Cuadros DF, Hallett TB, Bhatt S, Wabiri N, et al. Targeting the right interventions to the right people and places: the role of geospatial analysis in HIV program planning. *AIDS*. 2018;32:957–63.
13. Cuadros DF, Li J, Branscum AJ, Akullian A, Jia P, Mziray EN, et al. Mapping the spatial variability of HIV infection in sub-Saharan Africa: effective information for localized HIV prevention and control. *Sci Rep*. 2017;7:9093.
14. Coburn BJ, Okano JT, Blower S. Using geospatial mapping to design HIV elimination strategies for sub-Saharan Africa. *Sci Transl Med*. 2017;9:eaag0019.
15. Akullian A, Vandormael A, Miller JC, Bershteyn A, Wenger E, Cuadros D, et al. Large age shifts in HIV-1 incidence patterns in KwaZulu-Natal, South Africa. *Proc Natl Acad Sci U S A*. 2021;118:e2013164118.
16. Khalifa A, Stover J, Mahy M, Idele P, Porth T, Lwamba C. Demographic change and HIV epidemic projections to 2050 for adolescents and young people aged 15–24. *Null*. 2019;12:1662685.
17. Faust L, Yaya S. The effect of HIV educational interventions on HIV-related knowledge, condom use, and HIV incidence in sub-Saharan Africa: a systematic review and meta-analysis. *BMC Public Health*. 2018;18:1254.
18. Anderson S-J, Cherutich P, Kilongo N, Cremin I, Fecht D, Kimanga D, et al. Maximising the effect of combination HIV prevention through prioritisation of the people and places in greatest need: a modelling study. *Lancet*. 2014;384:249–56.
19. Schneider MT, Birger M, Haakenstad A, Singh L, Hamavid H, Chapin A, et al. Tracking development assistance for HIV/AIDS: the international response to a global epidemic. *AIDS*. 2016;30:1475–9.
20. Olakunde BO, Adeyinka DA, Ozigbu CE, Ogundipe T, Menson WNA, Olawepo JO, et al. Revisiting aid dependency for HIV programs in sub-Saharan Africa. *Public Health*. 2019;170:57–60.
21. Jewell BL, Mudimu E, Stover J, ten Brink D, Phillips AN, Smith JA, et al. Potential effects of disruption to HIV programmes in sub-Saharan Africa caused by COVID-19: results from multiple mathematical models. *Lancet HIV*. 2020;7:e629–40.
22. Nagelkerke NJD, Jha P, de Vlas SJ, Korenromp EL, Moses S, Blanchard JF, et al. Modelling HIV/AIDS epidemics in Botswana and India: impact of interventions to prevent transmission. *Bull World Health Organ*. 2002;80:89–96.
23. Long EF, Stavert RR. Portfolios of biomedical HIV interventions in South Africa: a cost-effectiveness analysis. *J Gen Intern Med*. 2013;28:1294–301.
24. Bershteyn A, Klein DJ, Eckhoff PA. Age-targeted HIV treatment and primary prevention as a “ring fence” to efficiently interrupt the age patterns of transmission in generalized epidemic settings in South Africa. *Int Health*. 2016;8:277–85.
25. Joint United Nations Programme on HIV/AIDS. *AIDSinfo*. UNAIDS; 2018. <http://aidsinfo.unaids.org/>. Accessed 18 June 2020.
26. Okano JT, Blower S. Sex-specific maps of HIV epidemics in sub-Saharan Africa. *Lancet Infect Dis*. 2016;16:1320–2.
27. Messina JP, Emch M, Muwonga J, Mwangalirwa K, Edidi SB, Mama N, et al. Spatial and socio-behavioral patterns of HIV prevalence in the Democratic Republic of Congo. *Soc Sci Med*. 2010;71:1428–35.
28. Palk L, Blower S. Geographic variation in sexual behavior can explain geospatial heterogeneity in the severity of the HIV epidemic in Malawi. *BMC Med*. 2018;16:22.
29. Tanser F, Barnighausen T, Cooke GS, Newell M-L. Localized spatial clustering of HIV infections in a widely disseminated rural South African epidemic. *Int J Epidemiol*. 2009;38:1008–16.
30. Bulstra CA, Hontelez JAC, Giardina F, Steen R, Nagelkerke NJD, Barnighausen T, et al. Mapping and characterising areas with high levels of HIV transmission in sub-Saharan Africa: a geospatial analysis of national survey data. *PLoS Med*. 2020;17:e1003042.
31. Stevens GA, Alkema L, Black RE, Boerma JT, Collins GS, Ezzati M, et al. Guidelines for accurate and transparent health estimates reporting: the GATHER statement. *Lancet*. 2016;388:e19–23.
32. Tatem AJ. WorldPop, open data for spatial demography. *Sci Data*. 2017;4:170004.
33. R Core Team. R: the R project for statistical computing. 2019. <https://www.r-project.org/>. Accessed 8 July 2020.
34. Burgert C, Colston J, Roy T, Zachary B. Geographic displacement procedure and georeferenced data release policy for the Demographic and Health Surveys; 2013.
35. UNAIDS. National HIV estimates file. 2019. <https://www.unaids.org/en/dataanalysis/datatools/spectrum-epp>.
36. GBD 2019 Demographics Collaborators. Global age-sex-specific fertility, mortality, healthy life expectancy (HALE), and population estimates in 204 countries and territories, 1950–2019: a comprehensive demographic analysis for the Global Burden of Disease Study 2019. *Lancet*. 2020;396:1160–203.
37. Bhatt S, Cameron E, Flaxman SR, Weiss DJ, Smith DL, Gething PW. Improved prediction accuracy for disease risk mapping using Gaussian process stacked generalization. *J Royal Soc Interface*. 2017;14:20170520.
38. Hastie T, Tibshirani RJ. Generalized additive models. London: Chapman & Hall; 1990.
39. Friedman JH. Greedy function approximation: a gradient boosting machine. *Ann Stat*. 2001;29:1189–232.
40. Tibshirani R. Regression shrinkage and selection via the lasso. *J R Stat Soc B Methodol*. 1996;58:267–88.
41. Kristensen K, Nielsen A, Berg CW, Skaug H, Bell BM. TMB: automatic differentiation and Laplace approximation. *J Stat Softw*. 2016;70:1–21.
42. Stein ML. Interpolation of spatial data: some theory for Kriging. New York: Springer-Verlag; 1999.
43. Lindgren F, Rue H, Lindström J. An explicit link between Gaussian fields and Gaussian Markov random fields: the stochastic partial differential equation approach. *R Stat Soc*. 2011;73:423–98.
44. Zaba BW, Carpenter LM, Boerma JT, Gregson S, Nakiyingi J, Urassa M. Adjusting ante-natal clinic data for improved estimates of HIV prevalence among women in sub-Saharan Africa. *AIDS*. 2000;14:2741–50.
45. Gouws E, Mishra V, Fowler TB. Comparison of adult HIV prevalence from national population-based surveys and antenatal clinic surveillance in countries with generalised epidemics: implications for calibrating surveillance data. *Sex Transm Infect*. 2008;84(Suppl 1):i17–23.
46. Marsh K, Mahy M, Salomon JA, Hogan DR. Assessing and adjusting for differences between HIV prevalence estimates derived from national population-based surveys and antenatal care surveillance, with applications for Spectrum 2013. *AIDS*. 2014;28:S497–505.
47. Marston M, Zaba B, Eaton JW. The relationship between HIV and fertility in the era of antiretroviral therapy in sub-Saharan Africa: evidence from 49 Demographic and Health Surveys. *Trop Med Int Health*. 2017;22:1542–50.
48. ESA-CCI Project. Land cover classification gridded maps from 1992 to present derived from satellite observations. 2020. <https://cds.climate.copernicus.eu/cdsapp#!/dataset/satellite-land-cover?tab=overview>. Accessed 30 Apr 2020.
49. Wickham H. ggplot2: elegant graphics for data analysis. New York: Springer-Verlag; 2016.
50. McGillen JB, Anderson S-J, Dybul MR, Hallett TB. Optimum resource allocation to reduce HIV incidence across sub-Saharan Africa: a mathematical modelling study. *Lancet HIV*. 2016;3:e441–8.
51. Hosek S, Pettifor A. HIV prevention interventions for adolescents. *Curr HIV/AIDS Rep*. 2019;16:120–8.
52. Schatz E, Seeley J, Negin J, Weiss HA, Tumwekwase G, Kabunga E, et al. “For us here, we remind ourselves”: strategies and barriers to ART access and adherence among older Ugandans. *BMC Public Health*. 2019;19:131.
53. De Neve J-W, Garrison-Desany H, Andrews KG, Sharara N, Boudreaux C, Gill R, et al. Harmonization of community health worker programs

- for HIV: A four-country qualitative study in Southern Africa. *PLoS Med.* 2017;14:e1002374.
54. Mbonu NC, van den Borne B, De Vries NK. Stigma of people with HIV/AIDS in Sub-Saharan Africa: a literature review. *J Trop Med.* 2009;2009:145891.
 55. World Health Organization. Consolidated guidelines on the use of antiretroviral drugs for treating and preventing HIV infection: recommendations for a public health approach. 2013. https://apps.who.int/iris/bitstream/handle/10665/85321/9789241505727_eng.pdf;jsessionid=FA819AE8F065C685D2C4CC768FEE7304?sequence=1. Accessed 20 Mar 2021.
 56. Sabapathy K, den Bergh RV, Fidler S, Hayes R, Ford N. Uptake of home-based voluntary HIV testing in sub-Saharan Africa: a systematic review and meta-analysis. *PLoS Med.* 2012;9:e1001351.
 57. Sharma M, Ying R, Tarr G, Barnabas R. Systematic review and meta-analysis of community and facility-based HIV testing to address linkage to care gaps in sub-Saharan Africa. *Nature.* 2015;528:577–85.
 58. Hatzold K, Gudukeya S, Mutseta MN, Chilongosi R, Nalubamba M, Nkhoma C, et al. HIV self-testing: breaking the barriers to uptake of testing among men and adolescents in sub-Saharan Africa, experiences from STAR demonstration projects in Malawi, Zambia and Zimbabwe. *J Int AIDS Soc.* 2019;22:e25244.
 59. Sia D, Onadja Y, Hajizadeh M, Heymann SJ, Brewer TF, Nandi A. What explains gender inequalities in HIV/AIDS prevalence in sub-Saharan Africa? Evidence from the demographic and health surveys. *BMC Public Health.* 2016;16:1136.
 60. Dean HD, Fenton KA. Addressing social determinants of health in the prevention and control of HIV/AIDS, viral hepatitis, sexually transmitted infections, and tuberculosis. *Public Health Rep.* 2010;125(4_suppl):1–5.
 61. Heestermaans T, Browne JL, Aitken SC, Vervoort SC, Klipstein-Grobusch K. Determinants of adherence to antiretroviral therapy among HIV-positive adults in sub-Saharan Africa: a systematic review. *BMJ Glob Health.* 2016;1:e000125.
 62. Tanser F, Bärnighausen T, Grapsa E, Zaidi J, Newell M-L. High coverage of ART associated with decline in risk of HIV acquisition in rural KwaZulu-Natal, South Africa. *Science.* 2013;339:966–71.
 63. Vandormael A, Akullian A, Siedner M, de Oliveira T, Bärnighausen T, Tanser F. Declines in HIV incidence among men and women in a South African population-based cohort. *Nat Commun.* 2019;10:5482.
 64. Hontelez JAC, de Vlas SJ, Baltussen R, Newell M-L, Bakker R, Tanser F, et al. The impact of antiretroviral treatment on the age composition of the HIV epidemic in sub-Saharan Africa. *AIDS.* 2012;26(Suppl 1 0 1):S19–30.
 65. CASCADE Collaboration. Differences in CD4 cell counts at seroconversion and decline among 5739 HIV-1-infected individuals with well-estimated dates of seroconversion. *J Acquir Immune Defic Syndr.* 2003;34:76–83.
 66. Goetz MB, Boscardin WJ, Wiley D, Alkasspoles S. Decreased recovery of CD4 lymphocytes in older HIV-infected patients beginning highly active antiretroviral therapy. *AIDS.* 2001;15:1576–9.
 67. Mills EJ, Bärnighausen T, Negin J. HIV and aging—preparing for the challenges ahead. *N Engl J Med.* 2012;366:1270–3.
 68. Coetzee L, Bogler L, De Neve J-W, Bärnighausen T, Geldsetzer P, Vollmer S. HIV, antiretroviral therapy and non-communicable diseases in sub-Saharan Africa: empirical evidence from 44 countries over the period 2000 to 2016. *J Int AIDS Soc.* 2019;22:e25364.
 69. Parikh SM, Obuku EA, Walker SA, Semeere AS, Auerbach BJ, Hakim JG, et al. Clinical differences between younger and older adults with HIV/AIDS starting antiretroviral therapy in Uganda and Zimbabwe: a secondary analysis of the DART trial. *PLoS One.* 2013;8:e76158.
 70. Gething P, Tatem A, Bird T, Burgert-Brucker CR. Creating spatial interpolation surfaces with DHS data. Rockville: ICF International; 2015.
 71. Golding N, Burstein R, Longbottom J, Browne AJ, Fullman N, Osgood-Zimmerman A, et al. Mapping under-5 and neonatal mortality in Africa, 2000–15: a baseline analysis for the Sustainable Development Goals. *Lancet.* 2017;390:2171–82.
 72. Graetz N, Friedman J, Osgood-Zimmerman A, Burstein R, Biehl MH, Shields C, et al. Mapping local variation in educational attainment across Africa. *Nature.* 2018;555:48–53.
 73. Blondeel K, Say L, Chou D, Toskin I, Khosla R, Scolaro E, et al. Evidence and knowledge gaps on the disease burden in sexual and gender minorities: a review of systematic reviews. *Int J Equity Health.* 2016;15:16.
 74. Woldeesenbet S, Kufa T, Lombard C, Manda S, Ayalew kassahun, Cheyip M, et al. The 2017 national antenatal sentinel HIV survey key findings, South Africa. 2019.
 75. Ministry of Health, National AIDS Control Council. Kenya AIDS response progress report 2016. 2016.
 76. Jiang H, Zhou Y, Tang W. Maintaining HIV care during the COVID-19 pandemic. *Lancet HIV.* 2020;7:e308–9.
 77. Global Administrative Areas. GADM maps and data. v.3.6. 2019. <https://gadm.org/>.

Publisher's Note

Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.